Analysis of genome-wide, cancer-associated mutation datasets in mouse and human



Jenny Mattison Wellcome Trust Sanger Institute Trinity College University of Cambridge

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Declaration

This thesis describes work carried out between April 2005 and September 2008 under the supervision of Dr Tim Hubbard and Dr David Adams at the Wellcome Trust Sanger Institute, while member of Trinity College, University of Cambridge. This dissertation is the result of my own work and contains nothing that is the outcome of work done in collaboration with others, except as specified in the text and Acknowledgements. No part of this dissertation nor anything substantially the same has been, or is being, submitted for any qualification at this or any other university. This dissertation does not exceed the page limit set by the Biology Degree Committee.

Jenny Mattison September 2008

Abstract

The complexity of human cancer genomes complicates the identification of those mutations that drive the tumourigenic process. Integrative analyses, particularly cross-species comparisons, provide a means of distinguishing likely driver mutations from the background of passenger mutations that arise in unstable cancer genomes. This thesis describes the analysis of human and mouse experimental datasets to identify human cancer gene candidates.

In mice, candidate cancer genes can be 'tagged' using insertional mutagens such as retroviruses and transposons. The analysis of more than 1,000 mouse tumours generated by insertional mutagenesis is described. Insertion sites are mapped to the mouse genome and are used to identify candidate cancer genes. The distribution of insertions within and around candidate genes is analysed to predict the likely mechanisms of mutagenesis and, therefore, the possible structure and function of the mutated gene products. Candidates are also characterised by comparison with other human and mouse cancer-associated mutation datasets, and co-operating cancer genes are identified in an attempt to better understand cancer gene pathways.

The mouse insertional mutagenesis results are then compared to genome-wide copy number data for human cancers. The Wellcome Trust Sanger Institute has generated comparative genomic hybridisation (CGH) data for ~700 human cancer cell lines using the Affymetrix 10K SNP array and, more recently, for ~600 human cancer cell lines using the high resolution Affymetrix SNP 6.0 array. Regions of copy number change in human cancers often encompass many genes, and it can be difficult to determine which genes contribute to the cancer phenotype. In this thesis, the human CGH data are processed into regions of copy number change and the mouse candidate cancer genes identified by retroviral insertional mutagenesis are used to narrow down the candidates in amplicons and deletions. The over-representation of mouse candidate oncogenes in regions of copy number gain suggests that a significant proportion of genes contributing to retrovirus-induced tumourigenesis in the mouse are also amplified in, and contribute to the development of, human cancers. Candidate oncogenes and tumour suppressor genes that are recurrently mutated in both human tumours and murine lymphomas are identified as strong candidates for a role in tumourigenesis.

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Table of Contents

Chapter 1	Introduction	1
1.1 Ou	tline of introduction	1
1.2 An	introduction to cancer	1
1.2.1	Definition and classification	1
1.2.2	Epidemiology	2
1.2.3	The multi-stage theory of carcinogenesis	4
1.2.4	The hallmarks of cancer	7
1.2.5	Cancer genes	7
1.2.6	Pathways in cancer	10
1.2.7	Treatment of cancer	13
1.3 Ge	nome-wide approaches for human cancer gene discovery	14
1.3.1	Gene resequencing	14
1.3.2	Gene expression profiling	17
1.3.3	Copy number analysis	18
1.3.4	Epigenetic profiling	29
1.3.5	Genome-wide mapping of transcription factor binding sites	31
1.4 Ca	ncer gene discovery in the mouse	34
1.4.1	The mouse as a model for studying cancer	34
1.4.2	Forward genetic screens in the mouse	39
1.5 Cr	oss-species comparative analysis for cancer gene discovery	58
1.6 Aii	ns of this thesis	61
•	Identifying insertion sites and candidate cancer genes by insertional	
U	is in the mouse	
	roduction	
	scription of the datasets	
2.2.1	The retroviral dataset	
2.2.2	The Sleeping Beauty dataset	69
2.2.3	Known cancer genes in the Cancer Gene Census	
2.3 Ma	pping the sequence reads using SSAHA2	70
2.4 Ac	counting for unmapped reads	73
2.5 Fil	tering the mapped reads	79

2.6 Ide	ntification and filtering of insertion sites
2.7 Estimating the coverage of the mutagenesis screens	
2.8 An	alysis of the distribution of insertions around mouse genes
2.9 Ass	signing insertions to genes
2.10 Id	entifying statistically significant common insertion sites
2.10.1	Monte Carlo simulations
2.10.2	Kernel convolution
2.10.3	Final set of candidate genes
2.11 D	iscussion
•	Analysis of mouse candidate cancer genes identified by insertional is
0	
	roduction
	mparative analyses between the insertional mutagenesis data and other cancer-
	108
3.2.1	Description of the datasets
3.2.2	Comparison with insertional mutagenesis data
	mparison of candidate cancer genes in the MuLV and Sleeping Beauty
	termining the mechanisms of MuLV insertional mutagenesis 125
3.4.1	Analysing the distribution of intragenic insertions 125
3.4.2	Analysing co-occurring insertions in candidate genes disrupted by MuLV
and T2	2/Onc
3.4.3	Identification of tumour suppressor genes inactivated by MuLV 138
3.4.4	Identifying retroviral insertions in regulatory features
3.4.5	Expression analysis of MuLV-induced tumours
3.5 Ide	ntification of co-operating cancer genes in the MuLV dataset 148
3.5.1	Genotype-specific cancer genes
3.5.2	Co-occurrence and mutual exclusivity of disrupted genes 153
3.6 Dis	scussion
Chapter 4	Using mouse candidate cancer genes to narrow down the candidates in

regions of copy number change in human cancers 161 4.1 Introduction 161

		101
4.2	Description of the datasets	162

	4.2	2.1	Mouse candidate cancer genes identified by retroviral insertional	
	mu	itage	enesis	162
	4.2	2.2	Copy number data for human cancer cell lines	164
	4.2	2.3	Copy number variants (CNVs)	166
4	.3	Pro	cessing the copy number data	166
4	.4	Cha	aracterising gains and losses in cancer genomes	173
4	.5	Cor	mparative analysis of mouse candidate cancer genes and CGH data fr	om
h	uma	ın ca	incers	175
	4.5	5.1	Global comparison	175
	4.5	5.2	Identification of individual candidates for a role in human cancer	190
4	.6	Cor	mparison of methods for calling gains and losses	208
4	.7	Glo	bal comparison of mouse candidate cancer genes and human CNVs .	215
4	.8	Dis	cussion	
~		_		
	-		Identifying human cancer genes in high-resolution copy number	
5			oduction	
5	.2		scription and processing of the datasets	
	5.2		High-resolution copy number data	
	5.2		Additional datasets	
	.3		mparative analysis of human high-resolution CGH data versus mouse	
ir			al mutagenesis data	
	5.3		Global comparison	
	5.3		Identifying individual cancer gene candidates	
5	.4		mparison between high-resolution and 10K CGH data	
5			ntification of co-operating cancer genes	
	5.5	5.1	Genotype-specific cancer genes	
	5.5	5.2	Co-occurrence of amplified and deleted candidate cancer genes	
5	.6	Dis	cussion	273
Cha	nto	r 6	Summary and Conclusions	276
Clia	ipte	10	Summary and Conclusions	270
Ref	eren	ices		
App	end	lices	5	337
թոհ	lice	tion	l	365

List of Figures

1.1	Summary of cancer incidence in 2004 and deaths from cancer in 2005 for the	
most co	ommon sites of cancer in males and females in the UK.	3
1.2	The clonal evolution of cancer.	5
1.3	Mutations in different genes in the same pathway can have an equivalent	
effect.		11
1.4	Array design and whole-genome sampling assay for the Affymetrix SNP	
array.		20
1.5	End sequence profiling of tumour DNA.	28
1.6	Overview of ChIP-PET for mapping transcription factor binding sites.	32
1.7	Generation of a conditional knockout allele in ES cells.	37
1.8	Structure of a retroviral provirus.	41
1.9	The mechanisms of mutagenesis of murine leukaemia virus include enhancer	
mutatio	on, promoter mutation and premature termination of gene transcription.	43
1.10	Isolation of retroviral insertion sites by inverse PCR and splinkerette PCR.	47
1.11	Structure of the Sleeping Beauty transposon.	54
2.1	Workflow for identifying mouse candidate cancer genes from sequencing rea	ds
genera	ted in a retroviral insertional mutagenesis screen.	65
2.2	The number of sequence reads per tumour before mapping.	68
2.3	The lengths of retroviral reads that are unambiguously mapped, unmapped, and	nd
unmap	ped and uncharacterised.	74
2.4	BLAST scores for uncharacterised unmapped reads.	78
2.5	The filtering process for single mapping reads.	81
2.6	Determining the exact insertion site and orientation of retroviral and transpos	on
insertio	ons in the mouse genome.	83
2.7	Insertions in the mouse aminoadipate-semialdehyde synthase (Aass) gene are	
PCR a	rtefacts that map to an LTR-like sequence in the mouse genome	85
2.8	A high proportion of insertions in control samples map to the Myc gene.	86
2.9	The number of insertions per tumour and reads per insertion.	88
2.10	The number of genes with insertions in 100 bp intervals up to 20 kb upstream	n in
the sen	se and antisense orientation and downstream in the sense and antisense orienta	ation
with re	espect to the gene.	91
2.11	Insertions around known cancer genes Pim1, Kit, Gata1 and Blm.	94

2.12	Insertions in $En2$ and $Foxf2$ are located at the splice junctions used to constru-	uct
the T2	2/Onc transposon and are contaminating sequences.	103
3.1	MuLV and T2/Onc insertions across the mouse genome.	120
3.2	Known and putative tumour suppressor genes identified in the Sleeping Beau	ıty
screer	1.	122
3.3	The distribution of MuLV insertions within candidate cancer genes.	126
3.4	Intragenic MuLV insertions in candidate cancer genes.	128
3.5	Co-occurring MuLV and T2/Onc insertions help to identify the mechanism of	of
mutag	genesis of genes Notch1, Rasgrp1 and Etv6.	133
3.6	Variation in the distribution of MuLV and T2/Onc insertions in Myb, Fli1 and	d
<i>Erg</i> m	nay reflect differences in the mechanisms of mutagenesis.	137
3.7	Smg6 and Foxp1 are putative tumour suppressor genes identified by MuLV	
inserti	ional mutagenesis.	141
3.8	Knockdown of QSK in human HeLa cells is associated with increased	
chrom	nosome lagging at anaphase.	159
4.1	The distance between the genomic coordinates of adjacent SNPs on the 10K	and
10K 2	2.0 SNP arrays.	167
4.2	The number of SNPs per human protein-coding gene on the 10K and 10K 2.	0
SNP a	arrays.	167
4.3	Altering the values for parameters in DNAcopy leads to differences in the re	gions
of cop	by number change detected by the algorithm, as demonstrated for ovarian cance	er cell
line 4	1M-CISR.	169
4.4	Graphical output from MergeLevels for human cancer cell lines 786-0 and A	N3-
CA.		171
4.5	The number of human cancer cell lines with segments of varying log ₂ -ratio	
follow	ving processing with DNAcopy and DNAcopy plus MergeLevels.	172
4.6	The number of human cancer cell lines with segments of varying copy numb	er
ratio f	following processing with DNAcopy plus MergeLevels.	172
4.7	Distribution of the lengths of amplicons, deletions and homozygous deletion	s in
713 h	uman cancer cell lines.	174
4.8	Overview of the method for identifying over-representation of the human	
orthol	logues of mouse candidate cancer genes in regions of human copy number	
chang	e.	176
4.9	Over-representation of human orthologues of genes nearest to CISs in ampli	cons

with boundaries extended beyond the first and last amplified SNP by a maximum

distan	ce of 0 kb, 200 kb, 500 kb, 1 Mb, 3 Mb, 5 Mb and up to the adjacent, non-ampl	ified
SNPs.		178
4.10	Over-representation of human orthologues of genes nearest to CISs in full-ler	gth
humar	n amplicons and shuffled full-length amplicons.	181
4.11	Over-representation of human orthologues of candidate cancer genes in regio	ns of
copy r	number change.	182
4.12	Over-representation of known oncogenes and known tumour suppressor gene	s in
region	s of copy number change in human cancer cell lines.	183
4.13	Over-representation of human orthologues of genes nearest to CISs with a P-	
value	of <0.001 in regions of copy number change in human cancer cell lines derived	
from s	solid tumours and haematopoietic and lymphoid cancers.	183
4.14	Over-representation of human orthologues of candidate cancer genes in region	ns of
copy r	number change in cancer cell lines derived from the upper aerodigestive tract,	
autono	omic ganglia, breast, large intestine, oesophagus and stomach.	185
4.15	Over-representation of human orthologues of genes nearest to CISs and genes	5
furthe	r from CISs in amplicons and deletions, where CISs have a <i>P</i> -value of <0.001 a	nd
< 0.05		187
4.16	Insertions appear to be associated with the gene nearest to the CIS, i.e.	
16000	014C10Rik and Slamf6, even though adjacent genes are also amplified.	189
4.17	Known human oncogenes EVI1, FGFR2 and KIT are amplified in human	
cance	r cell lines and are disrupted by retroviral insertional mutagenesis in mouse	
lymph	iomas.	193
4.18	Candidate oncogenes MMP13, SLAMF6 and RREB1 are amplified in human	
cance	r cell lines and are disrupted by retroviral insertional mutagenesis in mouse	
lymph	iomas.	195
4.19	Insertions assigned to <i>Heatr5a</i> may be associated with <i>Hectd1</i> or <i>EG544864</i> .	198
4.20	Candidate tumour suppressor genes WWOX and ARFRP2 are deleted in huma	n
cance	r cell lines and are disrupted by retroviral insertional mutagenesis in mouse	
lymph	iomas.	206
4.21	Under- and over-representation of human orthologues of candidate cancer get	nes
in regi	ions of copy number variation.	216
5.1	Distance between adjacent copy number probes across the human genome.	226
5.2	Characterisation of amplicons and deletions in 598 human cancer cell lines	
analys	ed using high-resolution SNP array CGH.	228

X

List of Tables

2.1	Characterisation of the insertional mutagenesis datasets.	66
2.2	The number of MuLV reads mapped using SSAHA2, with varying values for	-
parame	eters seeds and skip, and BLASTN.	72
2.3	Repeat elements that are over-represented and under-represented among	
unmap	ped reads compared with unambiguously mapped reads.	77
2.4	Summary of the proportions of unmapped and unambiguously mapping reads	5
that co	ntain vector sequences, or sequences of low complexity, low quality or short	
length.		77
2.5	Number of intergenic insertions up to 20 kb upstream and downstream of known	own
oncoge	enes and tumour suppressor genes from the Cancer Gene Census.	93
2.6	Maximum window sizes in kb for significant CISs for varying numbers of	
inserti	ons in the retroviral and Sleeping Beauty screens.	98
2.7	Comparison of the methods used to generate candidate cancer genes lists from	n the
retrovi	ral and <i>Sleeping Beauty</i> screens.	98
3.1	The human orthologues of mouse CIS genes can help to identify the critical	
gene(s) in regions of copy number change in acute lymphoblastic leukaemias from	
Mullig	han <i>et al.</i> (2007).	115
3.2	Over-represented GO terms among CIS genes identified using MuLV.	116
3.3	The predicted mutation types and mechanisms of mutagenesis based on the	
distrib	ution of MuLV insertions within and around candidate cancer genes.	131
3.4	Gene expression values for candidate cancer genes in insertion-containing	
tumou	rs compared with tumours that do not contain insertions.	147
3.5	Genes containing an over-representation or under-representation of insertions	s on a
given t	tumour background compared with all other backgrounds and compared with v	vild-
type in	sertions only.	150
3.6	Gene pairs in which insertions co-occur more often or less often than expecte	d by
chance	S.	155
4.1	Description of the lists of mouse candidate cancer genes used for comparison	with
human	cancer copy number data.	163
4.2	Tissues of origin of human cancer cell lines used in the 10K SNP array CGH	
analys	is.	165

4.3	The number of amplicons in which known cancer genes among genes nearest	to
CISs at	re identified when the amplicon boundaries are altered.	179
4.4	Genes that are nearest to CISs in mouse lymphomas and are also promising	
candid	ates for targets of amplification in human cancer cell lines.	192
4.5	miRNA genes that are nearest to CISs in mouse lymphomas and are amplified	1
and/or	deleted in human cancer cell lines.	198
4.6	Mouse genes that contain retroviral insertions within the coding region and ar	e
also pr	omising candidates for targets of amplification or deletion in human cancer cel	1
lines.		201
4.7	Mouse genes that contain retroviral insertions within the transcribed or transla	ated
region	and are also promising candidates for targets of deletion in human cancer cell	
lines.		203
4.8	Comparison of methods for detecting regions of copy number gain in 50	
randon	nly selected cancer cell lines.	212
4.9	<i>P</i> -values for the co-occurrence between genes from each gene list within CNV	/s
and reg	gions of copy number change in human cancer cell lines.	217
5.1	Tissues of origin of human cancer cell lines used in high-resolution copy num	ber
analysi	s.	225
5.2	Number of copy number probes per human autosome.	226
5.3	Lists of CIS genes that are in recurrent amplicons across all cell lines and acro	DSS
haemat	topoietic and lymphoid cancer cell lines only.	235
5.4	A list of CIS genes for which the maximum copy number across all cell lines	is
signific	cantly higher than expected by chance.	239
5.5	A list of CIS genes that are in recurrent deletions of copy number less than or	
equal t	o 0.6 across all cell lines and across haematopoietic and lymphoid cancer cell	
lines.		246
5.6	A list of CIS genes that are in recurrent deletions of copy number 0.3 or less	
across	all cell lines and across haematopoietic and lymphoid cancer cell lines only.	251
5.7	Comparison of the high- and low-resolution datasets based on the proportion of	of
CIS ge	nes and known cancer genes that are amplified and deleted.	258
5.8	A list of CIS genes that are in recurrent amplicons, recurrent deletions of copy	/
numbe	r 0.6 or less and recurrent deletions of copy number 0.3 or less in the 10K CGH	ł
dataset		260
5.9	A list of amplified and deleted CIS genes that are over- or under-represented i	n
cell lin	es that contain a mutation in TP53 or CDKN2A.	265

xiii

5.10 A list of CIS genes that are co-amplified or co-deleted across a significant number of human cancer cell lines. 269

5.11 A list of genes that are co-amplified, co-deleted or amplified and deleted across human cancer cell lines and are also co-disrupted by MuLV in mouse lymphomas. 271

Appendices

А	Human Ensembl genes and their mouse orthologues for known cancer genes i	n
the Car	ncer Gene Census.	337
B1	Sleeping Beauty CISs and predicted CIS genes obtained using the kernel	
convol	ution-based framework with a kernel width of 30 kb.	342
B2	MuLV CISs and predicted CIS genes obtained using the kernel convolution-b	ased
framew	vork with a kernel width of 30 kb.	342
C1	List showing other cancer-associated datasets in which the MuLV CIS genes	
appear		348
C2	List showing other cancer-associated datasets in which the Sleeping Beauty C	IS
genes a	appear.	353
D	Nearest and further genes from CISs with a <i>P</i> -value of <0.001 or <0.05 in lists	S
supplie	ed by the Netherlands Cancer Institute.	354
Е	Human cancer cell lines used in the 10K and SNP 6.0 CGH analyses.	361

Abbreviations

ALL	acute lymphoblastic leukaemia
AML	acute myeloid leukaemia
API	application programming interface
BAC	bacterial artificial chromosome
CGH	comparative genomic hybridisation
ChIP	chromatin immunoprecipitation
CIS	common insertion site
CML	chronic myelogenous leukaemia
CNV	copy number variation
COSMIC	Catalogue of Somatic Mutations in Cancer
CRUK	Cancer Research UK
DAS	distributed annotation system
ES	embryonic stem
EST	expressed sequence tag
ESP	end-sequence profiling
HGNC HUGC) Gene Nomenclature Committee
HMM	hidden Markov model
IARC	International Agency for Research on Cancer
IR/DR	inverted repeat/direct repeat
KC	kernel convolution
LINE	long interspersed nuclear element
LOH	loss of heterozygosity
LTR	long terminal repeat
MC	Monte Carlo
MCC	Matthew's Correlation Coefficient
MCR	minimal common region of amplification or deletion
MGI	Mouse Genome Informatics
	mammary tumour virus
MuLV	murine leukaemia virus
NCBI	National Center for Biotechnology Information
NKI	Netherlands Cancer Institute
PCR	polymerase chain reaction
PET	paired-end ditag sequencing
RTCGD	Retroviral Tagged Cancer Gene Database
SB	Sleeping Beauty
SNP	single nucleotide polymorphism
SINE	short interspersed nuclear element
TFBS	transcription factor binding site
UTR	untranslated region
VISA	viral insertion site amplification
WGSA	whole-genome sampling assay
WHO	World Health Organization
WTSI	Wellcome Trust Sanger Institute

Chapter 1 Introduction

1.1 Outline of introduction

This introduction presents the foundations of the work described in this thesis. Section 1.2 focuses on the importance of studying the genetic basis of cancer, beginning with an overview of the burden of cancer. This is followed by a synopsis of the major contributions to the current understanding of how cancer develops, and a description of the main classes of genes and some of the genetic pathways known to be involved in cancer development. The section concludes with a discussion of the contribution of cancer genetics to the development of drugs for cancer treatment. Section 1.3 discusses the use of genome-wide approaches in the identification of cancer genes in humans. Prior work on the analysis of mutations, gene expression and epigenetics in cancer genomes is outlined, and research into the analysis of copy number changes is described in greater detail. Methods to identify transcription factor binding sites, and therefore to elucidate regulatory pathways, are also discussed. Section 1.4 describes the role of the mouse in cancer research and focuses on the use of retroviral and transposon-mediated mutagenesis in the genome-wide discovery of novel cancer genes and collaborations between genes involved in cancer. A significant portion of the work presented in this thesis relates to the comparison of human and mouse datasets for cancer gene identification, and previous studies of this kind are discussed in Section 1.5. Finally, the aims and rationale of this thesis are presented in Section 1.6.

1.2 An introduction to cancer

1.2.1 Definition and classification

Cancer is a class of diseases manifesting as uncontrolled cell division that leads to invasion of surrounding tissues and spread to distant sites (metastasis). These malignant properties of cancers differentiate them from benign tumours, in which abnormal cell proliferation is usually confined locally. Most cancers are classified according to the tissue of origin. There are over 100 distinct types, and 4 broad categories: carcinoma, arising in epithelial cells; sarcoma, arising in connective or supportive tissue and soft

tissue; leukaemia, arising in blood-forming tissues; and lymphoma, arising in cells of the immune system. See Pelengaris and Khan (2006).

1.2.2 Epidemiology

Cancer is a leading cause of death worldwide, accounting for 13% of all deaths in 2005 (WHO, 2008). In developed countries, it is the second greatest cause of death after cardiovascular disease, while in less developed countries, it is the third greatest after infectious and cardiovascular diseases. In 2002, 24% of all deaths in the UK were caused by cancer, compared with 12% in Asia and just 4% in Africa (CRUK, 2008; Ferlay *et al.*, 2004). Economic growth in Asia is expected to cause a rise in the proportion of deaths from cancer, and yet, due to its population size, more than half of all deaths from cancer already occur in Asia (Ferlay *et al.*, 2004). The global population is growing and ageing and, as cancer is predominantly a disease of older people (CRUK, 2008), the number of cancer deaths is expected to increase by 45% between 2007 and 2030 (WHO, 2008).

More than a quarter of a million new cases of cancer are diagnosed each year in the UK, and the four most common cancers - breast, lung, colorectal and prostate - account for half of these. In 2004, the most common cancers in men and women were breast and prostate, respectively. However, in both sexes, lung cancer was the biggest killer, accounting for 22% of all cancer deaths in 2005 (CRUK, 2008; Figure 1.1).

It is estimated that around 35% of all deaths from cancer are preventable, and 9 main modifiable risk factors have been identified (Danaei *et al.*, 2005). The leading risk factor is smoking, which is thought to contribute to 21% of all preventable cancers. Others include alcohol use, diet, and physical inactivity. Environmental risk factors account for much of the striking geographical variation in the incidence of certain cancers, and migration studies indicate that reducing exposure to these factors could eliminate a high proportion of deaths from cancer. There is, for example, a heightened risk of developing stomach cancer in Japan (Parkin *et al.*, 2005), where risk factors include infection by *Helicobacter Pylori* (IARC, 1994) and a diet rich in salted foods (Tsugane, 2005). However, within one generation of settling in Hawaii, the incidence of stomach cancer among Japanese immigrants declines to levels comparable with the surrounding population (Peto, 2001).

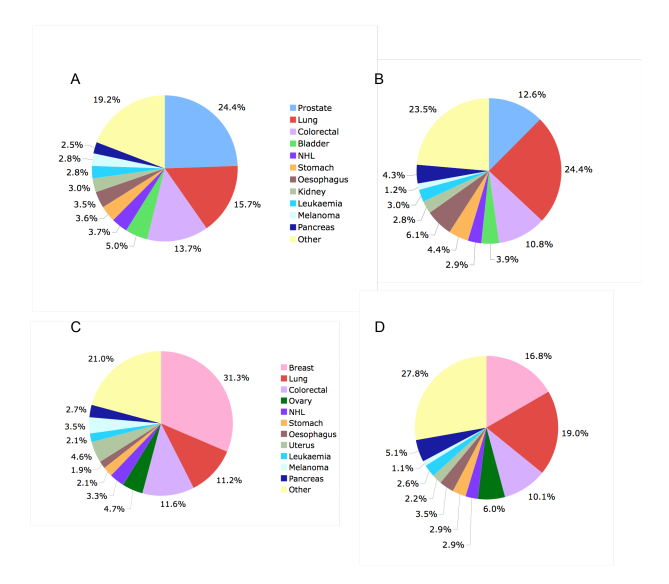


Figure 1.1. Summary of cancer incidence in 2004 and deaths from cancer in 2005 for the most common sites of cancer in males and females in the UK. Cancer incidence and mortality among males are shown in Figures A and B, respectively. Cancer incidence and mortality among females are shown in Figures C and D, respectively. The statistics for this figure were obtained from the Cancer Research UK CancerStats resource (CRUK, 2008).

While prevention could significantly reduce the burden of cancer, improvements in early diagnosis and treatment are also essential. Screening procedures that have reduced cancer mortality rates include the identification and removal of polyps in the colon (Weir *et al.*, 2003) and pre-cancerous cells in the uterus (Misra *et al.*, 1998), and widespread mammography screening for breast cancer (Shapiro, 1997). However, effective screening has been developed for only a handful of cancers, and advances in cancer treatment have been slower than for other chronic diseases, such as cardiovascular disease (Danaei *et al.*, 2005). A greater understanding of the genetic basis of cancers is essential for the development of effective treatments and diagnostic techniques.

1.2.3 The multi-stage theory of carcinogenesis

1.2.3.1 The somatic mutation theory

The theory that cancer is caused by somatic mutation can be traced back to Boveri (1926, 1914), who, extending the views of Hansemann (1890) and through his own work on aneuploidy in cancer cells, postulated that tumours originate from a single cell that has acquired chromosomal abnormalities. 35 years later, the multistage theory of carcinogenesis was borne, first postulated as two-stage carcinogenesis, in which an initiator and a promoter agent were proposed to be required for malignancy (Berenblum and Shubik, 1949), and later in the Armitage-Doll model, which suggested that six or seven independent, sequential, events were required (Armitage and Doll, 1954). Nowell (1976) proposed a model of clonal evolution, in which tumours evolve from a single cell through a series of stepwise genetic alterations within the original clone. He postulated that as the tumour progresses, genetically variant sublines emerge and the most favourable sublines, i.e. those with the greatest growth advantage, are selected (Figure 1.2).

An alternative theory for carcinogenesis, the tissue organisation field theory, proposes that rather than a cell acquiring the ability to proliferate uncontrollably through mutation, proliferation is in fact the default state of cells and cancer is caused by disruption to interactions between cells and tissues (Soto and Sonnenschein, 2004). There is, however, overwhelming support in favour of the somatic mutation theory for most cancer types.

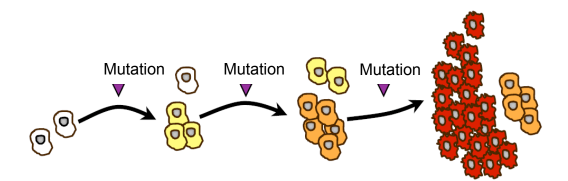


Figure 1.2. The clonal evolution of cancer. Tumours evolve from a single cell through a series of stepwise genetic changes within the original clone. Cells containing mutations that confer the greatest growth advantage are selected and become the dominant clone. Adapted from a figure supplied by D.J. Adams.

Chapter 1

1.2.3.2 The cancer stem cell hypothesis

In the original model of clonal evolution, events in all tumour cells can participate in the evolution of the tumour. However, the cancer stem cell hypothesis proposes that only cells that are capable of self-renewal, i.e. stem cells, contribute to tumour evolution and that these give rise to most of the cells with a more differentiated phenotype (for review, see Shipitsin and Polyak, 2008). The theory has some inconsistencies, but it is clear that putative cancer stem cells exist in most, if not all, cancer types, and xenotransplant assays have shown that stem cell-like tumour cells have a significantly higher potential to form tumours in irradiated NOD-SCID mice than do other cells from the same human tumour (Shipitsin and Polyak, 2008). Compared with well-differentiated tumours, poorly differentiated tumours overexpress genes that are normally enriched in embryonic stem (ES) cells (Ben-Porath et al., 2008). These genes include the transcriptional targets of NANOG, OCT4 and SOX2, which are key regulators of pluripotency and self-renewal in ES cells (see Loh et al., 2006). Wong et al. (2008) constructed a "module map" of stem cell genes, and showed that a subset of adult tissue stem cells shares a core gene expression program with ES cells, and that the ES cell-like program is frequently activated in human epithelial cancers. Other recent research has shown that the epithelialmesenchymal transition, which is often activated in tumour metastasis, is linked to the acquisition of epithelial stem cell-like properties (Mani et al., 2008).

1.2.3.3 Rate-limiting events in tumourigenesis

While it is widely accepted that cancer is caused by stepwise mutations, there are conflicting theories about how these mutations arise. The Armitage-Doll model suggests that mutations arise gradually over time, and that the number of rate-limiting events required for carcinogenesis can be inferred from the age-specific incidence of cancer and the rate of successive mutations in cells (Armitage and Doll, 1954). Cancers will not fit the model if the mutation rate is not constant, e.g. in smokers, where the mutation rate increases at the onset of smoking, or if the incidence does not increase with age, e.g. in childhood cancers (for review, see Knudson, 2001). However, the estimate of 5 to 7 mutations in colorectal cancer is compatible with the genetic model for colorectal tumourigenesis, in which at least four or five genes were proposed to be required for malignancy (Ashley, 1969; Fearon and Vogelstein, 1990).

More recent research suggests that a single rate-limiting step may be required for epithelial carcinogenesis, and that telomere crisis is one of the processes responsible for this step (Frieboes and Brody, 2005). The telomere crisis hypothesis proposes that mutations occur suddenly in cells with telomere dysfunction (Chin *et al.*, 2004; Maser and DePinho, 2002). In cells without active telomerase, telomeres erode and eventually cease to function. At this point, cells show massive genomic instability, including end-to-end fusions, non-reciprocal translocations, amplifications and deletions (Artandi *et al.*, 2000; O'Hagan *et al.*, 2002). This results in rapid cell senescence but some cells may escape by reactivating telomerase, and further mutations accumulate, leading to tumour progression (Maser and DePinho, 2002). Genomic instability is discussed in further detail in Sections 1.2.5.1.3 and 1.3.

1.2.4 The hallmarks of cancer

Hanahan and Weinberg (2000) proposed that all genetic alterations in cancer can be represented by six essential changes in cell physiology. These are "self-sufficiency in growth signals, insensitivity to antigrowth signals, evading apoptosis, limitless replicative potential, sustained angiogenesis, and tissue invasion and metastasis". The authors suggest that all tumours must acquire the same six capabilities, but that different genes may be mutated, and in a different order, even within cancers of the same type. The review by Hanahan and Weinberg is considered a seminal work, and the six "hallmarks of cancer" appear to be shared by most, if not all, malignancies.

1.2.5 Cancer genes

1.2.5.1 Classification

The term "cancer gene" will be used throughout this thesis to describe a gene for which mutations have been causally implicated in cancer. Cancer genes are often divided into 3 classes known as oncogenes, tumour suppressor genes and caretaker genes.

1.2.5.1.1 Oncogenes

In general, oncogenes play a role in accelerating cell growth and proliferation, but they may also contribute to loss of differentiation, avoidance of apoptosis, cell motility and

7

invasion (see Pelengaris and Khan, 2006). The normal counterparts of oncogenes, known as proto-oncogenes, mainly encode growth factors, growth factor receptors, signal transducers, transcription factors and regulators of cell death. Proto-oncogenes may become oncogenes through increased protein activity resulting from intragenic mutations that affect critical residues; increased protein concentration resulting from gene amplification, misregulation of gene expression or an increase in protein stability; or chromosomal translocations that increase inappropriate gene expression or produce a constitutively active fusion protein (Vogelstein and Kinzler, 2004). Oncogenes are dominant at the cellular level. One of the best known oncogenes is *MYC*, which appears to be activated in most human cancers at some stage during their development (see Pelengaris and Khan, 2006).

1.2.5.1.2 Tumour suppressor genes

In contrast to oncogenes, tumour suppressor genes act to limit the growth of tumours and inactivating mutations in these genes can lead to tumour development. Tumour suppressors inhibit cell proliferation by inducing growth arrest or apoptosis in response to DNA damage or hyperproliferative signals induced by oncogenes (see Pelengaris and Khan, 2006). They may be inactivated by missense mutations that alter sites required for protein activity; nonsense mutations that result in an inactive truncated protein; intragenic deletions and insertions; or epigenetic silencing (Vogelstein and Kinzler, 2004). Most tumour suppressor genes follow Knudson's "two-hit hypothesis", which proposes that both copies of the gene must be inactivated to confer a selective growth advantage on the cell (Knudson, 1971). Knudson applied his hypothesis to the identification of the first tumour suppressor gene, RB1. Compared with sporadic retinoblastoma, the hereditary form of this rare eye cancer arises earlier and is more often bilateral because cells already harbour one germline RB1 mutation and require only one additional somatic "hit" (Knudson, 1971). Some tumour suppressor genes are haploinsufficient, i.e. the loss of only one allele is required to confer a growth advantage. Haploinsufficiency of PTEN is sufficient for prostate cancer development, but progression is faster when both copies are inactivated (Trotman et al., 2003).

1.2.5.1.3 Caretaker genes

Caretakers maintain DNA integrity and their inactivation results in an increased tendency

8

to acquire mutations in other genes, including oncogenes and tumour suppressor genes (Vogelstein and Kinzler, 2004). Mutations in genes involved in repairing subtle mistakes during replication can cause microsatellite instability, which manifests as alterations in the length of short (1-4 bp) repetitive sequences called microsatellites (Loeb et al., 2003). Cells with microsatellite instability are particularly prone to mutation in the TGFBR2 tumour suppressor gene, and this is a common mechanism of disease in hereditary nonpolyposis colorectal cancer (HNPCC), in which patients have a germline mutation and a second, somatic, mutation in a mismatch repair gene, most often MSH2 or MLH1 (reviewed in Knudson, 2001). Much more common than microsatellite instability is chromosomal instability, which is caused by mutations in genes that are involved in largescale processes such as recombination and double-strand repair (Lengauer et al., 1998; Loeb et al., 2003). Chromosomal instability is characterised by gross chromosomal alterations, such as duplication or deletion of entire chromosomes (aneuploidy) or parts of chromosomes, and chromosomal rearrangement. Microsatellite and chromosomal instability are collectively known as genomic instability.

1.2.5.1.4 Genes with dual roles in cancer

The terms oncogene and tumour suppressor gene will be used to characterise genes described in this thesis. However, it should be noted that these terms are somewhat simplistic as the role of a protein may be dependent on the cellular context. Some mitogenic proteins have an intrinsic tumour suppressor activity such that inappropriate activation of the protein results in apoptosis of the mutated cell (Cobleigh et al., 1999). Activation of *Myc* in the pancreatic β cells of transgenic mice induces β cell proliferation but also induces apoptosis, which rapidly overwhelms the cell mass (Pelengaris et al., 2002). Likewise, the NOTCH1 receptor plays both oncogenic and tumour suppressive roles that reflect the pleiotropic effects of NOTCH1 signalling in different tissues (for review, see Radtke and Raj, 2003). NOTCH1 signalling is essential for maintaining haematopoietic stem cells and for committing haematopoietic progenitors to the T-cell lineage (Radtke et al., 1999). Aberrant NOTCH1 expression contributes to over 50% of cases of human T-cell acute lymphoblastic leukaemia (Weng et al., 2004). The involvement of NOTCH1 was established through the discovery of a translocation between chromosomes 7 and 9 that brings the dominant active cytoplasmic domain under the control of the $TCR\beta$ locus (Ellisen *et al.*, 1991), but point mutations and deletions are also implicated (Weng et al., 2004). In mice, Notch1 induces lymphomas by suppressing

p53 (Beverly *et al.*, 2005). However, Notch1 also functions as a tumour suppressor in mouse skin, where it participates in terminal differentiation by inducing *Waf1* and repressing Shh and Wnt signalling (Radtke and Raj, 2003).

1.2.5.2 Cancer Gene Census

In 2004, a census of genes in which mutations have been causally implicated in human cancer was compiled from the literature (Futreal *et al.*, 2004). It lists genes that are mutated by insertions, deletions or base substitutions in the coding region or splice sites, or by chromosomal translocations or copy number changes. Stringent criteria were applied to exclude genes in which reported mutations could be "passenger" mutations that do not confer any growth advantage.

The census indicates that mutations in more than 1% of human genes are implicated in cancer. Of the 291 genes listed in the original census, 90% have somatic mutations in cancer, 20% have germline mutations, and 10% have both. Chromosomal translocations are the most common class of somatic mutation in human cancer and almost all are dominant at the cellular level. Excluding translocations, there are equal numbers of recessive and dominant somatic mutations within the census list. The protein kinase domain is the most common domain encoded by genes in the census. Domains in proteins involved in transcriptional regulation and DNA maintenance and repair are also over-represented. See Futreal *et al.* (2004).

The Cancer Gene Census is frequently updated and the working list can be downloaded from http://www.sanger.ac.uk/genetics/CGP/Census/. It represents a valuable source of "known" cancer genes that will be utilised in this thesis.

1.2.6 Pathways in cancer

It is often more sensible to focus on the pathways that have been disrupted in cancer, rather than on individual genes. The p53 and RB1 pathways are thought to be inactivated in most, if not all, cancers. However, while *TP53*, which encodes p53, and *RB1* are often mutated, the same effect can be achieved by mutating a different gene in the pathway (see Vogelstein and Kinzler, 2004 and Figure 1.3).

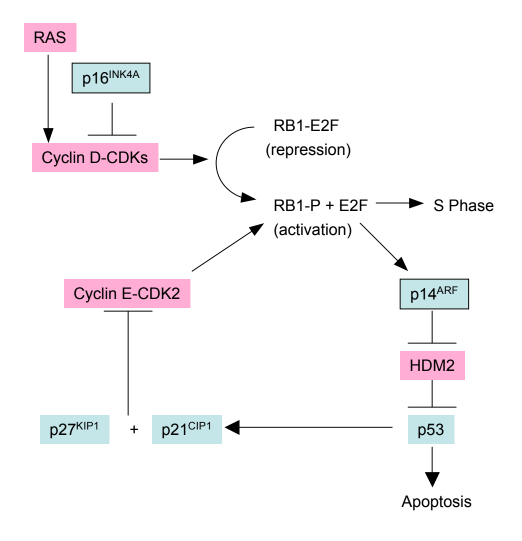


Figure 1.3. Mutations in different genes in the same pathway can have an equivalent effect. The figure shows a simple representation of the p53 and RB1 genetic pathways. The pathways are coupled through the *INK4A/ARF* locus, which encodes $p16^{INK4A}$ and $p14^{ARF}$, shown here in black boxes, and through $p21^{CIP1}$, which is activated by p53 and inhibits Cyclin E-CDK2 complexes in the RB1 pathway. Genes that are frequently inactivated in cancer are shown in blue; genes that are frequently activated in cancer are shown in pink. Adapted from Figure 1 in Lowe & Sherr (2003).

p53 is a transcription factor that inhibits cell growth and induces apoptosis in response to cellular stress, such as DNA damage or hyperproliferative signals induced by oncogenes (for review, see Vogelstein *et al.*, 2000). Many p53-responsive genes are involved in arresting cell proliferation at the G1/S and G2/M cell cycle transitions so that cells with DNA damage can be repaired before proceeding to DNA replication or mitosis (Vogelstein *et al.*, 2000). p53 is inhibited by the binding of HDM2 (known as Mdm2 in the mouse) to its N-terminal transactivation domain (Momand *et al.*, 2000). HDM2 also acts as an E3 ubiquitin ligase that targets itself and p53 for degradation by the ubiquitin-dependent proteasome pathway (Momand *et al.*, 2000). Overexpression of *HDM2* may have an equivalent effect to underexpression of *TP53*, and amplification of *HDM2* has been observed in a variety of tumours, including breast, lung and gastric cancers (Gunther *et al.*, 2000; Marchetti *et al.*, 1995a; Marchetti *et al.*, 1995b).

The RB1 pathway regulates cell proliferation by repressing the transcription of genes required for progression through the G1 phase of the cell cycle and for entry into S phase (Figure 1.3 and for review, see Weinberg, 1990). In mid-G1 phase, mitogenic signals from the RAS/MAP kinase pathway activate transcription of D-type cyclins, which bind to the cyclin-dependent kinases CDK4 and CDK6 and initiate phosphorylation of RB1. This results in the release of E2F transcription factors and their subunit partners, DP, from complexes with RB1, and the E2Fs activate transcription of genes required for cell cycle progression. Cyclin E-CDK2 complexes complete the phosphorylation of RB1. A further level of regulation is provided by cyclin-dependent kinase inhibitory (CDKI) proteins, which consist of the INK4 and CIP/KIP protein families. The INK4 proteins (p16^{INK4A}, p15^{INK4B}, p18^{INK4C} and p19^{INK4D}) inhibit CDKs, whereas the CIP/KIP proteins (p27^{KIP1} and p21^{CIP1}) stimulate assembly of the cyclin D-CDK4-6 complexes and inhibit cyclin E-CDK2 (for review, see Sherr, 2001). Inactivating p16^{INK4A}, p18^{INK4c}, p21^{CIP1} or p27^{KIP1} has a similar effect to inactivating RB1 (Sherr, 2001; Vogelstein and Kinzler, 2004). $p16^{INK4A}$ is inactivated by homozygous deletion, promoter methylation or, to a lesser extent, point mutation, in a large number of tumours (for review, see Liggett and Sidransky, 1998). Likewise, activation of CDK4 and cyclin D1 has an equivalent effect on the RB1 pathway, and these oncogenes are frequently amplified and overexpressed in cancer (Vogelstein and Kinzler, 2004).

Cancer pathways are not standalone entities. As well as regulating the RB1 pathway, $p21^{CIP1}$ is one of the major transcriptional targets of p53 (Vogelstein *et al.*, 2000). In

addition, the p53 and RB1 pathways are coupled through the *INK4A/ARF* (or *CDKN2A*) locus, which uses alternative reading frames to encode two tumour suppressors: p16^{INK4A}, described above, and p14^{ARF} (also known as p19^{ARF} or ARF), which activates p53 by sequestering HDM2 (Quelle *et al.*, 1995; Sherr, 2001; Figure 1.3). The *INK4A/ARF* locus is frequently mutated in human cancer but mutations in *TP53* and *INK4A/ARF* are often mutually exclusive, e.g. in human glioblastoma (Fulci *et al.*, 2000). This suggests that inactivating both loci may not provide any additional growth advantage. However, expression and genotypic analysis of *Trp53*, *Arf* and *Mdm2* in Myc-induced murine lymphomas showed that *Mdm2* was overexpressed in a significant proportion of *Arf*-deficient tumours, while loss of both *Arf* and *Trp53* in primary pre-B cells results in a greater growth advantage than the loss of one gene alone (Eischen *et al.*, 1999).

1.2.7 Treatment of cancer

The main forms of cancer treatment, often used in combination, are surgery, radiotherapy and chemotherapy. Some cancers respond well to these treatments, e.g. testicular cancer has a high cure rate following chemotherapy, but others, such as lung cancer, show a much lower response (CRUK, 2008). Radiotherapy and chemotherapy can have considerable side effects as neither specifically targets cancer cells.

A greater understanding of the genetic basis of cancer has initiated the development of more effective therapies that specifically target deregulated gene expression and signalling pathways in cancer cells. Gleevec (imatinib) targets the BCR-ABL oncoprotein, which causes 95% of cases of chronic myelogenous leukaemia (CML) and $\sim 20\%$ of cases of acute lymphoblastic leukaemia (ALL) (Deininger and Druker, 2003; Faderl et al., 1999). Gleevec stabilises a catalytically inactive form of BCR-ABL (Nagar et al., 2002). It also inhibits four other tyrosine kinases (KIT, PDGFRA, PDGFRB and ARG) but shows minimal side effects (Buchdunger et al., 1996; Druker et al., 1996; Okuda et al., 2001). Treatment has an 89% response rate in chronic CML after 5 years (Druker et al., 2006), and an initial, but not durable, response rate of 52% in patients who have progressed to blast crisis, the terminal phase of the disease (Sawyers et al., 2002). Gleevec has also proved effective in the treatment of gastrointestinal stromal tumours (GISTs) by targeting KIT (Joensuu et al., 2001; van Oosterom et al., 2001) and PDGFRA (Apperley et al., 2002). Other tyrosine kinase inhibitors include Herceptin (trastuzumab), which targets the HER2/ERBB2 receptor in breast cancer (Cobleigh et al., 1999), and

Iressa (gefitinib), which targets the epidermal growth factor receptor (EGFR) in lung adenocarcinomas and non-small cell lung cancers (Fukuoka *et al.*, 2003).

As with traditional therapies, there is evidence that cancer cells can develop resistance to targeted therapies (Balak et al., 2006; Engelman et al., 2007; Gorre et al., 2001; Kobayashi et al., 2005; Nagata et al., 2004; Shattuck et al., 2008), necessitating the development of new drugs for targeted combination therapy (Baselga, 2006). However, the results outlined above demonstrate that targeting a single, critical gene in a complex tumour can elicit a dramatic response. Success of such a treatment depends on the targeted kinase being required for growth and survival of the tumour throughout its evolution (a notion known as "oncogene addiction" (Weinstein, 2002)). The mutation status of other genes can also influence drug response. For example, breast tumours that harbour an amplification of *HER2/ERBB2* are less responsive to trastuzumab if they also harbour an oncogenic *PIK3CA* mutation or have low *PTEN* expression (Berns *et al.*, 2007). Likewise, lung cancers that contain KRAS mutations are resistant to treatment with EGFR inhibitors because KRAS acts further downstream in the EGFR pathway (Pao et al., 2005). Due to huge variation in the genetic basis of different cancers, each targeted therapy will be effective against only a subset of cancers. This necessitates the identification of many different drug targets, and fundamentally relies on the identification and characterisation of mutated genes in cancer.

1.3 Genome-wide approaches for human cancer gene discovery

The elucidation of the human genome sequence and developments in high-throughput techniques for genome-wide analysis have allowed for profiling of entire cancer genomes. This section discusses the large-scale technologies that are available for detecting alterations and, ultimately, for identifying cancer genes in human cancer genomes.

1.3.1 Gene resequencing

Advances in DNA sequencing technology have enabled the identification of recurrent intragenic mutations across multiple cancer genomes. Davies and colleagues (2002) screened the coding sequence and intron-exon junctions of BRAF for mutations in more than 900 human cancer cell lines and primary tumours, and found somatic missense mutations in 66% of malignant melanomas and in a smaller proportion of many other

human cancers. 80% of *BRAF*-mutated melanomas were found to contain a V599E substitution, which is thought to constitutively activate the kinase by mimicking phosphorylation (Davies *et al.*, 2002). An inhibitor has recently been developed that selectively targets the V599E gene product, and so selectively targets BRAF in tumour cells (Tsai *et al.*, 2008).

As the cost of sequencing has diminished, it has become possible to perform larger scale screens to look for mutations in multiple genes across multiple tumours. The first systematic mutational study of a complete gene family was performed by Bardelli and coworkers (2003), who identified 7 candidate cancer genes in a screen of the tyrosine kinase gene family in 182 colorectal cancers. A further study of mutations in the tyrosine phosphatase gene family identified 6 putative tumour suppressor genes that were mutated in 26% of the colorectal cancers analysed (Wang et al., 2004). Resequencing of the phosphatidylinositol 3-kinase (PI3K) gene family revealed one member, PIK3CA, that is frequently mutated in tumours of the colon, breast, brain and lung, with most mutations clustering within the helical or catalytic domain (Samuels and Velculescu, 2004). Mutations have since been identified in additional tumour types, such as hepatocellular carcinomas (Bachman et al., 2004) and ovarian cancers (Campbell et al., 2004; Levine et al., 2005). A screen of serine/threonine kinases showed that 40% of colorectal tumours harbour a mutation in 1 of 8 PI3K-pathway genes (Parsons et al., 2005). The PI3K pathway regulates a wide range of cellular functions that are important in cancer, including growth, proliferation, survival, angiogenesis and migration (Brugge et al., 2007).

Studies at the Wellcome Trust Sanger Institute have centred around the resequencing of coding regions from all 518 genes of the protein kinase family. A study of 25 breast cancers revealed diverse patterns of mutation, with variation in the number of mutations and in the identity of mutated genes, such that no commonly point-mutated kinase gene was identified (Stephens *et al.*, 2005). A study of 33 lung cancers reached similar conclusions (Davies *et al.*, 2005). While both studies showed an over-representation of nonsynonymous substitutions, as predicted for "driver" mutations that confer a selective growth advantage on the cell, most of the mutations are likely to be "passenger" mutations that do not contribute to tumourigenesis. Protein kinase resequencing at the Sanger Institute has culminated in the identification of 921 base substitution somatic mutations in 210 diverse human cancers (Greenman *et al.*, 2007). Putative driver

mutations were identified in 119 genes but 83% of mutations were predicted to be passengers. Cancers showed variation in mutation prevalence, with many of the cancer types with highest prevalence originating from high turnover, surface epithelia that are most exposed to mutagens (Greenman *et al.*, 2007). Cancers also showed different "mutational signatures", which often reflect differences in mutagenic exposure. For example, most lung cancers have a high proportion of C:G > A:T transversions, which are caused by exposure to tobacco carcinogens (Davies *et al.*, 2005).

The first study to approach the scale of a genome-wide screen involved resequencing the coding regions of all (~13,000) consensus coding sequence (CCDS) genes in 11 breast and 11 colorectal cancers (Sjoblom et al., 2006). Each cancer was found to harbour an average of 93 mutated genes, of which at least 11 (189 candidates in total) were thought to be driver mutations. Many of the functional groups and pathways enriched for candidate cancer genes were unique to one or other cancer type, suggesting differences in the tumourigenic process in breast and colorectal cancers (Lin et al., 2007). There have been claims that the statistical analysis performed in this screen was flawed, in part because they used a different dataset to estimate background mutation rates, which can vary between and within cancer genomes, and because the sample size was small (Getz et al., 2007). However, the findings of this study are in agreement with those of Greenman et al. (2007) in suggesting that the genomic landscape of human cancers is more complex than previously thought (Kaiser, 2006). The study has since been expanded to include all of the human RefSeq (Pruitt et al., 2007) genes and a larger number of breast and colorectal cancers (Wood et al., 2007). Each tumour contained an average of 15 potential driver mutations and most of these were in genes that were mutated in fewer than 5% of tumours, therefore recapitulating the conclusions of the previous studies.

Although statistical methods can provide a prediction of the likely driver and passenger mutations within a cancer, there is a strong rationale for using functional assays to test the predictions. Frohling and coworkers (2007) resequenced the coding exons and splice junctions of the receptor tyrosine kinase *FLT3* in samples from patients with acute myeloid leukaemia (AML). They found that out of 9 mutants with candidate driver mutations, only 4 were able to transform cells in culture (for review, see Futreal, 2007).

The Wellcome Trust Sanger Institute Catalogue of Somatic Mutations in Cancer (COSMIC) collates and displays somatic mutation information relating to human cancers

(Forbes *et al.*, 2006). At the time of writing (May 2008, COSMIC release 37), the database contained mutation data for around 4,770 genes from ~260,000 tumours. Gene resequencing is also a major component of the \$50 million 3-year pilot phase of the Cancer Genome Atlas (http://cancergenome.nih.gov/), a large-scale collaboration between the National Cancer Institute (NCI) and the National Human Genome Research Institute (NHGRI).

1.3.2 Gene expression profiling

Gene expression arrays can be used to analyse the transcription of thousands of genes simultaneously. There are two main types: cDNA arrays, where clones corresponding to the transcripts to be analysed are spotted onto a matrix, and oligonucleotide arrays, where oligonucleotides corresponding to the transcripts are synthesised onto a matrix along with mismatch control oligonucleotides. A new approach has also been developed, in which of is the abundance transcripts measured directly using Illumina (http://www.illumina.com) sequencing technology. In two-colour microarray expression analysis, the sample of interest and a control sample are differentially labelled with fluorescent dyes and are hybridised onto the array, which is then scanned to determine the ratio of fluorescence intensities for each gene. The ratio represents the relative amounts of transcript in the sample. Unsupervised clustering of the expression data for multiple samples can be used to subcategorise cancers. For example, lung cancers cluster into known histological subtypes that are predictive of patient survival (Beer et al., 2002; Bhattacharjee et al., 2001; Garber et al., 2001). Gene expression profiles may also provide an indication of the genes involved in oncogenesis in a given tumour. Lung cancers harbouring a mutation in KRAS have a characteristic expression profile that can be used in their identification (Sweet-Cordero et al., 2005). Analysis of gene expression does not provide any insights into the underlying genetic changes and it can be affected by physiological variation, such as the degree of inflammatory response or hypoxia (Eden et al., 2004). However, it is important as a complementary approach to other methods of cancer profiling, such as mutational and copy number analysis. Integrative approaches involving gene expression and copy number analysis are discussed in the following section.

Chapter 1

1.3.3 Copy number analysis

1.3.3.1 DNA copy number changes

Changes in DNA copy number result from chromosomal aberrations such as deletions and duplications, non-reciprocal translocations and gene amplifications. Copy number variations (CNVs) have been identified in all humans studied (Feuk *et al.*, 2006), and a genome-wide study of 270 apparently healthy individuals from four diverse populations identified almost 1,500 germline copy number variable regions encompassing 12% of the human genome (Redon *et al.*, 2006). CNVs accounted for ~18% of the total detected variation in gene expression between individuals, suggesting that they make a considerable contribution to phenotypic variation (Stranger *et al.*, 2007). In the context of cancer, genomic instability results in the acquisition of somatic copy number aberrations that may contribute to tumourigenesis through the amplification of oncogenes and/or loss of tumour suppressor genes. Genomic instability is also referred to in Sections 1.2.3.3 and 1.2.5.1.3.

Chromosome instability, which manifests as alterations in chromosome number (aneuploidy), seems to arise early in tumourigenesis but increases with tumour progression (for review, see Lengauer *et al.*, 1998). Fridlyand and coworkers (2006) found that shorter or altered telomeres were associated with greater numbers of amplifications but that the frequency of low-level changes was associated with altered expression of genes involved in mitosis, cell cycle, DNA replication and repair, and included many genes that are direct targets of E2F (Fridlyand *et al.*, 2006). This suggests that the RB1 pathway (see Section 1.2.6) contributes to chromosome instability, as hypothesised by Hernando *et al.* (2004) (Fridlyand *et al.*, 2006). Advanced tumours tend to reach a stable state, which, in the form of cancer cell lines, are stable over many generations and in different laboratories, suggesting that they have evolved to an optimal state (Albertson *et al.*, 2003).

1.3.3.2 Using CGH to detect copy number changes

Large alterations in copy number were initially detected and quantified using metaphase spreads in a technique known as comparative genomic hybridisation (CGH) (Kallioniemi *et al.*, 1992). In CGH, cancer and normal genomic DNA are differentially labelled with fluorochromes and are co-hybridised to normal metaphase chromosomes. Cot-1 DNA is

added to suppress hybridisation to repetitive elements in the genome. The ratio of fluorescence intensities at any chromosomal position is approximately proportional to the ratio of copy numbers of the cancer and normal DNA at that position (reviewed in Pinkel *et al.*, 1998). CGH profiles can be viewed and compared using the NCBI Cancer Chromosomes database, which integrates three databases of chromosomal aberrations in cancer: the SKY/M-FISH & CGH Database, the Mitelman Database of Chromosome Aberrations in Cancer, and the Recurrent Chromosome Aberrations in Cancer database (Knutsen *et al.*, 2005). Rearrangement breakpoints are linked to the underlying genome assembly. However, the tool is limited to cytogenetic resolution because CGH cannot detect changes of less than 20 Mb or distinguish changes that are close together, and it cannot determine exact genomic coordinates (Pinkel *et al.*, 1998).

Array CGH is a higher resolution, high-throughput version of conventional CGH, in which differentially labelled cancer and reference samples are hybridised to an array made from large genomic clones, e.g. bacterial artificial chromosomes (BACs), or cDNAs (for review, see Albertson and Pinkel, 2003; Pinkel *et al.*, 1998; Pollack *et al.*, 1999). The copy number is measured at each probe on the array, and can be mapped directly to the genome. A disadvantage of array CGH is that it cannot detect loss of heterozygosity (LOH), which has traditionally been identified using methods involving microsatellites and restriction fragment length polymorphisms (RFLPs) that are not suitable for large scale analyses (see Thomas *et al.*, 2006).

Single nucleotide polymorphism (SNP) arrays are the most recent development in copy number analysis. SNPs account for most of the genetic variation in the human genome (Stranger *et al.*, 2007) and they occur, on average, every 100-300 base pairs along the genome. The Affymetrix GeneChip Mapping Assay (http://www.affymetrix.com) is a commonly used procedure that combines a whole-genome sampling assay (WGSA) with high-density SNP arrays (Kennedy *et al.*, 2003; Matsuzaki *et al.*, 2004). WGSA is used to reduce the complexity of the sample, and involves ligating an adapter to restriction-digested DNA, which enables PCR amplification using a single primer that is complementary to the adapter (Figure 1.4B). The amplified DNA is then fragmented, labelled and hybridised to the array. SNPs within the amplified DNA are used as probes on the array, therefore ensuring that all probes are informative (Bignell *et al.*, 2004). In the Affymetrix GeneChip Mapping 10K assay, which uses an array containing 11,555 SNPs, WGSA involves a single restriction enzyme, *XbaI* (Kennedy *et al.*, 2003).

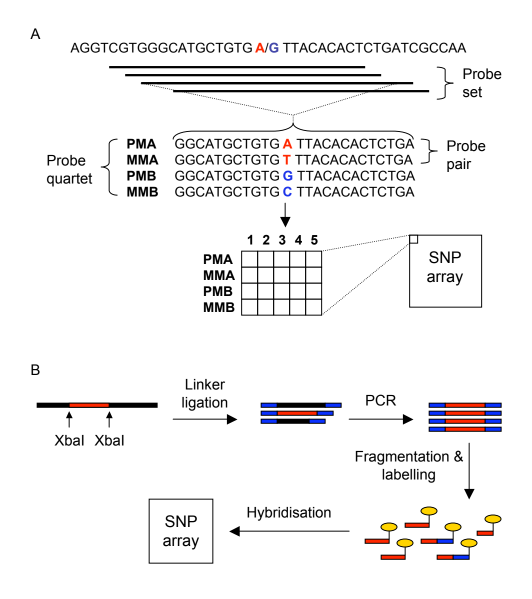


Figure 1.4. Array design (A) and whole-genome sampling assay (B) for the Affymetrix SNP array. A. A SNP in the DNA sequence is shown in red/blue. The SNP is represented in the array by a probe set, which comprises multiple probe quartets that differ from one another in the position of the polymorphic site relative to the centre of the probe. Each probe quartet consists of four 25mer oligonucleotides in the form of two probe pairs, which comprise a perfect match (PM) probe and a mismatch (MM) probe corresponding to each SNP allele (A and B). B. Genomic DNA is digested with a restriction enzyme, shown here as *XbaI*, and a linker (shown in blue) is ligated to the digested DNA. The DNA is PCR amplified using a primer that binds to the linker. Amplified DNA is fragmented, labelled and hybridised to the array.

Regions of the genome in which the *XbaI* site is rare will be under-represented in the array (Bignell *et al.*, 2004). The higher resolution 100K SNP array therefore use two restriction enzymes, *XbaI* and *HindIII*, which produce complementary SNP densities (Matsuzaki *et al.*, 2004). Each SNP in an Affymetrix array is represented by a "probe set" comprising multiple "probe quartets". Each probe quartet consists of four 25mer oligonucleotides in the form of two "probe pairs" comprising a perfect match probe and a mismatch probe corresponding to each SNP allele (Figure 1.4A). Probe quartets differ from one another in offset, i.e. the position of the polymorphic site relative to the centre of the oligonucleotide, and orientation (reviewed in Xiao *et al.*, 2007). Normal and tumour DNA are hybridised to different arrays, therefore avoiding the need for matched samples and allowing for a pool of normal samples to be used as a control (Bignell *et al.*, 2004; Figure 1.4C). As in other forms of array CGH, the copy number at each probe can be inferred from the intensity of fluorescence of hybridised sample DNA (Bignell *et al.*, 2004; Zhao *et al.*, 2004).

Commercially available arrays now range in resolution from 10,000 to ~1 million SNPs across the genome. SNP arrays therefore provide the potential for fine mapping of copy number changes, enabling the identification of small aberrations and accurate mapping of chromosomal breakpoints. Furthermore, the SNPs can be genotyped and compared to a normal sample to identify regions of LOH. This permits the identification of complex changes such as LOH without decrease in copy number and decrease in copy number without LOH (Bignell *et al.*, 2004; Raghavan *et al.*, 2005; Zhao *et al.*, 2004). Such changes are common, as demonstrated in pancreatic and cervical cancer cell lines, where the proportion of LOH associated with copy-reduction was found to be just 32% (Calhoun *et al.*, 2006) and 25% (Kloth *et al.*, 2007), respectively.

CGH signal intensities must be normalised to account for technical bias while still retaining biologically relevant changes. Normalisation of array CGH data has generally involved the use of methods originally developed for normalising gene expression microarray data (for review, see Quackenbush, 2002). Cross-slide and within-slide normalisation are used to transform the data such that all arrays, and all the spots on each array, are comparable. In median normalisation, all values are multiplied by a constant factor so that all arrays have a median \log_2 ratio of 0. Lowess, or Loess, normalisation accounts for spot intensity biases and other dependencies such as the location of the spot on the array and the use of different print tips. The data are linearised by subtracting a

Lowess regression curve. A number of additional methods for dealing with spatial effects in expression microarray data are reviewed in Neuvial *et al.* (2006).

In general, array CGH must be more stringent than gene expression analysis because it is required to detect single copy changes and, while the copy number, unlike the expression level, of a gene is expected to be identical in two samples, this is often not the case due to tumour heterogeneity and the presence of contaminating stromal cells (Khojasteh et al., 2005). Khojasteh and coworkers (2005) proposed a multi-step normalisation process specifically for dealing with array CGH data. A "spatial segmentation" algorithm has also been developed to account for array CGH-specific spatial effects designated "local spatial biases", where clusters of spots show a shift in signal, and "continuous spatial gradient", where there is a smooth gradient in signal across the array (Neuvial et al., 2006). Staaf and coworkers (2007) showed that copy number imbalances correlate with intensity in array CGH data and that normalisation of expression data erroneously corrects for biologically relevant gains in copy number. They have therefore developed a normalisation algorithm that prevents suppression of copy number ratios by stratifying the data into separate populations representing discrete copy number levels (Staaf et al., 2007). Array CGH data are also affected by a genome-wide technical artefact termed "spatial autocorrelation", or "wave", for which the peaks and troughs are aligned across samples but the amplitude, and for some samples, the direction, varies (Marioni et al., 2007). Removal of the wave using a Lowess curve led to an increase in the number of biologically relevant CNVs detected in array CGH data from normal individuals (Marioni *et al.*, 2007).

Affymetrix have developed a number of procedures for normalising SNP array CGH data. As described above, each SNP on an Affymetrix array is represented by a probe set comprising multiple probe pairs (Figure 1.4A). Fluorescence on the mismatch probes represents non-specific hybridisation, and the data can be corrected by subtracting the mismatch from the perfect match intensity for each probe pair. The corrected intensities are then averaged across the probe set. The data can be globally normalised by multiplying the average intensity of the experimental array, i.e. the array to which the cancer sample is hybridised, by a normalisation factor to make it numerically equivalent to the average intensity of the control array, to which a normal sample is hybridised. Intensity ratios are calculated by dividing the average intensity for each SNP in the experimental array by the equivalent value in the control array. Three software packages that are commonly used for processing copy number data on Affymetrix SNP arrays are Copy Number Analyser for GeneChip arrays (CNAG, Nannya *et al.*, 2005), DNA-Chip Analyzer (dChip, Zhao *et al.*, 2004) and Affymetrix GeneChip Chromosome Copy Number Analysis Tool (CNAT, Huang *et al.*, 2004). These are compared and reviewed in Baross *et al.* (2007), who concluded that the detection of all real CNVs from a 100K array necessitated the combined use of multiple procedures.

The next step, following normalisation, is to identify regions of copy number change within the CGH data. Many different approaches have been developed for segmenting the genome into regions of homogeneous copy number. These include change-point analysis, where the genome is segmented at points where the copy number changes significantly (Olshen et al., 2004; Venkatraman and Olshen, 2007), Hidden Markov Models (HMMs) (Engler et al., 2006; Marioni et al., 2006; Nannya et al., 2005; Rueda and Diaz-Uriarte, 2007; Shah et al., 2006; Stjernqvist et al., 2007), hierarchical clustering along chromosomes (Wang et al., 2005) and smoothing methods (Hsu et al., 2005; Huang et al., 2007). There are also a number of web-based applications, such as ADaCGH (Diaz-Uriarte and Rueda, 2007) and CGHweb (Lai et al., 2008), for viewing and comparing outputs from multiple algorithms. Further methods have been developed to identify copy number changes specifically in SNP array CGH data, which has increased noise at the probe level compared with BAC array CGH (Yu et al., 2007), and a number of these infer allele-specific copy numbers (Huang et al., 2006a; LaFramboise et al., 2005; Lamy et al., 2007; Nannya et al., 2005; Yu et al., 2007). Some of the methods for detecting copy number changes are discussed in further detail in Section 4.6.

Finally, having identified regions of copy number change, the statistical power can be increased by examining the region across many samples. Unlike for CNVs in normal samples, cross-sample analysis of copy number changes in cancer is hampered by the large size of many rearrangements, variation in the location of breakpoints between samples, and sample heterogeneity that prevents accurate estimation of the copy number (Marioni *et al.*, 2007). A handful of methods have been developed to identify recurrent regions of copy number change in tumours: CMAR (Rouveirol *et al.*, 2006), STAC (Diskin *et al.*, 2006), H-HMM (Fiegler *et al.*, 2007) and KC-SMART (Klijn *et al.*, 2008). The latter is the only algorithm that does not discretise the data into 3 states (loss, gain and no change), which can lead to undetected copy number changes in heterogeneous tumours (Klijn *et al.*, 2008).

Chapter 1

1.3.3.3 Analysis of copy number changes in cancer genomes

CGH can detect an euploidy, gene amplifications and deletions, and non-reciprocal translocations in cancer genomes. Gene amplifications are gains in copy number of restricted regions of DNA (Bignell *et al.*, 2007) that contribute to tumourigenesis by increasing the transcript levels, and therefore the protein levels, of oncogenes (Schwab, 1999). Gene amplification is the major mechanism of oncogenesis for a number of cancer genes, including *MYCN*, which is amplified in ~30% of advanced neuroblastomas (Seeger *et al.*, 1985). Amplified genes represent a promising target for cancer therapy, as demonstrated in breast cancers harbouring an amplified *HER/ERBB2* receptor gene (Cobleigh *et al.*, 1999, see Section 1.2.7).

Deletions are an important mechanism for inactivating tumour suppressor genes, including *PTEN* (Li *et al.*, 1997) and *CDKN2A* (*INK4A/ARF*) (Orlow *et al.*, 1995). A genome-wide analysis of homozygous deletions in over 600 cancer cell lines showed that deletions occur in regions with fewer genes and repeat elements but higher flexibility compared with the rest of the genome (Cox *et al.*, 2005). A significant proportion occur in regions that are prone to chromosome breakage, and some of the genes in these "fragile sites", such as *WWOX* and *FHIT*, show similar mutational patterns to known tumour suppressor genes, so it is not clear whether or not these genes are causally implicated in cancer (Futreal *et al.*, 2004).

Like gene expression analysis, copy number profiling can be used to subcategorise cancers. It can distinguish three subtypes of glioblastoma (Maher *et al.*, 2006), and separates leiomyosarcomas into a distinct cluster from gastrointestinal stromal tumours, which, until recently, were classified as the same tumour type (Meza-Zepeda *et al.*, 2006). It also provides predictive power in breast cancer prognosis, where a poor prognosis is indicated by high-level amplification (Chin *et al.*, 2006), extensive chromosome instability (Fridlyand *et al.*, 2006) and/or the presence of multiple, closely spaced amplicons, or "firestorms", on a single chromosome arm (Hicks *et al.*, 2006). Copy number profiles can also help to stage a tumour, such as in cervical cancer, where gain of chromosome 3q is associated with the transition from severe dysplasia to invasive carcinoma (Kersemaekers *et al.*, 1998). Furthermore, studies in ovarian cancer have revealed an association between drug response and the presence of copy number changes associated with drug sensitivity or resistance (Bernardini *et al.*, 2005; Kim *et al.*, 2007a).

The amplification of genes involved in drug metabolism or inactivation is commonly observed in cultured cells as a means of acquiring drug resistance (Lengauer *et al.*, 1998).

While many cancer genomes have been analysed for copy number changes, there has been limited progress in determining the functional significance of altered regions. One successful approach involves identifying recurrently altered regions that are specific to particular tumour types. This enables the identification of "lineage addiction" cancer genes, which may target essential lineage-specific survival functions and therefore represent promising therapeutic targets (Garraway and Sellers, 2006). Two such genes are the melanoma-specific oncogene MITF, which is selectively amplified and overexpressed in 20% of melanomas (Garraway et al., 2005), and NKX2-1, which lies in the minimal amplified region of a lung-cancer-specific amplicon on chromosome 14q13.3 found in up to 20% of lung cancers (Kendall et al., 2007; Weir et al., 2007). Genes TTF1 and NKX2-8 are usually co-amplified with NKX2-1 in the 14q13.3 amplicon and all three genes have been shown to co-operate in lung tumourigenesis (Kendall et al., 2007). The co-occurrence and mutual exclusivity of copy number alterations at different loci may also reflect co-operating and complementary cancer genes, respectively. For example, gains of *ERBB2* and *CCNE1* frequently co-occur in bladder cancer, while *CCND1* and *E2F1*, which function in the same pathway, are mutually exclusive (Veltman *et al.*, 2003).

The identification of cancer genes in regions of copy number change can be challenging because changes often span large regions of the genome that encompass many genes and may include many attractive candidates. Gains of more than one copy may have involved multiple evolutionary events and the critical gene may reside at the highest peak in copy number, as demonstrated for oncogenes *CYP24* and *ZNF217* in breast cancer (Albertson *et al.*, 2000). Measurement of gene expression is also important for evaluating candidate cancer genes. *SPANXB* was identified as the putative critical gene in an Xq duplication in acute lymphoblastic leukaemias with an *ETV6/RUNX1* translocation since it was the only gene with high and uniform overexpression across all samples (Lilljebjorn *et al.*, 2007). While gene expression and gene dosage are rarely perfectly correlated, many studies, such as the comparison of array CGH and gene expression data in breast cancers, have shown good correlation (Hyman *et al.*, 2002; Pollack *et al.*, 2002). However, genes that are amplified are not necessarily overexpressed, as demonstrated by Kloth and colleagues (2007), who did not observe a genome-wide correlation between copy number and gene expression in cervical cancer cell lines. Gene expression is influenced by factors other

than gene dosage, such as the availability of transcription and regulatory factors, DNA methylation and chromatin conformation, and the presence of miRNAs (Kloth *et al.*, 2007).

The integration of copy number analysis with gene resequencing also facilitates cancer gene identification. Mullighan and colleagues (2007) performed a genome-wide analysis of genetic alterations in 242 paediatric acute lymphoblastic leukaemias (ALL) using 100K and 250K SNP arrays. They found mutations in genes that regulate late B lymphocyte development in 40% of B-progenitor ALL cases. *PAX5* mutations, which included deletions, point mutations and translocations, were identified in 32% of cases (Mullighan *et al.*, 2007). ALL genomes are relatively stable, but genomes harbouring different translocations show variability in the number of copy number changes, which may reflect differences in the number of events required for tumourigenesis (Mullighan *et al.*, 2007; Wang and Armstrong, 2007). The integration of resequencing data, and epigenetic data (see Section 1.3.4), can facilitate the identification of tumour suppressor genes in regions of LOH, where the other allele may be inactivated by point mutation or epigenetic changes.

The identification of human cancer genes is aided by the integration of complementary genome-wide analyses of human cancers, but the integration of cancer-associated mutation datasets from other species, particularly the mouse, provides an even more powerful approach for cancer gene discovery. Cross-species comparisons are discussed in Section 1.5.

1.3.3.4 Limitations of CGH and alternative strategies

Limitations of CGH-based approaches include difficulties in determining the ploidy of the sample and identifying the location of rearranged sequences in the cancer genome. However, the ploidy and location of larger rearrangements (> 10 Mb) can be discerned by combining CGH with G-banding or Spectral Karyotyping (SKY) (Watson *et al.*, 2007). CGH may also struggle to detect low level changes and changes in heterogeneous samples, e.g. primary cancers containing normal stromal cells, and it is affected by low-copy reiterated sequences, including gene paralogues (for full review, see Pinkel and Albertson, 2005).

A further limitation of CGH is that while it can detect nonreciprocal, or unbalanced, translocations, which result in the gain or loss of DNA and often cause the inactivation of tumour suppressor genes (Mitelman *et al.*, 2004), it cannot detect reciprocal, or balanced, translocations. These result in fusion transcripts or transcriptional deregulation due to the positioning of an intact gene next to promoter and/or enhancer elements of another gene. It has recently been discovered that cytogenetically balanced translocations are frequently associated with focal copy number alterations, suggesting that high-resolution array CGH may in fact be capable of detecting a proportion of balanced translocations in cancer (Watson *et al.*, 2007). However, truly balanced translocations cannot be identified.

Balanced translocations are often initiating events in tumourigenesis that are essential for tumour development, and they therefore represent promising therapeutic targets (see Section 1.2.7). Until recently, it was thought that balanced translocations predominated in haematopoietic tumours, but an assessment of data in the Mitelman Database of Chromosome Aberrations in Cancer suggests that they also play an important role in epithelial tumourigenesis (Mitelman et al., 2004). Furthermore, human solid tumours appear to contain large numbers of gene fusions (Volik et al., 2006) and a quarter of the breakpoints detected in 3 breast cancer cell lines were found to be balanced (Howarth et al., 2008). The high-throughput identification of balanced translocations has been hindered because translocation breakpoints cannot be amplified by PCR (Howarth et al., 2008). Genome-wide techniques for identifying translocations include array painting, in which chromosomes are sorted and DNA is amplified and hybridised to DNA microarrays (Howarth et al., 2008), and informatics approaches, such as the algorithm developed by Tomlins and coworkers (2005) that used RNA expression data to identify candidate gene fusions in prostate cancers. The EML4-ALK fusion was identified in nonsmall cell lung cancers by paired-end sequencing (Soda et al., 2007).

End-sequence profiling (ESP) can be used to precisely map all types of genomic rearrangements, including balanced translocations (Volik *et al.*, 2003). ESP involves constructing a BAC library from the cancer genome and sequencing the ends of clones to identify rearrangements, which map to locations in the reference genome that are of abnormal distance or orientation (Volik *et al.*, 2003; Figure 1.5). The method can also identify fusion transcripts (tESP) and can be targeted to specific amplicons (Volik *et al.*, 2006). Complete sequencing of the BACs enables detailed analysis of the structure of genomic rearrangements and elucidation of the mechanisms of rearrangement.

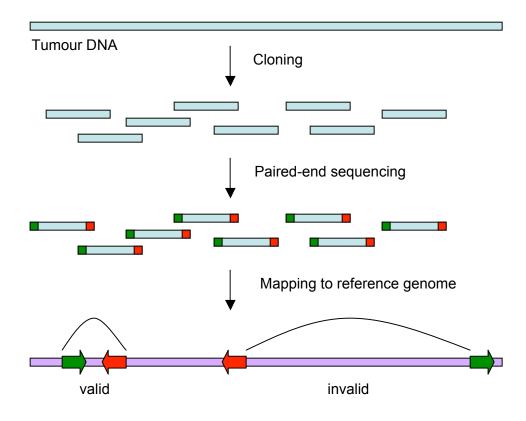


Figure 1.5. End sequence profiling of tumour DNA. 100-250 kb regions of the tumour genome are cloned and a 500 bp region at the end of each clone is sequenced. The ends are mapped to the human reference genome. Ends that are an abnormal distance apart or in an abnormal orientation, shown here as "invalid", are indicative of rearrangements within the tumour genome. Redrawn with minor modifications from Figure 1 of Raphael *et al.* (2008).

ESP-based analysis of 4 cancer amplicons revealed evidence for sister chromatid breakfusion-bridge cycles, the excision and reintegration of double minutes (extrachromosomal DNA), and more complex architectures involving clusters of small genomic fragments (Bignell *et al.*, 2007). Break-fusion-bridge cycles are initiated by a double-strand chromosomal break, which, following DNA synthesis, results in sister chromatids with identical free DNA ends that fuse to one another to prevent apoptosis. An anaphase bridge is formed during chromatid separation in mitosis, and this results in a new doublestrand break and reinitiation of the cycle (McClintock, 1941).

ESP analysis of 6 epithelial cancers, including primary tumours from brain, breast and ovary, plus a metastatic prostate tumour and 2 breast cancer cell lines, revealed extensive chromosomal rearrangements, some of which appeared to be recurrent (Raphael *et al.*, 2008). Despite the benefits of this strategy, sequencing large numbers of clones across many cancer genomes is costly and impractical. However, Bashir and colleagues (2008) have derived a formula to maximise the probability of detecting fusion genes with the least amount of sequencing. The formula depends on the distribution of gene lengths and the parameters of the sequencing strategy used (Bashir *et al.*, 2008). A high-throughput alternative to ESP, which involves massively parallel sequencing of the ends of randomly sheared DNA, has recently been applied to the genome-wide analysis of somatic and germline rearrangements in 2 lung cancers (Campbell *et al.*, 2008). The analysis revealed a wide spectrum of rearrangements, as well as providing high-resolution copy number information. Paired-end sequencing is an attractive strategy for the complete characterisation of rearrangements in cancer.

1.3.4 Epigenetic profiling

Epigenetic changes are chemical modifications to the DNA or histones that change the structure of chromatin but do not alter the DNA sequence. If chromatin is in the condensed conformation, transcription factors cannot access the DNA and genes are therefore not expressed, whereas genes in open chromatin can be expressed as required. DNA methylation and changes in chromatin conformation have both been implicated in tumourigenesis. DNA methylation of CpG islands, which are located in promoter regions, can result in gene "silencing" by preventing transcription factor binding. It can also repress gene expression by recruiting methyl-binding domain proteins, which

associate with histone deacetylases (HDACs). HDACs mediate chromatin condensation by deacetylating histones. See Pelengaris and Khan (2006).

Aberrant DNA methylation of *CDKN2A* has been observed in a wide range of common cancer types (Herman *et al.*, 1995; Merlo *et al.*, 1995), while *VHL* and *BRCA1* are silenced by methylation in a significant proportion of kidney (Herman *et al.*, 1994) and breast and ovarian cancers (Esteller *et al.*, 2000), respectively. *VHL* and *BRCA1* are also frequently mutated in cancer, but for other tumour suppressor genes, such as *RASSF1A*, promoter hypermethylation appears to be the principal mechanism for inactivation (for review, see Jones and Baylin, 2002).

Detection of DNA methylation relies on the ability to distinguish cytosine from 5-This can be achieved using restriction enzymes that restrict only methylcytosine. unmethylated DNA, or by using sodium bisulfite, which converts unmethylated cytosines to uracil, or by immunoprecipitation of methylated DNA using 5-methylcytosine-specific antibodies or methyl-binding domain proteins (see Down et al., 2008). All three approaches can be applied to the genome-wide detection of DNA methylation through the use of oligonucleotide arrays. However, restriction enzyme-based methods are limited to the analysis of CpG sites that contain the recognition site for the enzyme in use, while bisulfite conversion reduces the complexity of the DNA and so reduces the number of unique probes that can be used on the array (Down et al., 2008). Bisulfite conversion and methylated DNA immunoprecipitation have also been combined with next-generation sequencing in techniques known as BS-seq (Cokus et al., 2008) and MeDIP-seq (Down et al., 2008), respectively. Histone modifications can be detected using chromatin immunoprecipitation (ChIP), which is described in Section 1.3.5.

Large genomic regions, such as an entire chromosome arm, can show aberrant methylation in cancer (Frigola *et al.*, 2006), and there is evidence to suggest that some cancers show a CpG island methylator phenotype (CIMP). CIMP+ colorectal cancers have significantly more hypermethylation at CpG islands, including an increased incidence of *CDKN2A* and *THBS1* methylation (Toyota *et al.*, 1999), and they are characterised by a methylated mismatch repair gene, *MLH1*, which gives rise to microsatellite instability (Weisenberger *et al.*, 2006; see Section 1.2.5.1.3 for a description of microsatellite instability). Genes that are reversibly repressed by Polycomb proteins in embryonic stem cells are significantly over-represented amongst constitutively

hypermethylated genes in colorectal cancers (Widschwendter *et al.*, 2007). This provides support for the theory of a stem cell origin of cancer (Section 1.2.3.2). A detailed discussion of the epigenomics of cancer is beyond the scope of this thesis, which focuses on changes in cancer that alter the DNA sequence. Epigenomics approaches are reviewed in Callinan and Feinberg (2006) and, for a detailed review of epigenomics and its relevance to the cancer stem cell hypothesis, see Jones and Baylin (2007).

1.3.5 Genome-wide mapping of transcription factor binding sites

The mapping of transcription factor binding sites (TFBS) across the whole genome can help to elucidate gene regulatory networks. Chromatin immunoprecipitation (ChIP) is a powerful approach for analysing TFBS in living cells (Wei *et al.*, 2006). Cells are treated with formaldehyde to mediate the formation of cross-links between DNA and proteins. The chromatin is then fragmented by sonication and an antibody against the transcription factor of interest is used to immunoprecipitate the transcription factor bound to DNA (see Loh *et al.*, 2006). The precipitated DNA can be used to probe a DNA microarray in a high-throughput method known as ChIP-chip. This approach has been used to map TFBS in the yeast genome (Ren *et al.*, 2000). For more complex genomes, it has been necessary to restrict analysis to specific regions, such as promoter regions or individual chromosomes (Boyer *et al.*, 2005; Cawley *et al.*, 2004; Horak *et al.*, 2002; Weinmann *et al.*, 2002), but more recent analyses have used ChIP-chip to survey the entire genome (Kim *et al.*, 2005b; Lee *et al.*, 2006).

An alternative approach involves cloning and sequencing the precipitated DNA fragments, and then mapping the sequences to the genome. Initially, this involved the sequencing of individual fragments sampled from the DNA pool (Hug *et al.*, 2004; Weinmann *et al.*, 2001). However, high coverage is required to distinguish real binding sites from background DNA, and this has been achieved at reduced cost by sequencing a "tag" from each DNA fragment by serial analysis of gene expression (SAGE) (Chen and Sadowski, 2005; Impey *et al.*, 2004; Kim *et al.*, 2005a; Roh *et al.*, 2005). To overcome the problems of ambiguity associated with mapping short tags, Wei and coworkers (2006) developed an approach called ChIP-PET, in which ChIP is coupled with paired-end ditag (PET) sequencing so that both the 5' and 3' ends of each DNA fragment are sequenced (Figure 1.6). This method was applied to the unbiased global mapping of 542 p53 binding sites in the human genome (Wei *et al.*, 2006). The functions of p53 target genes

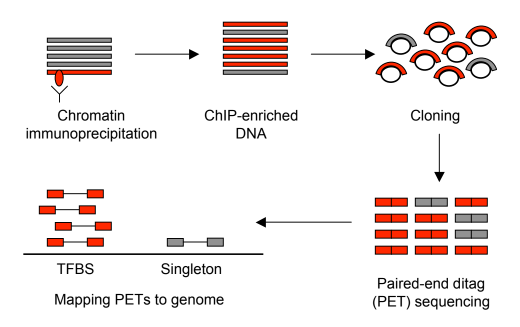


Figure 1.6. Overview of ChIP-PET for mapping transcription factor binding sites. In chromatin precipitation (ChIP), the chromatin is fragmented by sonication and an antibody against the transcription factor of interest is used to precipitate the transcription factor bound to DNA. The ChIP-enriched DNA is cloned and the ends of each clone are sequenced to create a library of paired-end ditags (PETs). The PETs are mapped to the reference genome. Multiple PETs mapping to a single location indicate the presence of a transcription factor binding site (TFBS) at that location. Redrawn with modifications from Figure 1 of Loh *et al.* (2006).

included known roles of p53, such as apoptosis, DNA repair and transcription regulation, but also novel functions, such as cell adhesion and mobility (see Wei *et al.*, 2006).

Loh and coworkers (2006) applied the ChIP-PET technology to the global mapping of Oct4 and Nanog binding sites within mouse embryonic stem (ES) cells. Oct4 and Nanog are required for the maintenance of ES cell pluripotency and self-renewal and may play an important role in cancer (see Section 1.2.3.2). Approximately 1,000 and 3,000 high confidence binding sites were identified for Oct4 and Nanog, respectively, and the presence of one or other binding site was found to be associated with genes that are repressed and induced during differentiation. The target genes include known effectors of ES cell fate, such as *Foxd3* and *Setdb1*, genes required for maintaining pluripotency, including *Esrrb* and *Rif1*, and *Mycn*, which is involved in ES cell self-renewal and proliferation. Most of the Oct4 binding sites also bind Sox2, suggesting that Oct4 and Sox2 co-operate in regulating gene expression.

ChIP-PET has also been used in human B cells to identify more than 4,000 potential binding sites for Myc, of which 668 were identified as direct targets of Myc regulation (Zeller *et al.*, 2006). Many of the target genes are involved in protein synthesis and cell metabolism, which is consistent with a role for Myc in controlling cell size. A large number of transcription factors were also identified. This study showed a weak overlap with other analyses of Myc binding sites, reflecting the current limitations of ChIP-PET, such as the limited sensitivity of PET detection, the experimental noise associated with ChIP, and the fact that the analysis only describes a snapshot of transcription factor binding at a particular moment in time (Zeller *et al.*, 2006). A comparative study of STAT1 binding sites identified by ChIP-chip and ChIP-PET found a considerable overlap between methods, but each method also identified unique sites, suggesting that higher accuracy could be achieved by using both techniques (Euskirchen *et al.*, 2007).

The most advanced method for identifying TFBS is ChIP-seq, in which the DNA fragments isolated by ChIP are amplified and sequenced using next-generation sequencing technology. ChIP-seq requires less starting material and involves fewer steps, making it faster and less prone to error. ChIP-seq using Solexa massively parallel sequence identified STAT1 binding sites in human HeLa S3 cells with an estimated sensitivity of 70-92% and specificity of at least 95% (Robertson *et al.*, 2007).

1.4 Cancer gene discovery in the mouse

1.4.1 The mouse as a model for studying cancer

1.4.1.1 Background

The mouse is a leading model system for cancer research because it has a rapid reproduction rate and breeds well in captivity and, owing to its small size, it can be maintained in large numbers in limited space (see Frese and Tuveson, 2007). It is also genetically and physiologically similar to human. In light of these factors, the mouse genome has been sequenced and annotated to a high standard, second only to that of human (Waterston *et al.*, 2002).

The mouse was initially used as a cancer model through tumour transplantation within inbred strains, but following the discovery of the immunodeficient "nude" mouse and, later, the severe combined immunodeficient (SCID) mouse, it became possible to transplant human tumours into the mouse, creating xenograft models. Such models can be used to rapidly assess tumour tissue and cell lines *in vivo* but they do not fully recapitulate the behaviour of an endogenous tumour because many features of the tumour microenvironment, such as stromal cells, vasculature and immune cells, are missing. The tumour xenograft is also likely to be less heterogeneous than the endogenous tumour because cells in culture are under high selective pressure. These factors have contributed to the limited success of xenograft models in drug development (for review, see Sharpless and Depinho, 2006)

Many inbred strains that spontaneously develop cancer at high frequency have been established, and these, as well as mice that have been treated with a mutagen, are useful for studying the properties of endogenous cancers *in vivo*. They have been used to identify cancer genes and to assess the effects of carcinogens and therapeutic compounds. However, these models may be biased towards specific types of tumour that show variable penetrance and latency and do not accurately reflect common human cancers (Frese and Tuveson, 2007).

1.4.1.2 Genetically engineered mouse models

Genetically engineered mouse models represent a major advance in cancer research that allows for the study of gene function *in vivo* and for the creation of models that more accurately recapitulate human cancers. Genetically engineered models can be classified as transgenic or endogenous (Frese and Tuveson, 2007).

1.4.1.2.1 Transgenic models

Transgenic mice can be created to study the effect of overexpressing an oncogene or a dominant-negative tumour suppressor gene, which encodes a mutant tumour suppressor that can inactivate the wildtype protein. Transgenic mice can be generated by pronuclear microinjection, in which a construct containing the gene of interest (transgene) is microinjected into the mouse oocyte after fertilisation and randomly integrates into the genome, usually in tandem copies. If the transgenic cells contribute to the germ line, the genetic change can be transmitted to the next generation, producing mice that are fully transgenic and establishing a strain. Many genes involved in cancer development are also essential for mouse development. Therefore, to prevent embryonic lethality and to restrict overexpression to specific tissues, the construct containing the gene of interest also contains promoter elements designed for spatial and temporal restriction of gene expression. For example, the Tet-On and Tet-Off systems (Baron and Bujard, 2000) promote gene expression in the presence and absence, respectively, of doxycycline, a non-toxic analogue of tetracycline, while fusing the gene of interest to a gene encoding the oestrogen receptor binding domain results in an inactive protein that is activated upon treatment with Tamoxifen (Eilers et al., 1989).

Limitations of the microinjection method include the possibility that, because the transgene integrates randomly, it could disrupt other genes, resulting in a phenotype that does not reflect the function of the gene of interest (for review, see Muller, 1999). In addition, the tendency of the transgene to integrate in multiple copies could result in excessive overexpression that is toxic to the animal (Muller, 1999). However, transgenic mice have made a significant contribution to cancer research. In the earliest examples, mouse models were used to demonstrate the role of oncogenes in cancer. For example, tissue-specific overexpression of the Myc oncogene in mammary glands and B-cells resulted in the generation of mice prone to breast cancer (Stewart *et al.*, 1984) and

lymphomas (Adams *et al.*, 1985), respectively. Overexpression of dominant-negative mutant tumour suppressor genes has also proved effective, e.g. a gene encoding mutant type II transforming growth factor beta (Tgf β) receptor has been shown to accelerate chemically induced tumourigenesis in the mammary gland and lung (Bottinger *et al.*, 1997).

1.4.1.2.2 Endogenous models

A knockout mouse can be created to study the effect of inactivating a tumour suppressor gene. In this method, a targeting vector is transfected into embryonic stem (ES) cells, which are harvested from the inner cell mass of mouse blastocysts. The vector must share homology with the region of the mouse gene that is being targeted, i.e. the tumour suppressor gene of interest, and must also contain genes for selection, such that only cells in which the vector DNA has replaced the endogenous DNA by homologous recombination will survive. The surviving ES cells are injected back into a blastocyst, and will contribute to all cell lineages, including the germ line (Robertson *et al.*, 1986). The targeting vector can be engineered to knock out the whole gene or part of a gene, or small changes can be introduced into the gene sequence. Alternatively, the complete gene under the control of a strong promoter can be introduced to create a knockin mouse for overexpressing oncogenes. By targeting a single copy to the genome, this overcomes the problems associated with pronuclear microinjection. (For review, see Muller, 1999).

As with transgenic mice, mutations can be spatiotemporally regulated. Conditional mouse models frequently use the Cre-lox system from bacteriophage P1, in which Cre recombinase catalyses recombination between loxP sites (Sauer and Henderson, 1988), and the intervening DNA is deleted or inverted, depending on the orientation of the sites (Lakso *et al.*, 1992). loxP sites can therefore be placed on either side of a gene region to remove that region in the presence of Cre (Figure 1.7). Large-scale chromosomal deletions and inversions can also be generated by placing loxP sites further apart on the chromosome (Kmita *et al.*, 2000; Smith *et al.*, 2002), while chromosomal translocations can be created by placing a loxP site at each breakpoint (Forster *et al.*, 2003). Conditional oncogene expression can be achieved by inserting a stop cassette, which is flanked by loxP sites, between the promoter and the first exon such that Cre-mediated excision of the cassette results in expression of the gene (de Alboran *et al.*, 2001; Jackson *et al.*, 2001).

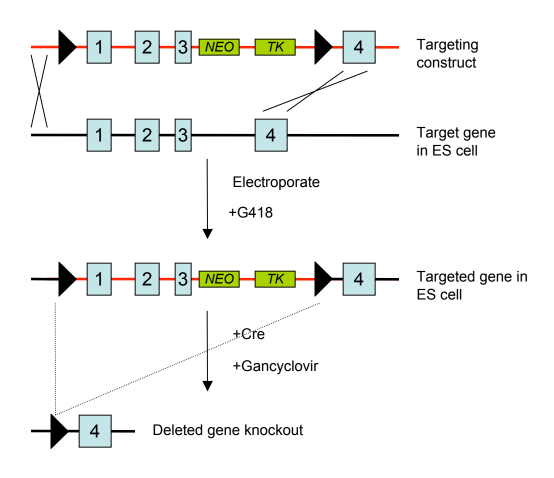


Figure 1.7. Generation of a conditional knockout allele in ES cells. A targeted gene construct is designed that contains loxP sites flanking the region of the gene to be deleted as well as genes for selection. Upon introduction into ES cells, DNA in the construct replaces endogenous DNA in the target gene by homologous recombination. The addition of G418 selects for cells that express the *Neomycin* gene, and therefore contain the knockout construct. The addition of Cre results in recombination between the loxP sites, removing the region of the gene containing exons 1, 2 and 3 and the *Neomycin* and *tk* genes. Gancyclovir kills cells expressing *tk*, and therefore selects cells in which recombination has occurred and the gene has been knocked out.

Unlike the conditional expression systems in transgenic mice, once Cre recombinase has been expressed, the change is irreversible, and there is evidence to suggest that Cre can be cytotoxic, perhaps due to recombination at pseudo-loxP sites (see Jonkers and Berns, 2002). In addition, the Cre-lox system cannot generate conditional point mutations, and this represents a significant limitation since point mutations and deletions do not always produce the same phenotype (Frese and Tuveson, 2007). However, the Cre-lox system has proved invaluable in creating models that would otherwise not arise or survive. For example, homozygous Brcal and Brca2 knockouts die early in embryogenesis, and heterozygous mice are not tumour-prone, but mice harbouring a Cre-mediated deletion of Brcal (Xu et al., 1999) or Brca2 and Trp53 (Jonkers et al., 2001) in the adult mammary gland do develop mammary tumours. Likewise, Trp53 mutations have been identified in many types of human cancer, but if *Trp53* is mutated in all cells, the mouse is most likely to develop lymphomas or sarcomas. Conditional Trp53 mutations can be used to create models for human cancers that are driven by TP53 mutation in other tissues (Jonkers and Berns, 2002). The Flp/FRT system from Saccharomyces cerevisiae is an alternative to Cre-lox that works in a similar way.

1.4.1.3 Mouse models in drug discovery

Mouse models that faithfully recapitulate human cancers are important for developing and testing therapeutic drugs. Studies on a mouse model for acute promyelocytic leukaemia (APL) have resulted in the development of an effective, retinoic-acid-based treatment for the disease (Lallemand-Breitenbach *et al.*, 1999; Soignet and Maslak, 2004). Mouse models can also be used to identify predictive markers of disease response and progression, and to understand drug toxicity and resistance. They have proved particularly useful in the study of oncogene addiction, which is an important consideration in drug target validation (see Section 1.2.7). Mouse models have demonstrated the requirement for persistent expression of *Hras*, *Myc*, *Bcr-Abl*, *Erbb2* and *Fgf7* in the maintenance of melanoma (Chin *et al.*, 1999), haematopoietic tumours (Felsher and Bishop, 1999), B-cell lymphoma and leukaemia (Huettner *et al.*, 2000), breast cancer (Xie *et al.*, 1999), and lung cancer (Tichelaar *et al.*, 2000), respectively.

1.4.1.4 Mouse models in cancer gene discovery

The methods described in Section 1.3 can also be applied to the identification of candidate cancer genes in the mouse. For example, array CGH has been used to identify regions of copy number change in mouse models of malignant melanoma (O'Hagan *et al.*, 2003) and pancreatic islet carcinomas (Hodgson *et al.*, 2001). However, as with human cancers, by the time the cancer has presented, it is difficult to distinguish the important driver mutations from the background of passenger mutations.

The genetically engineered mouse models discussed thus far are useful for studying the function of a particular gene or for representing a specific human cancer, but the tumours in these models do not evolve naturally. In general, the initiating event, i.e. the engineered mutation, is present throughout a tissue, whereas in natural tumourigenesis, the tumour develops from one mutated cell (see Section 1.2.3). Likewise, in mouse models used to study the combined action of multiple genes in cancer, the genes of interest are usually simultaneously mutated, whereas "natural" tumours progress through a multi-step process, where mutations are gradually acquired. Finally, many mouse models are designed to show high penetrance and short latency to keep costs down, but as a result they may not possess many of the co-operating oncogenic events that would eventually be acquired by a naturally evolving tumour (for review, see Frese and Tuveson, 2007; Sharpless and Depinho, 2006).

It is important that the mutations in mouse models used to identify novel cancer genes reflect the mutations found in human cancers, and this requires more accurate modelling of the natural evolution of tumours.

1.4.2 Forward genetic screens in the mouse

Forward genetic screens using somatic mutagens are a powerful approach for cancer gene discovery in which tumours undergo a process of evolution that mirrors that of human tumour formation. They allow for relatively unbiased, genome-wide identification of both novel cancer genes and collaborations between genes involved in cancer. Chemical mutagenesis is highly efficient but mutations are very difficult to identify. Insertional mutagenesis by retrovirus or transposon is an effective alternative approach in which the mutagen acts as a molecular tag for easy identification of the mutated allele.

1.4.2.1 Retroviral insertional mutagenesis

1.4.2.1.1 Mechanisms of mutagenesis

The slow transforming retroviruses murine leukaemia virus (MuLV) and mouse mammary tumour virus (MMTV) have been widely used for insertional mutagenesis in the mouse. Unlike acute transforming retroviruses, which induce tumours by expression of a viral oncogene, slow transforming retroviruses do not carry an oncogene, and tumours are induced by mutations caused by insertion of the retrovirus into the host genome. Consequently, tumours develop with a longer latency of 3-12 months, compared with 2-3 months for acute transforming retroviruses (Uren *et al.*, 2005). MMTV was identified as a causative agent in several strains of mice that were prone to mammary tumours, while MuLV was identified as a causative agent in the lymphoma-prone AKR mice (see Weiss, 2006). The principal dataset used in this thesis was generated using MuLV, and this mutagen is therefore the main focus of the background provided herein.

Retroviruses infect host cells by binding of the viral envelope proteins to cell surface receptors. Once the retrovirus has inserted into the host genome, forming a provirus, it will produce viral envelope proteins that occupy the cell surface receptors and prevent reinfection of the same cell. However, recombination with endogenous viral sequences results in the production of envelope proteins that bind to other receptors. This, combined with the fact that many proviruses have defective envelope coding sequences, enables retroviruses to reinfect the same cell, resulting in the accumulation of mutations. Mutations that confer a growth advantage on the cell co-operate in tumour formation, and the process therefore recapitulates the multi-step progression of human tumours (for review, see Mikkers and Berns, 2003; Uren *et al.*, 2005, see also Section 1.2.3).

The MuLV provirus consists of viral genes flanked by two long terminal repeats (LTRs), which are composed of three parts: U3, R and U5 (see Uren *et al.*, 2005; Figure 1.8). Elements within the LTRs drive expression of the viral genes but can also disrupt host genes. U3 contains enhancer and promoter sequences, while R contains transcription start and termination sites. High levels of viral transcription and, therefore, host gene disruption, will only occur in cells containing transcription factors that bind to U3. The propensity of MuLV to induce T- and B-cell lymphomas can be attributed to its dependence upon T- and B-cell-specific transcription factors, including *Runx*, *Ets* and *Myb* (see Neil and Cameron, 2002).

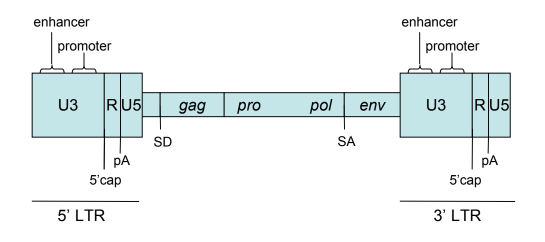


Figure 1.8. Structure of a retroviral provirus. The provirus contains two long terminal repeats (LTRs) flanking the genes required for viral assembly. Elements within the LTRs drive transcription of the viral genes but can also induce mutation of nearby cellular genes. Splicing of a viral splice donor (SD) or cryptic splice donor (not shown) to a splice acceptor or cryptic splice acceptor in the first intron or 5' UTR of a cellular gene results in the formation of a chimeric transcript, in which the cellular gene is coupled to the viral promoter. Splicing of a splice donor or cryptic splice donor in a cellular gene to a viral splice acceptor (SA) or cryptic splice acceptor (not shown) can cause premature termination of gene transcription owing to the presence of polyadenylation signals (pA) and cryptic polyadenylation signals (not shown) in the LTR. Adapted from Figure 1 of Uren *et al.* (2005). Figure is not to scale.

Retroviruses can mutate host genes in a number of different ways. The most common mechanism is enhancer mutation, where one of the U3 enhancers upregulates expression of host genes, which may be some distance away from the retroviral insertion (Figure 1.9A). Most proviruses causing enhancer mutations are found upstream of the mutated gene in the antisense orientation or downstream in the sense orientation. Several possible explanations for the directionality of the enhancer are that upregulation of the host gene may be impeded if the viral promoter intercepts the viral enhancer and host gene, or that viral enhancers may only be functional if they are not transcribed (Clausse *et al.*, 1993; see Uren *et al.*, 2005). *Myc* and *Gfi1* are frequent targets of enhancer mutation in retroviral insertional mutagenesis (Akagi *et al.*, 2004; Corcoran *et al.*, 1984; Selten *et al.*, 1984). *Myc* is mutated in many types of human cancer. It encodes a transcription factor that is thought to regulate the expression of 15% of all genes, including genes involved in cell division, cell growth and apoptosis (see Gearhart *et al.*, 2007). Gfi1 is a zinc finger transcriptional repressor that is involved in cell fate determination and differentiation, including in T- and B-cells (Rathinam and Klein, 2007; Yucel *et al.*, 2003).

An alternative mechanism of mutagenesis is promoter mutation, where the retrovirus inserts in the sense orientation into the promoter region of a host gene (Figure 1.9B). This uncouples the host gene from its own promoter and places it under the control of the viral promoters, resulting in the production of elevated levels of the wildtype protein from chimeric transcripts comprising part of the viral sequence and the complete coding region of the host gene (Mikkers *et al.*, 2002). Promoter mutations led to identification of *Evi1* as a potential oncogene (Copeland and Jenkins, 1990; Mucenski *et al.*, 1988a; Mucenski *et al.*, 1988b). *EVI1* encodes a zinc finger transcription factor that is frequently overexpressed in human myeloid malignancies. It is involved in several recurrent rearrangements, including 2 translocations that result in the fusion transcripts *AML1/MDS1/EVI1* and *ETV6/MDS1/EVI1*, where *MDS1* and *EVI1* are also expressed as a readthrough transcript in normal tissues (for review, see Wieser, 2007).

The retrovirus contains a polyadenylation signal within the R region of the LTR and a cryptic polyadenylation signal in the antisense orientation. Therefore, intragenic retroviral insertions in both orientations can cause premature termination of gene transcription. Insertions within the 3' UTR that truncate a transcript such that mRNA-destabilising motifs are removed will give rise to a more stable transcript and, as a result, increased levels of the wildtype protein (see Uren *et al.*, 2005; Figure 1.9C). Oncogenes

42

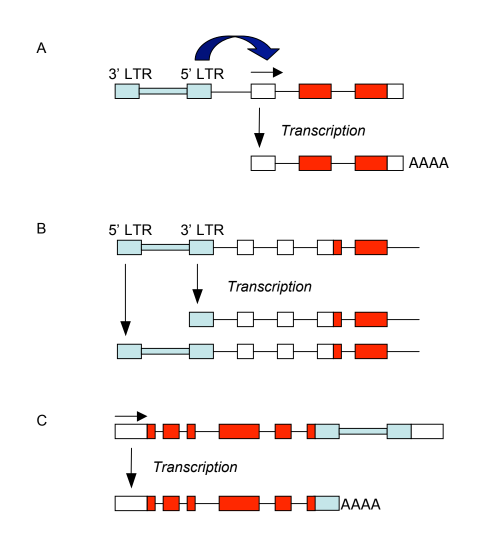


Figure 1.9. The mechanisms of mutagenesis of murine leukaemia virus include enhancer mutation (A), promoter mutation (B) and premature termination of gene transcription (C). The provirus is shown in blue; coding and non-coding exons are shown in red and white, respectively. A. An enhancer element in the 5' LTR of murine leukaemia virus (MuLV) can cause upregulation of nearby cellular genes. Oncogenic insertions of this type are most frequently found upstream and in the antisense orientation with respect to the cellular gene(s) that they are mutating. B. Insertion of MuLV into the promoter region of a cellular gene results in chimeric transcripts that are produced at higher levels than the endogenous gene transcript. C. Intragenic MuLV insertions can cause premature termination of gene transcription, resulting in either gene upregulation or gene inactivation. The figure shows an insertion within the 3' UTR region, which may remove mRNA-destabilising motifs, thereby stabilising the gene transcript. Adapted from figures in Uren *et al.* (2005).

Pim1 and *Mycn* are frequently mutated in this way (Cuypers *et al.*, 1984; Selten *et al.*, 1985; van Lohuizen *et al.*, 1989). *PIM1* encodes a serine/threonine kinase that is frequently overexpressed in human prostate cancer (Dhanasekaran *et al.*, 2001), while *MYCN* encodes a transcription factor related to *MYC* that is amplified in a variety of human tumours, most notably neuroblastomas (Brodeur *et al.*, 1984, 1985).

Intragenic insertions can also activate a gene by causing C-terminal or N-terminal truncation of the encoded protein. Insertions in oncogenes *Myb* and *Notch1* cause both N-terminal and C-terminal truncations (Rosson *et al.*, 1987; Uren *et al.*, 2005). C-terminally truncated Notch1 lacks the destabilising PEST domain and is therefore produced at increased levels, while N-terminal truncations remove the extracellular domain, resulting in a constitutively active intracellular domain expressed from the viral promoter or from a cryptic promoter in *Notch1* (Hoemann *et al.*, 2000). Activating mutations within the extracellular and PEST domains of NOTCH1 have been observed in human T-cell acute lymphoblastic leukaemia (Weng *et al.*, 2004), in which NOTCH1 plays an important role (see Section 1.2.5.1.4 for further details). Analysis of the distribution of insertions within an oncogene may therefore help to explain how the gene is mutated in human cancer.

Intragenic insertions may also cause gene inactivation, either through premature termination of transcription or by disrupting gene splicing (see Uren et al., 2005). It is therefore possible to identify tumour suppressor genes by retroviral insertional mutagenesis, although they are found much less frequently than oncogenes because both copies of the gene must be inactivated. Mutation at the NfI locus is observed in acute myeloid leukaemias in BXH2 mice (Largaespada et al., 1996), which contain MuLV insertions (Bedigian et al., 1984), while in an insertional mutagenesis screen of Blmdeficient mice, 11 genes met the criteria for tumour suppressor genes, including *Rbl1* and *Rbl2*, which are paralogues of *Rb1* (Suzuki *et al.*, 2006). *Blm*-deficient mice have a mutation in the RecQ protein-like-3 helicase gene (Ellis et al., 1995) and show a predisposition to cancer due to increased frequencies of mitotic recombination (Luo et al., 2000). There is an increased likelihood of finding tumour suppressor genes in these mice because they have a higher probability of a normal allele being lost so that only one insertion is required to inactivate the gene (Luo et al., 2000). However, candidate tumour suppressor genes still only accounted for 5% of all genes identified in the screen by Suzuki et al (2006). In theory, insertional mutagenesis screens should have a better

chance of finding haploinsufficient tumour suppressor genes, but none have yet been unambiguously identified (Uren *et al.*, 2005).

Insertional bias could also account for the paucity of tumour suppressor genes identified in retroviral screens. MuLV shows a strong preference for integration near to the transcription start sites of actively transcribed genes (Wu *et al.*, 2003) and is therefore less likely to disrupt a gene by intragenic insertion. However, it is possible that promoter mutations could also cause gene inactivation, as CpG islands in the retroviral LTRs are methylation targets, and DNA methylation could "spread" to CpG islands in the host gene, resulting in gene silencing (see Touw and Erkeland, 2007). Retroviruses prefer to insert into open chromatin (Muller and Varmus, 1994; Pryciak and Varmus, 1992), but different retroviruses show different target site preferences, suggesting that virus-specific interactions are involved (Mitchell *et al.*, 2004). DNA sequence does not seem to influence target site selection (Bushman *et al.*, 2005). The tendency for MuLV to insert into the promoter region indicates that the retrovirus interacts with cellular proteins bound near start sites (Mitchell *et al.*, 2004; Wu *et al.*, 2003).

1.4.2.1.2 Identifying candidate cancer genes

The retroviral insertions act as tags for identifying the mouse genes that are mutated by insertional mutagenesis, and sequencing of the mouse genome and the development of high-throughput genomic techniques have made it possible to identify hundreds or thousands of insertions in a single screen. Insertion sites were initially identified using methods that involved Southern blot analysis and genomic library screening, followed by genome walking to find the mutated gene (see Neil and Cameron, 2002; Uren *et al.*, 2005). However, these have been replaced by PCR-based methods, in which mouse genomic DNA flanking the insertion sites is amplified and is then mapped back to the genome. One such method, known as viral insertion site amplification (VISA) involves using a PCR primer designed to bind to the MuLV LTR and a degenerate, restriction-site-specific primer that enables amplification of the DNA between the insertion and a nearby restriction site (Hansen *et al.*, 2000; Weiser *et al.*, 2007). In inverse PCR and linker-mediated PCR-based methods, the genomic DNA is restriction-digested prior to PCR amplification.

In inverse PCR (Figure 1.10A), the digested genomic DNA is allowed to ligate to itself to form a circular template. PCR primers bind to the retroviral DNA and point out towards the genomic sequence, resulting in amplification of genomic DNA directly flanking the retrovirus (Ochman *et al.*, 1988; Triglia *et al.*, 1988). Only DNA fragments that are a suitable length for efficient circularisation and for PCR amplification will be detected (Uren *et al.*, 2005).

In linker-mediated PCR, rather than the digested DNA ligating to itself, it is ligated to a linker, and this enables shorter insertions to be identified. One primer is designed to bind to the linker, and the other binds to the retroviral sequence. A number of methods have been developed, each with a different approach for avoiding amplification of DNA that has linkers at both ends but contains no retroviral DNA. Vectorette PCR involves the use of a double-stranded linker with a cohesive end, designed for ligation to restricted DNA, and a central region with a mismatch (Riley *et al.*, 1990). The primer is the same sequence as the mismatched part of the upper strand, and this prevents initiation of priming from the linker until the complementary strand has been synthesised by priming from within the retroviral insertion. However, this method suffers from non-specific annealing of the primers and 'end-repair' priming, in which the ends of unligated linkers initiate priming and enable PCR amplification without involving the retroviral-specific primer (see Devon *et al.*, 1995). Any errors that cause amplification of DNA that is not flanking an insertion will lead to the false identification of insertion sites.

An improved method uses splinkerettes, which incorporate a hairpin structure on the bottom strand, rather than a mismatch sequence (Devon *et al.*, 1995; Figure 1.10B). The primer has the same sequence as the upper strand and, as with vectorette PCR, cannot anneal until the complementary strand has been synthesised. The stable hairpin does not enable end-repair priming and only the upper strand can act as a non-specific primer. In all the PCR-based methods, insertions are only identified if target sites for the chosen restriction endonuclease are close enough to the insertion for the intervening region to be amplified. Coverage can be improved by using multiple restriction endonucleases (Uren *et al.*, 2005).

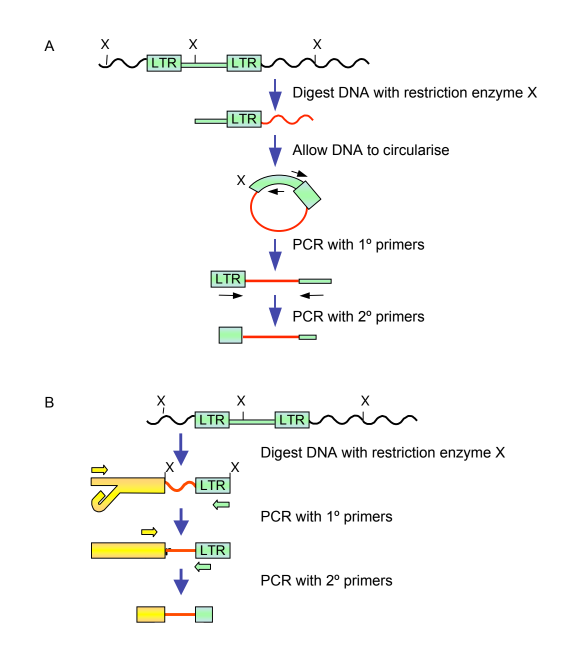


Figure 1.10 Isolation of retroviral insertion sites by inverse PCR (A) and splinkerette PCR (B). In inverse PCR, tumour DNA is digested using restriction enzyme X and the restricted DNA is allowed to circularise. Genomic DNA flanking retroviral insertions are amplified using PCR primers that bind within the insertion and point out towards the genomic DNA. A second round of PCR is performed using nested primers. The amplified DNA is sequenced and mapped to the mouse reference genome. Splinkerette PCR follows a similar procedure, except that instead of circularising the digested DNA, a splinkerette adapter (shown in yellow) is ligated to digested tumour DNA and genomic DNA flanking the retroviral insertions is amplified using PCR primers that bind to the adapter and the retroviral LTR.

Once the insertion-flanking genomic DNA has been amplified, the PCR products must be separated for sequencing. In the past, products were separated using agarose or polyacrylamide gels, but rare insertions are likely to be missed, and gel extraction is painstaking and subjective. An alternative method is to subclone the PCR products directly into a vector. By shotgun cloning the total mixture, it is possible to maintain the relative proportions of insertions from the starting material. However, it also means that more sequencing will be required to capture the rare insertions (see Uren et al., 2005). The VISA approach sequences PCR products directly, without subcloning, which reduces the risk of sequencing contaminating products (Weiser et al., 2007). The latest method parallel sequencing technology massively from 454 Life uses Sciences (http://www.454.com), in which fragmented genomic DNA is ligated to short adapters that are used for purification, amplification and sequencing. The DNA is denatured and immobilised onto beads, where PCR amplification and sequencing occur. This approach is extremely high-throughput, does not rely on cloning and is capable of detecting rare insertions. However, it can encounter problems when dealing with repetitive regions and long runs of a single nucleotide.

The next step is to map the sequenced DNA to the genome using a DNA alignment algorithm. For large screens, it is an advantage to be able to find high quality alignments quickly (Uren et al., 2005). The Sequence Search and Alignment by Hashing Algorithm (SSAHA2, Ning et al., 2001) converts the genome into a hash table, which can then be rapidly searched for matches. Sequences in the database (the mouse genome) are preprocessed into consecutive k-tuples of k contiguous bases and the hash table stores the position of each occurrence of each k-tuple. The query sequence (sequenced DNA) is also split into k-tuples and the locations of all occurrences of these sequences in the database, i.e. the "hits", are extracted from the hash table. The list of hits is sorted, and the algorithm searches for runs of hits in the database that match those in the query sequence. Having identified regions of high similarity, sequences are fully aligned using cross match (Green, unpublished), which is based on the Smith-Waterman-Gotoh alignment algorithm (Gotoh, 1982; Smith and Waterman, 1981). Because the database is hashed, search time in SSAHA2 is independent of database size, provided k is not too small. SSAHA2 is therefore three to four orders of magnitude faster than the BLAST alignment algorithm (Altschul et al., 1990), which scans the database and therefore performs at a speed that is directly related to database size (Ning et al., 2001).

48

As the PCR mixture is shotgun cloned and preferably sequenced to a high depth, an insertion site may be represented by more than 1 sequence read. Reads from a single tumour that map to the same genomic region must therefore be clustered into single insertion sites. Like the mutations in human cancer, tumour DNA will contain both insertions that drive oncogenesis (oncogenic insertions) and insertions that are passengers (background insertions). In theory, most identified insertions should be oncogenic because these, and particularly the earliest events in tumourigenesis, should be present in most, if not all, tumour cells, whereas background insertions that occur early in tumour development in a cell containing oncogenic insertions could also be highly represented in the final tumour (see de Ridder *et al.*, 2006).

Clustering of insertions from different tumours into common insertion sites (CISs) helps to distinguish oncogenic and background insertions. In theory, background insertions should be randomly distributed across the genome. Therefore, for small-scale screens, a gene in the vicinity of a cluster of insertion sites in different tumours is a strong candidate for a role in cancer. Methods for identifying statistically significant CISs, i.e. regions that are mutated by insertions in significantly more tumours than expected by chance, have involved generating a random distribution of insertions across the genome and obtaining an estimate of the number of false CISs in windows of fixed size using Monte Carlo simulation (Suzuki *et al.*, 2002) or the Poisson distribution (Mikkers *et al.*, 2002). These methods can be used to define the maximum window size in which insertions must fall to be considered non-randomly distributed. However, for larger scale screens, the window must be decreased to a size that is smaller than the spread of insertions within a single CIS so that many CIS are missed (de Ridder *et al.*, 2006). In addition, the above methods assume that insertions are randomly distributed and take no account of insertional biases, as mentioned in Section 1.4.2.1.1 (Wu *et al.*, 2006).

A more recent approach for CIS detection overcomes these problems by using a kernel convolution (KC)-based framework, which calculates a smoothed density distribution of inserts across the genome (de Ridder *et al.*, 2006). The scale (kernel size) can be varied so that CISs of varying widths can be identified. Decreasing the kernel size may identify separate CISs affecting the same gene, while increasing the kernel size will identify CISs where insertions are widely distributed in or around a gene. The method can be used for large-scale studies because it keeps control of the probability of detecting false CISs. The

threshold for significant CISs is based on the alpha-level defined by the user and on a null-distribution of insertion densities obtained by performing random permutations. A background distribution, such as the location of transcription start sites, can be provided to correct for insertional biases. See de Ridder *et al.* (2006).

The final step is to identify the genes that are being mutated by insertions within CISs, which are known in this thesis as "CIS genes". This may be relatively straightforward for intragenic insertions, but for enhancer mutations, which may have long distance effects, it is often difficult to identify the mutated gene unequivocally. Measuring the expression and transcript size of candidate genes in insertion-containing tumours can shed some light, but animal models and analysis of the orthologues in human cancer data are required for more conclusive evidence (Uren *et al.*, 2005).

A number of screens have been performed in recent years that have each identified hundreds of insertion sites (Hwang *et al.*, 2002; Johansson *et al.*, 2004; Li *et al.*, 1999; Lund *et al.*, 2002; Mikkers *et al.*, 2002; Slape *et al.*, 2007; Stewart *et al.*, 2007; Suzuki *et al.*, 2006; Suzuki *et al.*, 2002; Theodorou *et al.*, 2007; Uren *et al.*, 2008; Weiser *et al.*, 2007). The results of many screens have been collated and stored in the Retroviral Tagged Cancer Gene Database (RTCGD; http://rtcgd.abcc.ncifcrf.gov/) (Akagi *et al.*, 2004). At the time of writing, the database contains 503 CISs from 29 screens (database accessed May 2008). Users can search for individual genes of interest, or for CISs identified using particular mouse models and/or in particular tumour types. Genes with the most CISs are *Gfil* and *Myc*, with 82 and 77 insertions across all screens, respectively.

1.4.2.1.3 Identifying co-operating cancer genes

Retroviral insertional mutagenesis is a powerful tool for identifying genes that collaborate in tumour development. Collaborations can be identified by analysing the co-occurrence of CISs in individual tumours. For example, proviral activation of *Meis1* and *Hoxa7* or *Hoxa9* is strongly correlated in myeloid leukaemias from BXH2 mice (Bedigian *et al.*, 1984; Nakamura *et al.*, 1996). *Meis1* and *Hoxa9* are targets of translocation in human pre-B leukaemia (Kamps *et al.*, 1990) and acute myeloid leukaemia (AML) (Calvo *et al.*, 2002), respectively, and they are frequently co-expressed in human AML (Lawrence *et al.*, 1999). Both genes encode homeodomain transcription factors that bind to Pbx, and Meis1-Pbx and Hox-Pbx complexes have been shown to co-occupy the promoters of leukaemia-associated genes, such as *Flt3* (Wang *et al.*, 2006a).

A two-dimensional Gaussian Kernel Convolution method has recently been developed for identifying cooperating mutations in insertional mutagenesis data (de Ridder *et al.*, 2007). It is based on the kernel convolution framework used for identifying CISs (discussed in Section 1.4.2.1.2). The method has been applied to the data in RTCGD and, as well as finding previously characterised interactions, such as *Meis1* and *Hoxa9/Hoxa7*, it also finds novel interactions, such as *Rasgrp1* and *Cebpb*, which are both known to play a role in *Ras*-induced oncogenesis (de Ridder *et al.*, 2007).

As retroviral-induced tumours are oligoclonal, it is difficult to prove that tagged genes are in the same cell, and therefore that they collaborate (Largaespada, 2000). In an alternative approach, retroviral screens are performed on transgenic mice overexpressing known oncogenes, and knockout mice harbouring inactivated tumour suppressor genes, to identify genes that collaborate with the overexpression of oncogenes, and loss of tumour suppressor genes, respectively. For example, 35% of B-cell lymphomas generated in MuLV-infected $E\mu Myc$ transgenic mice, in which Myc is overexpressed in B-cell progenitors under the control of the immunoglobulin heavy chain enhancer, have an insertion in *Pim1* or the polycomb group protein *Bmi1* (van Lohuizen *et al.*, 1991). Bmi1 collaborates with Myc by inhibiting Cdkn2a (Ink4a/Arf), and therefore inhibiting Mycinduced apoptosis (Jacobs et al., 1999). In concurrence with these findings, Myc insertions were identified in 20% of tumours from MuLV-infected Cdkn2a-deficient mice, but none contained insertions in Bmil (Lund et al., 2002). Insertional mutagenesis also identifies genes that can functionally complement one another in tumour development. For example, in MuLV-infected EµMyc mice, activation of Pim2 increases from 15% to 80% in compound mutant mice lacking *Pim1* expression (van der Lugt et al., 1995), while Pim3 is selectively activated in mice lacking Pim1 and Pim2 expression (Mikkers et al., 2002). Pim1 is a coactivator of Myc that is required for expression of around 20% of all Myc target genes (Zippo et al., 2007). Pim kinases also appear to suppress Myc-induced apoptosis, but it is not clear whether this mechanism or Myc coactivation is responsible for the co-occurrence of *Pim1* and *Myc* mutations observed in lymphomagenesis (for review, see Naud and Eilers, 2007)). Piml also collaborates with Myc in human prostate cancers (Ellwood-Yen et al., 2003).

Retroviral screening of a mouse model for human myeloid leukaemia has identified 6 CIS genes, including *Plag1* and *Plagl2*, which co-operate with the oncogenic fusion gene *CBFB-MYH11* (Castilla *et al.*, 2004). This screen used a replication-defective retrovirus, cloned amphotropic virus 4070A, to limit the number of mutations and therefore to show that mutation of only one or a few genes was sufficient to induce tumorigenesis. Other studies using replication-competent viruses report 3-6 insertions in a single tumour (Mikkers *et al.*, 2002; Suzuki *et al.*, 2002) but, as mentioned above, retroviral-induced tumours are oligoclonal and it is therefore difficult to make a reliable estimate of the number of insertions in a tumour clone (see Neil and Cameron, 2002).

1.4.2.1.4 Generating tumours of different types

As discussed in Section 1.4.2.1.1, the dependence of retroviruses on cell-type-specific transcription factors limits the range of tumours that they can induce. There have been some successful attempts to alter the propensity of MuLV for T-cell lymphomas by using an $E\mu Myc$ transgenic mouse, which results in predominantly B-cell lymphomas (van Lohuizen *et al.*, 1991), and by expressing platelet derived growth factor B-chain $(PDGF\beta)$ from an MuLV-based retrovirus to generate mice with glioblastomas, which require activation of PDGF receptors for tumourigenesis (Johansson *et al.*, 2004). Mutations in the retroviral LTR may also lead to a change in tumour type, but manipulated viruses have a tendency to revert to wildtype (Uren *et al.*, 2005). In addition, MuLV and other retroviruses cannot infect nondividing cells, and infection is inefficient in slowly replicating cells and in tissues that have a basement membrane or mucin layer (Wang *et al.*, 2002a; Yamashita and Emerman, 2006). Transposon-mediated insertional mutagenesis is an alternative method that provides the possibility of generating a wider spectrum of tumours.

1.4.2.2 Transposon-mediated insertional mutagenesis

Like retroviruses, transposons are genetic elements that can mobilise within the genome. They are classified according to their mechanism of transposition. DNA transposons move by a "cut and paste" mechanism, in which they are excised from one site in the genome and integrated into another. Retrotransposons transpose via an RNA intermediate and are classified into LTR retrotransposons, which encode reverse transcriptase and transpose in a similar manner to retroviruses, and non-LTR retrotransposons, which are transcribed by host RNA polymerases and may or may not encode reverse transcriptase (Kapitonov and Jurka, 2008).

1.4.2.2.1 Sleeping Beauty

While DNA transposons are actively mobile in plants and invertebrates, all of the elements that have been so far identified in vertebrates are non-functional (Uren *et al.*, 2005). However, they can be mobilised in the mouse by using an invertebrate DNA transposon or by reconstructing a degenerate vertebrate transposon. *Sleeping Beauty* (SB) is a synthetic transposon derived from dormant DNA transposons of the Tc1/Mariner family in the genomes of salmonid fish. An active transposon, named SB10, was synthesised by directed mutagenesis on the basis of a consensus sequence obtained by aligning 12 degenerate transposon sequences from 8 species (Ivics *et al.*, 1997). SB consists of two inverted repeat/direct repeat (IR/DR) elements of ~230 bp each, flanking a cargo sequence (Collier *et al.*, 2005; Figure 1.11). Transposition occurs via binding of a transposase enzyme to two sites in each IR/DR (Izsvak *et al.*, 2000). All four binding sites are required for transposition and, in general, the closer the IR/DRs, the higher the transposition efficiency (Izsvak *et al.*, 2000). Higher levels of transposition have been achieved by introducing point mutations into the transposase, producing, for example, the SB11 (Geurts *et al.*, 2003) and SB12 (Zayed *et al.*, 2004) transposases.

The utility of SB for oncogenic insertional mutagenesis was first demonstrated in two studies published in 2005 (Collier *et al.*, 2005; Dupuy *et al.*, 2005). In both studies, transposons were introduced into mice by pronuclear injection of a linear plasmid containing one copy of the transposon, which forms a multicopy concatemer of variable length at a single site in the mouse genome. SB was mobilised by crossing these mice to mice expressing a transposase from a ubiquitous promoter. Collier and coworkers (2005) used a transgene containing the SB10 transposase under the control of the CAGGS promoter to mobilise around 25 T2/Onc transposons (Figure 1.11), while Dupuy *et al.* (2005) used the more active SB11 version knocked into the endogenous *Rosa26* locus to mobilise 150-350 copies of the T2/Onc2 transposon.

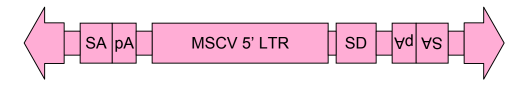


Figure 1.11. Structure of the *Sleeping Beauty* **transposon.** The presence of splice acceptors (SA) and polyadenylation signals (pA) in both orientations enables premature termination of gene transcription from intragenic insertions in both orientations. The transposon also contains the murine stem cell virus (MSCV) 5' LTR and a splice donor (SD) site that can induce promoter mutations in cellular genes. Elements for mutagenesis are flanked by 2 IR/DR elements, shown as arrows, which are required for transposon mobilisation. Redrawn and adapted from Figure 1a of Collier *et al.* (2005).

T2/Onc and T2/Onc2 were engineered to contain elements for mutagenesis much like those in retroviruses. The cargo of both transposons contains the 5' LTR of the murine stem cell virus (MSCV) followed by a splice donor, as well as splice acceptors followed by polyadenylation sites in both orientations. The transposons are therefore capable of disrupting genes by promoter mutation, N-terminal and C-terminal truncation and gene inactivation but, unlike retroviruses, they show low enhancer activity (Dupuy et al., 2005). T2/Onc and T2/Onc2 are essentially the same, except that T2/Onc2 contains a larger fragment of the Engrailed splice acceptor and the IR/DRs have been optimised for transposase binding (Dupuy et al., 2005). In the study by Dupuy and coworkers (2005), there was a high rate of embryonic lethality and, of the 24 T2/Onc2;Rosa26SB11 mice that survived to weaning, all developed cancer, most commonly T-cell lymphomas but also other haematopoietic malignancies plus a few cases of medulloblastomas and intestinal and pituitary neoplasias. Some mice had 2 or 3 types of cancer and all died within 17 weeks. In contrast, in the study by Collier et al. (2005), mice on a wildtype background did not develop tumours, but those on an Arf-null background developed sarcomas at an accelerated rate. The difference between the two studies most likely reflects the differences in transposon copy number and in transposase expression and activity (Collier and Largaespada, 2007). Transposase expression in CAGGS-SB10 mice has since been shown to be low and variegated in most tissues, probably due to epigenetic silencing of the transgene, while transposase expression is high in nearly all cell types in Rosa26SB11 mice (Collier and Largaespada, 2007). However, transposase is expressed in the testes of CAGGS-SB10 mice, which show high rates of transposition in the male germline (Collier and Largaespada, 2007; Dupuy et al., 2001).

Transposons, like retroviruses, can be used to identify co-operating cancer genes. For example, *Braf* was frequently mutated in *Arf*-null mice, suggesting that these genes co-operate in tumour formation (Collier *et al.*, 2005), while of the six T-cell tumours containing *Notch1* mutations, three also contained insertions mutating *Rasgrp1*, and 2 of these contains *Sox8* mutations, suggesting that these three genes also co-operate (Dupuy *et al.*, 2005).

While a number of the genes identified in the haematopoietic malignancies of T2/Onc2;Rosa26SB11 mice had been previously identified in retroviral mutagenesis, other genes had not (Dupuy *et al.*, 2005). This indicates that transposon-mediated mutagenesis is a complementary approach for cancer gene discovery, and may reflect

differences in insertional bias. While MuLV shows a strong preference for inserting near transcription start sites (Wu et al., 2003), SB shows a less pronounced preference and shows no preference for actively transcribed genes (Yant et al., 2005). SB inserts at TA dinucleotides and therefore shows a bias towards AT-rich sites, particularly those with the consensus sequence ANNTANNT (Carlson et al., 2003; Vigdal et al., 2002). However, most significant is the strong tendency of SB to transpose to sites close to the concatemer. This phenomenon, known as "local hopping", results in a non-random distribution of insertions that hampers CIS detection. Another potential hindrance to cancer gene identification is the ability of transposons to excise themselves and reinsert multiple times. SB leaves a small footprint upon excision, and it is possible that, at least in exons, this could continue to cause gene disruption that would not be identifiable (Collier and Largaespada, 2007). Likewise, the excision in some cells of transposons that had been critical for tumour development could result in a more heterogeneous tumour in which cancer gene identification would be more complicated. However, it is possible that such an event would be deleterious and that the cell would be eliminated (Collier and Largaespada, 2007) and, as SB transposition efficiency is higher for methylated (Yusa et al., 2004) and heterochromatic (Ikeda et al., 2007) transposons, excision of transposons involved in gene disruption may be relatively rare. A further drawback of SB, and possibly other DNA transposons, is that transposition induces genomic rearrangements, including deletions and inversions near to the transposon concatemer, and tumourigenesis could therefore be initiated by genes disrupted by these rearrangements rather than by mobilised transposons (Geurts et al., 2006).

One of the key benefits of using a transposon such as SB for insertional mutagenesis is that the mutagenic elements can be modified to control the types of mutation that occur. For example, modifying the cargo to enable only truncating mutations could increase the likelihood of identifying tumour suppressor genes (Collier and Largaespada, 2007). Tissue-specific promoters can be integrated as cargo, making transposons an attractive mutagen for cancer gene discovery in specific cancer types (Dupuy *et al.*, 2006). Spatial and temporal transposition could also be achieved by introducing a lox-stop-lox cassette between the SB transposase promoter and cDNA, such that transposition is induced upon the addition of Cre (Dupuy *et al.*, 2006).

Identification of cancer genes in SB mutagenesis follows much the same procedure as for retroviruses. Largaespada and Collier (2008) have developed a technique that uses

linker-mediated PCR, as described in Section 1.4.2.1.2, but that enables PCR amplification of DNA flanking both sides of the transposon to maximise coverage. Primers were designed to bind to the IR/DR sites and to synthetic adapters. Unlike in retroviral mutagenesis, tumour cells contain a concatemer of non-transposed elements. To avoid repeated cloning of the junctions between these elements, "blocking" primers can be used that bind to the plasmid DNA flanking each transposon in the concatemer but that have blocked 3' ends to prevent polymerase extension. Alternatively, after linker ligation, the DNA can be redigested with an endonuclease that cuts within the flanking plasmid DNA so that the primer binding sites are separated onto different molecules. (See Largaespada and Collier, 2008).

1.4.2.2.2 Alternative mutagens for transposon insertional mutagenesis

The active invertebrate transposons *piggyBac* and *Minos* are the only other DNA transposons that have so far been mobilised in the mouse (Collier and Largaespada, 2007). The *piggyBac* transposon, isolated from the cabbage looper moth, mobilises in mouse somatic cells and in the germline, and it can carry a larger cargo than SB (Ding *et al.*, 2005). The coding sequence of *piggyBac* has been codon-optimised to enable higher levels of transposition in the mouse, and inducible versions have been generated by fusing the transposon to the ERt² oestrogen receptor ligand-binding domain (Cadinanos and Bradley, 2007). Unlike SB, it shows a strong preference for inserting into genes in the mouse (Ding *et al.*, 2005) and in human cell lines (Wilson *et al.*, 2007). The *Minos* transposon, from *Drosophila hydei*, has attracted interest because it shows a low insertional bias and high transposition efficiency in a range of animals (for review, see Pavlopoulos *et al.*, 2007). However, it has so far shown only weak *in vivo* activity in the mouse (Drabek *et al.*, 2003; Zagoraiou *et al.*, 2001).

Retrotransposons are also gaining attention as potential insertional mutagens. Long interspersed nuclear elements (LINEs) are non-LTR retrotransposons that are transcribed into mRNA by RNA polymerase II and encode two proteins that are essential for transposition (Moran *et al.*, 1996): a protein that binds to single-stranded RNA (Hohjoh and Singer, 1997) and a protein with reverse transcriptase and endonuclease activity (Feng *et al.*, 1996; Mathias *et al.*, 1991). 17% of the human genome is composed of LINE-1 (L1) elements (Lander *et al.*, 2001). Transcription of endogenous L1 elements is generally inefficient but there are a small number of highly active "hot L1s", which were

used to generate a transgenic mouse model of L1 retrotransposition that showed a higher frequency of de novo somatic L1 insertions (Babushok et al., 2006). A 200-fold increase in transposition in the mouse germline has also been achieved by codon optimisation of the human L1 coding region (Han and Boeke, 2004). L1 mobilises by a "copy and paste" mechanism. It is therefore an attractive mutagen for forward genetic screens because, unlike DNA transposons, it is capable of self-expansion and the original insertion remains intact, aiding identification of mutated genes (Bestor, 2005; Collier and Largaespada, 2007). In addition, it appears to show no preference (An et al., 2006), or only a slight preference (Babushok et al., 2006), for inserting into genes and there is no local hopping because the RNA intermediate must exit and re-enter the nucleus before inserting into the genome. However, most L1 insertions are truncated at the 5' end (Babushok et al., 2006), potentially resulting in the loss of promoters, splice acceptors and polyadenylation signals required for mutagenesis (Collier and Largaespada, 2007). Controlled insertional mutagenesis using L1 derivatives has not yet been reported and *Sleeping Beauty* remains the preferred transposon for cancer gene discovery.

1.5 Cross-species comparative analysis for cancer gene discovery

Important biological sequences, such as gene coding regions and regulatory elements, are conserved in evolution. Cross-species comparative sequence analysis may therefore potentially help in the characterisation of known cancer genes. Comparison of intronic sequences in human and mouse BRCA1 led to the identification of two evolutionarily conserved regulatory elements in the second intron that, when mutated, had opposite effects on gene expression (Wardrop and Brown, 2005). However, cross-species comparative analysis also provides an extremely powerful approach for identifying novel genes and gene collaborations involved in cancer formation. As discussed in Section 1.3, the human cancer genome is highly complex. Many genes and pathways have been implicated in tumourigenesis, and most human cancers exhibit genomic instability, leading to the acquisition of genetic alterations that drive tumourigenesis but also many passenger mutations that do not contribute to the tumour phenotype. Distinguishing driver and passenger mutations is a major challenge. However, the molecular mechanisms that govern important biological processes are conserved in evolution, and cancer-associated mutation data from other species can therefore be used as a filter for identifying genes that represent strong candidates for a role in human cancer.

Genome-wide expression data for human tumours can be difficult to interpret, and a number of studies have therefore used cross-species comparative analysis to identify conserved expression signatures that are important in tumourigenesis. Expression profiles of intestinal polyps from patients with a germline mutation in *APC* were compared to those from *Apc*-deficient mice and the conserved signature showed an overrepresentation of genes involved in cell proliferation and activation of the Wnt/ β -catenin signalling pathway (Gaspar *et al.*, 2008). Likewise, comparison of expression profiles for human lung adenocarcinoma and a mouse model of *Kras2*-mediated lung cancer led to the identification of a *KRAS2* expression signature that was not identified by analysing *KRAS2*-mutated human tumours alone (Sweet-Cordero *et al.*, 2005). More recently, a mutated *Kras*-specific signature that can be used to classify human and mouse lung tumours on the basis of their *KRAS* mutation status has been identified by comparing *KRAS2*-mutated human cancer cells to mouse somatic cells containing knocked-in mutant *Kras* (Arena *et al.*, 2007).

Mouse prostate cancers induced by human *MYC* have an expression signature that defines a set of "*Myc*-like" human prostate tumours and includes overexpression of the oncogene *Pim1* (Ellwood-Yen *et al.*, 2003). Rat prostate tumours also have a similar expression profile to human prostate tumours, and have been used to identify conserved genes that are differentially expressed in both species in response to treatment with the chemopreventive agent Selenium (Schlicht *et al.*, 2004). The mouse is therefore not the only cancer model that has been used for cross-species comparison. The greater the evolutionary distance between the species, the greater the likelihood that conserved changes in gene expression contribute to the cancer phenotype. An expression signature in zebrafish liver tumours is more consistently associated with human liver tumours than with other human tumour types and, since human and zebrafish are distantly related, genes in the conserved signature are strong candidates for a role in cancer development (Lam *et al.*, 2006).

Another approach for cross-species analysis involves comparing the CGH profiles of human tumours to the CGH profiles of tumours generated from a mouse model of the corresponding human cancer. Such studies take advantage of the conserved synteny between the human and mouse genomes (Waterston *et al.*, 2002). Comparison of CGH profiles for human neuroblastomas with profiles for tumours and cell lines from a *MYCN* transgenic mouse model of neuroblastoma have shown that many genetic aberrations are

conserved between species (Cheng et al., 2007; Hackett et al., 2003). Likewise, 80% of aberrations detected by array CGH in tumour cells of the mouse model for epithelial ovarian cancer are conserved in human epithelial ovarian cancer (Urzua et al., 2005), and epithelial carcinomas in mice with telomere dysfunction show numerous copy number changes in regions syntenic to those in human cancers (O'Hagan et al., 2002). Zender and coworkers (2006) used array CGH to identify regions of copy number change in the tumours of a mouse model for hepatocellular carcinoma. The CGH profiles were compared to array CGH data for human hepatocellular carcinomas to identify minimally conserved amplicons, and genes that showed increased expression in both species were chosen as candidate cancer genes. The authors identified 2 oncogenes, cIAP1 and Yap, that act synergistically in a focal amplicon on mouse chromosome 9qA1, which is syntenic to an 11q22 amplicon in human tumours. Kim *et al.* (2006b) used a comparable approach to identify *Nedd9* as a candidate for a role in promoting melanoma metastasis. A focal amplicon comprising 8 genes, including Nedd9, was identified on chromosome 13 in 2 metastatic cell lines derived from a *Ras* mouse model of nonmetastic melanoma. 36% of metastatic melanomas contained a much larger amplicon in a syntenic region on human chromosome 6p25-24, and 35-52% of metastatic melanomas showed significant overexpression of NEDD9, with more advanced tumours showing higher levels.

Comparison of human cancers with mouse models of cancer relies on the use of mouse models that accurately recapitulate the human cancer (Tomlins and Chinnaiyan, 2006). While *cIAP1* and *Yap* overexpression was found to be important in *p53^{-/-};Myc*-induced hepatoblasts in the study by Zender et al. (2006), neither gene contributed to tumourigenesis in p53^{-/-}; Akt or Ras hepatoblasts. Likewise, Nedd9 did not contribute to melanoma metastasis in the absence of *Ras* or *Raf* activation (Kim *et al.*, 2006b). Crossspecies comparison of genomic profiles for a particular cancer may therefore require some prior knowledge of the genetic events that drive tumourigenesis in that cancer so that an appropriate mouse model can be generated. However, cross-species analysis can also facilitate the selection of a suitable mouse model. Lee and coworkers (2004) used unsupervised hierarchical clustering of expression data from human and mouse hepatocellular carcinomas to identify the mouse models that provided the best fit for human cancers. Mouse and human tumours that clustered together due to similar expression profiles also shared phenotypic characteristics, such as proliferation rate and prognosis (Lee *et al.*, 2004). Most genetically engineered mouse models do not show the high levels of chromosome instability associated with human cancers. Mice that are

engineered with telomere dysfunction, or defects in DNA damage checkpoints or DNA repair, may therefore represent better models for comparative oncogenomics (Maser *et al.*, 2007). Comparative analysis of copy number alterations in chromosomally unstable murine T-cell lymphomas and human solid tumours identified recurrent aberrations in the mouse that are conserved in human T-cell acute lymphoblastic leukaemias but also in other human tumour types (Maser *et al.*, 2007).

Candidate cancer genes can also be identified by comparing expression and CGH profiles for human tumours with mouse insertional mutagenesis screens. Genes in expression signatures associated with distinct subclasses of human acute myeloid leukaemia were significantly correlated with genes nearest to insertion sites in a Graffi 1.4 MuLV mouse model and with candidate leukaemia genes in BXH2 and AKXD mouse models (Erkeland et al., 2006). There was little overlap between the candidates identified by Graffi 1.4 and BXH2/AKXD, demonstrating that retroviral screens involving multiple models and viruses may be required for a more effective cross-species comparison (Touw and Erkeland, 2007). Amplified regions in human pancreatic cancer have also been shown to contain more CIS in retrovirus-induced murine lymphomas and leukaemias than expected by chance (Aguirre et al., 2004). As discussed in Section 1.4, insertional mutagenesis "tags" the mutated gene, therefore facilitating cancer gene identification. In contrast, copy number alterations in human cancer can be very large, encompassing many genes, and no systematic approach currently exists for identifying the critical genes within these regions (Degenhardt et al., 2008). Thus comparative analysis of oncogenic insertions in mouse tumours and CGH data for human tumours is potentially a very powerful approach for narrowing down the candidates in regions of copy number change.

1.6 Aims of this thesis

The elucidation of the human genome sequence and the advent of high-throughput technologies for characterising cancer genomes have led to the discovery that the cancer genome is far more complex than previously thought. Genome-wide, cancer-associated mutation datasets can be generated at increasing speed and diminishing cost, yet identifying the mutations that contribute to the cancer phenotype remains a challenge. Integrative analyses, particularly cross-species comparisons, provide a means of distinguishing likely driver mutations from the background of passenger mutations that arise in unstable cancer genomes. The identification of cancer genes in regions of copy

number change is especially problematic since such regions are often large and encompass many potential candidates. Forward genetic screens are purported to be a powerful tool for cancer gene discovery in the mouse, but how relevant are they to human cancer?

This thesis describes work undertaken to compare large-scale datasets generated by mouse insertional mutagenesis and CGH analysis of human cancer cell lines. The main aims of this project are to narrow down the candidate cancer genes in regions of copy number change in human cancers and, in so doing, demonstrate the utility of forward genetic screens in the mouse for the identification of human cancer genes. Chapter 2 describes the steps taken to identify mouse candidate cancer genes from a retroviral insertional mutagenesis dataset generated from 1,005 mouse tumours and a smaller transposon-mediated insertional mutagenesis dataset generated from 73 mouse tumours. Chapter 3 describes detailed analyses of the mouse candidate genes, including comparisons with numerous human and mouse cancer-associated mutation datasets, as well as an analysis of the types of mutations occurring in each candidate and the identification of collaborating cancer genes. Chapter 4 describes the work undertaken to identify regions of copy number change in Affymetrix 10K SNP array CGH data for 713 human cancer cell lines, and then to identify candidate cancer genes within these regions by comparison with mouse candidates from the retroviral screen. In Chapter 5, higher resolution Affymetrix SNP 6.0 CGH data generated from a subset of the same cell lines is used, again to identify putative cancer genes, but also for comparison with the lower resolution data to demonstrate the superiority of the high-resolution data for cancer gene discovery. Analyses that attempt to identify genes that co-occur, and therefore potentially co-operate, in both human and mouse cancers are also described. Finally, conclusions drawn from the analyses are presented in Chapter 6.

Chapter 2 Identifying insertion sites and candidate cancer genes by insertional mutagenesis in the mouse

2.1 Introduction

When a retroviral or transposon insertional mutagen inserts into the mouse genome, it acts as a molecular tag that facilitates the identification of genes that it disrupts. As discussed in Section 1.4.2.1.2, the elucidation of the mouse genome sequence and the development of high-throughput, PCR-based technologies for insertion site identification have allowed for larger scale mutagenesis screens that can identify a higher proportion of insertions across larger numbers of tumours. 1,005 mouse tumours were generated in a retroviral insertional mutagenesis screen performed by the Netherlands Cancer Institute (NKI). Murine leukaemia virus (MuLV) was used as the insertional mutagen, and insertions into the mouse genome were identified using splinkerette PCR (see Section 1.4.2.1.2). In a separate study at the University of Minnesota, 73 mouse tumours were generated by insertional mutagenesis using the Sleeping Beauty T2/Onc transposon (see Section 1.4.2.2.1). Genomic DNA flanking the retroviral and transposon insertion sites was sequenced at the Wellcome Trust Sanger Institute. This chapter begins with a description of the retroviral and transposon insertional mutagenesis datasets. While I did not contribute to the generation of tumours or sequence reads, all statistics are the result of my own analyses. A dataset of known cancer genes, compiled by the Sanger Institute Cancer Genome Project, is also described. This is followed by an account of the work undertaken to process the sequence reads into insertion sites, to filter out erroneous reads and insertion sites, and to measure the coverage of the screen. A relatively high proportion of reads could not be mapped, and the nature of non-mapping reads was therefore investigated. The remainder of the chapter focuses on the methods used to identify candidate cancer genes in the vicinity of mapped insertions. The identification of genes that are being mutated by retroviral insertions is complicated by the presence of enhancer mutations that may act at long range (see Section 1.4.2.1.2). Insertions were assigned to genes by defining rules based on an analysis of the distribution of insertions around mouse genes. Statistically significant common insertion sites (CISs) were defined using Monte Carlo simulations (Suzuki et al., 2002) and a kernel convolution-based

framework (de Ridder *et al.*, 2006), and CIS genes identified by the two approaches were compared. Data from the retroviral screen forms the principal mouse dataset used in this thesis, and is therefore discussed in greater detail than data from the transposon screen. The main steps involved in identifying candidate cancer genes from retroviral sequence reads are summarised in Figure 2.1. Unless otherwise stated, *P*-values provided in this chapter were generated using the Chi-squared test for independence.

2.2 Description of the datasets

2.2.1 The retroviral dataset

Mice of the *FVB* strain were engineered with a range of genetic backgrounds in order to identify cancer genes that collaborate with the loss of tumour suppressor genes (see Section 1.4.2.1.3). 1,005 tumours were generated, of which 22.7%, 12.5% and 23.0% were on a $p19^{4RF-/-}$ (*Cdkn2a^{-/-}*), $p53^{-/-}$ or wildtype genetic background, respectively. The remaining tumours were generated on a background deficient in *p15*, *p16*, *p21* or *p27*, or a combination of these (Table 2.1A). Equal numbers of males and females were used (500 each of males and females, 1 hermaphrodite and 4 unknown). The vast majority (at least 90.9%) of tumours originated in the spleen, thymus or lymph nodes (Table 2.1B). The 1-tailed Fisher Exact Test was performed to determine whether genetic background or gender was associated with particular tumour types. Wildtype and $p19^{-/-}$ genetic backgrounds were over-represented in tumours of the thymus $(P=4.67 \times 10^{-6} \text{ and }$ $P=9.87 \times 10^{-5}$, respectively), while among tumours of the spleen, there was an overrepresentation of $p53^{-/-}$ (P=5.05x10⁻⁵) as well as wildtype and $p19^{-/-}$ genetic backgrounds $(P=0.0240, \text{ and } P=1.84 \times 10^{-4}, \text{ respectively})$. Lymph node tumours were over-represented in $p16^{-/2}p19^{-/2}$ mice (P=1.10x10⁻⁵) and in mice with a deficiency in p21 or p21 and p27 $(p21^{-/-}, P=2.32x10^{-13}; p21^{-/-}p27^{+/-}, P=4.34x10^{-4}; p21^{-/-}p27^{-/-}, P=2.17x10^{-4}; p21^{+/-}p27^{+/-}, P=4.34x10^{-4}; p21^{-/-}p27^{+/-}, P=2.17x10^{-4}; p21^{-/-}p27^{+/-}, P=4.34x10^{-4}; p21^{-/-}p27^{-/-}, P=2.17x10^{-4}; p21^{-/-}p27^{+/-}, P=4.34x10^{-4}; p21^{-/-}p27^{+/-}; P=4.34x10^{-4}; P=4.34x10^{-4};$ P=0.0128). It is possible that these results represent a subjective bias in the selection of Alternatively, they may indicate that different genetic backgrounds are tumours. predisposed to different tumour types. Most striking was the over-representation of the $p16^{-/2}p19^{-/2}$ genotype among tumours in the liver ($P=1.04 \times 10^{-32}$). At least 24 of the 33 liver tumours have been identified as tumours of the liver nodule. These are commonly observed in $p16^{-/2}p19^{-/2}$ mice infected with MuLV and may be lymphomas that have spread to the liver or they may be histiocytic sarcomas, which are a poorly-defined class of haematopoietic neoplasm (Lund et al., 2002). There was no significant difference

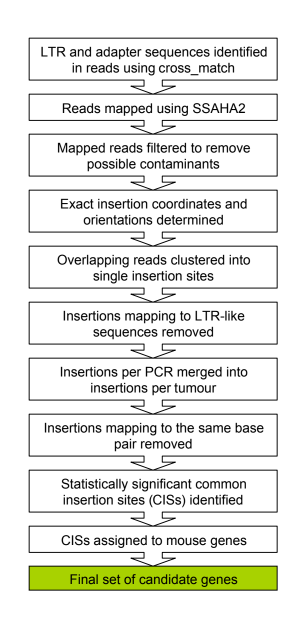


Figure 2.1. Workflow for identifying mouse candidate cancer genes from sequencing reads generated in a retroviral insertional mutagenesis screen.

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	Number of
Genotype	tumours
wildtype	231
p19-/-	228
p53-/-	126
p16-/-, p19-/-	91
p15-/-	55
p21-/-, p27+/-	54
p21-/-	43
p27+/-	38
p21-/-, p27-/-	36
p27-/-	36
p16+/-, p19+/-	26
p21+/-, p27+/-	17
p15-/-, p21-/-	15
p21+/-, p27-/-	5
p53+/-	2
p21+/-	2
Total	1005

	Number of
Tissue	tumours
spleen	468
thymus	227
lymph node	125
spleen; lymph node	71
unknown	52
liver	33
thymus; spleen	15
spleen nodule	4
spleen; liver	3
kidney nodule	2
scapular tumour	1
uterine tract	1
uterine tumour	1
fascial lymphoma	1
uterine tumour; lymph node	1
Total	1005

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	Number of
Tissue	tumours
spleen	38
thymus	22
lymph node	10
brain tumour	2
unknown	1
Total	73

Table 2.1. Characterisation of the insertional mutagenesis datasets. (A) The number of tumours from mice with different genetic backgrounds in the MuLV screen. (B) The number of tumours of each tissue type in the MuLV screen. (C) The number of tumours of each tissue type in the *Sleeping Beauty* T2/Onc screen.

В

between the number of males and females with tumours from different tissues or genetic backgrounds.

Following the isolation of tumour DNA, most samples were subjected to two separate splinkerette PCRs using different restriction enzymes, Sau3AI and Tsp509I, in order to increase the number of insertions that could be identified in the screen (see Section 1.4.2.1.2). The PCR products were shotgun cloned, and 96 reads were sequenced per PCR. Everything described from this point onwards is the result of my own work. The reads were converted to a CAF (Common Assembly Format) file, which contains the DNA sequence, base quality, and the coordinates of sequencing and cloning vector sequences within the read. The CAF file was then converted to FASTA format, in which the vector sequences were masked. The resulting dataset comprised 159,303 sequence reads from 2,060 PCRs. 14,767 reads from 199 PCRs were discarded because they were of unknown identity or had been flagged as invalid due to possible sample mix-up, no obvious tumour when killed, or contaminated or low quality PCR. The remaining 144,536 reads included 134,985 that were generated from 1,734 PCRs performed on 1,005 mouse tumours. For 62% of tumours, the dataset contained reads obtained from 2 PCR experiments, i.e. using both restriction enzymes, while for 33% of tumours, reads were only available for a single experiment. The remaining 5% of tumours were subjected to 3 or 4 PCRs, in which additional reactions using Sau3AI and/or Tsp509I were performed. The number of reads per tumour is shown in Figure 2.2. To facilitate the identification of PCR artefacts, 1,180 reads were also generated from 24 PCRs performed on uninfected mice. Finally, 8,371 reads were generated from 103 PCRs performed on samples that were harvested from mice 5 or 10 days post-MuLV infection. There has been limited time for cell re-infection, and thus for tumour initiation and progression, in these "short infection time" mice. A high proportion of insertions in samples from these mice are therefore expected to map to sites in the genome where the virus prefers to insert ("hotspots") and that may not contribute to tumourigenesis.

Cross_match (Green, unpublished) was used to identify and mask the retroviral LTR (5'-GCTAGCTTGCCAAACCTACAGGTGGGGTCTTTCA-3') and splinkerette adapter (5'-CCACTAGTGTCGACACCAGTCTCATTCAGCCAC-3') in order to prevent erroneous mapping of reads to regions of the mouse genome that resemble these sequences. The minimum length of the perfectly matching sequence (minmatch) and the minimum alignment score (minscore) were each set to 10. These parameters were used

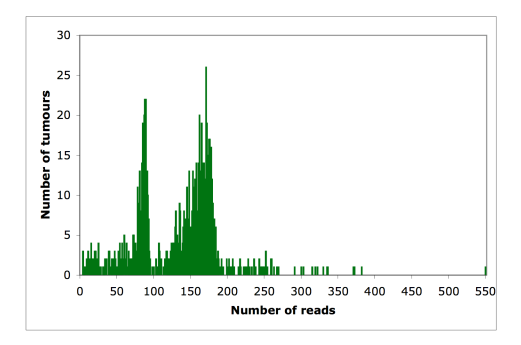


Figure 2.2. The number of sequence reads per tumour before mapping. Up to 96 reads were sequenced for each PCR. The bimodal distribution reflects the fact that 62% of tumours were subjected to 2 linker-mediated PCRs, while for 33% and 5% of tumours, 1 PCR or more than 2 PCRs, respectively, were performed.

for all cross_match runs, unless otherwise specified. Among the reads from tumour DNA, 110,318 (81.7%) contained LTR and adapter sequences, 9,534 (7.1%) contained an LTR but no adapter, 12,592 (9.3%) contained an adapter but no LTR, and 2,541 (1.9%) contained neither.

2.2.2 The Sleeping Beauty dataset

This smaller screen comprised 73 tumours, of which 60 were from wildtype mice on a mixed *C57BL/6J/FVB* background, and 13 were from *Bloom (Blm)*-deficient mice from the same strain. *Blm*-deficient tumours may be more likely to harbour mutations that inactivate tumour suppressor genes (see Section 1.4.2.1.1). These tumours, and 31 wildtype tumours, were generated from a transposon array (LC76) located on chromosome 1. The remaining wildtype tumours were generated from an array (LC68) on chromosome 15. As in the MuLV screen, tumours developed almost exclusively in the spleen, thymus and lymph node (Table 2.1C) since mice have a propensity for these tumour types.

Insertions were cloned using linker-mediated PCR in which genomic DNA flanking both sides of the insertion was amplified to maximise insertion site identification (see Section 1.4.2.2.1). The restriction enzymes *BfaI* and *NlaIII* were used to clone DNA flanking the 5' and 3' IR/DRs, respectively. As in the retroviral screen, PCR products were shotgun cloned and 96 reads were sequenced. All work described hereafter is my own. The initial dataset comprised 16,674 sequences. Although steps were taken to minimise the amplification of transposons within the concatemer (see Section 1.4.2.2.1), the sequence data inevitably contain some reads that map to the concatemer. Transposons in the concatemer are flanked by the sequence 5'-TATAGGGATCC-3' and therefore any reads containing this sequence are likely to represent transposons that have not mobilised. 89 concatemer sequences were removed using cross match (Green, unpublished).

The presence of the transposon IR/DR provides evidence that the genomic DNA is directly flanking an insertion. Using cross_match, IR/DR elements were identified and masked in 15,630 reads (94.2% of the total), and the rest were discarded. The linker, which was identified in 12,209 (78.1%) of the remaining reads, and extra vector sequence from the PROMEGA pGEM-T easy vector T7 promoter-multiple cloning site-SP6 promoter were also screened out with cross_match. 3,716 reads (23.8%) contained fewer

than 25 bp of unmasked sequence after screening, and as these would be too short for mapping, they were removed from the dataset. Tumour details were not available for a further 1,123 reads, and so these were also removed. The final dataset comprised 10,791 reads generated from 138 PCRs. This included 60 tumours for which genomic DNA flanking both sides had been amplified and sequenced, and 11 tumours for which only one side had been amplified. For the remaining 2 tumours, both sides had been amplified, and PCR had been performed twice on one or both sides.

2.2.3 Known cancer genes in the Cancer Gene Census

The Cancer Gene Census is a list of genes for which there is strong evidence of a role in cancer (Futreal et al., 2004; see Section 1.2.5.2). The complete working list dated 13/02/2007 was downloaded from http://www.sanger.ac.uk/genetics/CGP/Census/. The Ensembl (Hubbard et al., 2007) Perl Application Programming Interface (API) was used to extract the Ensembl identifiers for each gene in the list from Ensembl version 48. Ensembl provides annotation on a selection of eukaryotic genomes, and it has been used throughout this project to obtain information about the mouse and human genomes. The API provides standardised methods for accessing data in the Ensembl MySQL databases through Perl scripts and it insulates developers from changes at the database level. From the 363 genes in the Cancer Gene Census, 354 human Ensembl genes were identified. 352 mouse Ensembl genes have a human orthologue in the Cancer Gene Census. 314 mouse genes have an orthologue with somatic mutations in cancer and 67 have an orthologue with germline mutations, including 32 that have an orthologue with both mutation types. The orthologues of 285 mouse genes bear mutations that are dominant at the cellular level, 66 bear recessive mutations, of which 2 are X-linked, and 1 has both dominant and recessive mutations. 205 have been implicated in leukaemia and/or lymphoma, 102 have been implicated in epithelial tumourigenesis and 84 have been implicated in mesenchymal tumourigenesis. The most common type of mutation is translocation, which affects the orthologues of 263 mouse genes. A list of the human cancer genes with mouse orthologues is provided in Appendix A.

2.3 Mapping the sequence reads using SSAHA2

As discussed in Section 1.4.2.1.2, SSAHA2 (Ning *et al.*, 2001) is a fast DNA alignment algorithm that is suited to mapping large numbers of insertions to the mouse genome.

The parameters of SSAHA2 were adjusted to maximise the number of mapped reads, and therefore to identify as many insertions as possible. A test set of 25,000 reads from the retroviral screen was mapped to the NCBI m34 mouse genome assembly. SSAHA2 preprocesses the query sequence (the read) and the subject (sequences in the NCBI m34 database) into consecutive k-tuples of k contiguous bases, called the word size or k-mer. Lowering the k-mer increases the sensitivity, and therefore yields more hits, but it also increases CPU time, and a k-mer of 13 or 14 is generally recommended for large databases, such as genome assemblies. The default k-mer of 12 was used for all runs of SSAHA2, since this offers a small gain in sensitivity without impacting too heavily on the speed. The "seeds" parameter defines the number of exact words that must match in the subject. Lowering the seeds increases the sensitivity, resulting in a higher proportion of low (<95%) identity and ambiguous mappings, but also more high identity unambiguous mappings (Table 2.2A). Initially, seeds 3 was chosen because seeds 2 yielded only 8 additional high identity unambiguous mappings and required more CPU time. Bv default, sequences are processed into consecutive k-mers with no overlap. Reducing the parameter "skip" increases the overlap between k-mers and should provide greater sensitivity. For seeds 3, decreasing skip to 4 (8 base overlap) and 6 (6 base overlap) did not increase numbers of high identity, single mapping reads. For higher seeds, numbers did increase but were lower than for seeds 3 alone (Table 2.2B). SSAHA2 with seeds 3 vielded more mappings than NCBI BLASTN (Altschul et al., 1990; Table 2.2A) and was significantly faster. BLASTN parameters were set for moderately sized (~500 bp) genomic DNA (-G 1, -E 3, -W 30, -F 'm D', -U, -e 1e-20).

The full set of 144,536 retroviral reads was mapped to the NCBI m36 mouse build using SSAHA2 with seeds 3 and default values for all other parameters. Alignments with low identity were not segregated in this larger analysis because they may simply represent sequencing reads of poor quality and, if they are erroneous, they should be picked up in the filtering process (see Section 2.5). 86,290 reads (59.7%) mapped to a single location, 28,484 (19.7%) mapped to multiple locations, and 29,762 (20.6%) did not map at all. Further runs of SSAHA2 were performed with lower seeds to map as many of the unmapped reads as possible. 3,866 (13.0%) of unmapped reads could be mapped using seeds 2, and the same results were obtained with seeds 1. This is surprising, since the difference between seeds 3 and 2 was minimal when the 25,000-read test dataset was used. In the test set, analysis with seeds 2 did increase the number of alignments with <95% identity (Table 2.2A), and it is therefore likely that a proportion of the additional

А

SSAHA2 seeds				
5	4	3	2	BLAST
13470	13894	14158	14164	14010
7060	6002	4971	3870	5960
1837	2110	2414	3253	1365
2633	2994	3457	3713	3665
25000	25000	25000	25000	25000
	5 13470 7060 1837 2633	541347013894706060021837211026332994	543134701389414158706060024971183721102414263329943457	543213470138941415814164706060024971387018372110241432532633299434573713

В

	seeds 3			9	seeds 5	
Mapping	default	skip 4	skip 6	default	skip 4	skip 6
Single	14158	13699	13854	13470	13875	14004
None	4971	4187	4044	7060	3959	5096
Low	2414	3301	3281	1837	3301	2413
Multiple	3457	3813	3821	2633	3865	3487
Total	25000	25000	25000	25000	25000	25000

Table 2.2. The number of MuLV reads mapped using SSAHA2, with varying values for parameters seeds and skip, and BLASTN. (A) Lowering the number of seeds increases the number of reads mapped by SSAHA2. (B) Increasing the overlap between *k*-mers decreases the number of reads mapped using seeds 3 but increases the number mapped using seeds 5. Mapping types are Single (read maps to a single location in the genome), None (read unmapped), Low (read maps with an identity lower than 95%) and Multiple (read maps to multiple locations in the genome).

unambiguous mappings obtained using SSAHA2 with seeds 2 on the entire dataset have a low identity. The difference may also reflect developments in the algorithm in the time between the two analyses. The default minimum Smith-Waterman score is 30, and reducing this to 20 further increased the number of unmapped reads that could be mapped to a single location using seeds 2 to 4,382 (14.7%). The final set of mappings comprised 90,672 reads (62.7%) that mapped unambiguously and 29,769 (20.6%) that mapped to multiple locations. 24,095 (16.7%) remained unmapped.

Based on the observations for the retroviral dataset, the 10,791 reads of the *Sleeping Beauty* dataset were mapped to NCBI m36 using SSAHA2 with default parameters plus seeds 2 and score 20. 5,470 (50.7%) mapped to a single genomic location, 1,859 (17.2%) mapped to multiple locations, and 3,462 (32.1%) did not map at all.

2.4 Accounting for unmapped reads

Even after maximising the number of reads that could be mapped using SSAHA2, there was still a high proportion of unmapped reads in both the retroviral and *Sleeping Beauty* datasets. The lengths of the 96,072 single-mapping, and 24,095 non-mapping, retroviral reads are shown in Figures 2.3A and 2.3B, respectively. Since it is not known which part of the read, if any, is genomic DNA, all bases that were not masked as vector, LTR or linker were counted. 2,143 (8.9%) of the unmapped reads were exactly 132 base pairs in length and a high proportion of these shared an identical sequence flanked by LTR and splinkerette sequences. One read of length 132 bp was submitted to SSAHA2 and BLASTN on the Ensembl website (http://www.ensembl.org/). As expected, there were no matches to NCBI m36 using SSAHA2 with near exact or no optimisation. Using BLASTN optimised for near exact matches (-E 10 -B 100 -filter dust -RepeatMasker -W 15 -M 1 -N -3 -Q 3 -R 3), there were 96 hits, all of which were low scoring. The hit with the lowest E-value and P-value (both 4.2×10^{-7}) was an alignment of 50 bp with a score of 22 and 86% identity to chromosome 8:126312491-126312540. The sequence was also submitted to the Ensembl Trace Server (http://trace.ensembl.org), which contains millions of single-pass DNA sequencing reads from over 1,000 different species. The full length of the read matched with 100% identity to 6 clones from the free-living nematode species Pristionchus pacificus. Since it was unclear how DNA from this organism would have become incorporated into the screen, a 132 bp read was also submitted to NCBI VecScreen (http://www.ncbi.nlm.nih.gov/VecScreen/), which

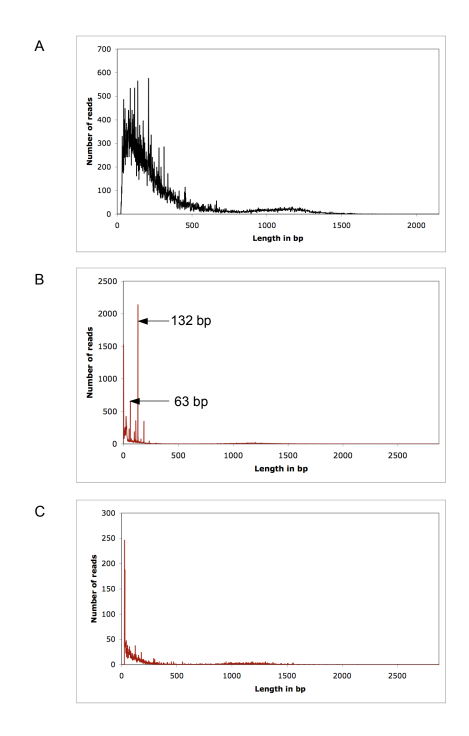


Figure 2.3. The lengths of retroviral reads that are unambiguously mapped (A), unmapped (B), and unmapped and uncharacterised (C). The reads of length 63 bp and 132 bp, which underwent further investigation, are shown.

searches for vector contamination in nucleic acid sequences by using BLAST to query the UniVec database. The entire sequence aligned to the MuLV retroviral vector pLNL6 with 100% identity and no gaps. Therefore, it appears that the 132 bp sequences are composed entirely of retroviral sequence, to which an adapter has been ligated.

There were 657 sequences of length 63 bp. One such sequence was submitted to BLASTN optimised for near exact matches, and one hit – an alignment of 18 bp with 100% identity to chromosome 15:90360616-90473373 – was obtained. The highest scoring hit obtained in a search against the Trace Server was just 75.9% identity, to a sequence from an unknown source. A VecScreen search revealed a 100% identity match along the entire length of the unmasked sequence to the cloning vector pBR322. Since these reads contain an adapter sequence, it is likely that they represent contamination during linker-mediated PCR.

Other reads containing the pLNL6 and pBR322 vector sequences were identified using cross_match. 19.4% of unmapped reads contained the pLNL6 sequence, while a further 4.8% contained the pBR322 sequence. In contrast, only 0.32% and 0.08% of reads mapping to a single location had matches to pLNL6 and pBR322, respectively. RepeatMasker (Smit *et al.*, 1996-2004) was used to identify repeat regions within the remaining unmapped reads. 13.6% contained low-complexity regions. Such regions are difficult to sequence, and these reads may have failed to map because they were not correctly sequenced. Alternatively, low-complexity regions may be the result of polymerase stuttering, where the polymerase transcribes the same nucleotide multiple times during PCR amplification, and the read may therefore no longer bear a close enough resemblance to the corresponding region of the mouse genome. The proportion of low-complexity regions was significantly lower in reads that mapped unambiguously (6.2%, P=0).

A further 26.6% of unmapped retroviral reads were below the minimum length (25 bp) that could be mapped using SSAHA2 with the chosen parameter values. Of the remaining sequences, 0.73% comprised more than 50% Ns (i.e. unknown nucleotides), and 1.2% contained other types of repeat element identified by RepeatMasker. This compared with 0.11% and 0.18%, respectively, for reads mapping to a single location (P=0 for both). The 2-tailed Fisher Exact test was used to determine whether there was any significant difference between the numbers of each type of repeat identified by

75

RepeatMasker in the mapped and unmapped reads. All types of low-complexity region were over-represented in the unmapped reads, as were simple repeats and a selection of retrovirus-related repeat elements (HAL1, GSAT_MM, L1MEd, LTR/ERV1, MuLV-int, MuRRS-int, RLTR4_MM-int and RLTR6-int, see Table 2.3A). This supports the theory that many of the reads could not be mapped either because they contain low complexity regions, or because they contain retroviral sequence and may not contain any genomic DNA. There were also numerous under-represented repeat elements among the unmapped reads (Table 2.3B). These included elements that one would expect to find in genomic DNA, such as 4.5SRNA and LINEs and SINEs, and elements that are specific to the genomes of rodents, such as the endogenous LTR MTE2a, and to the mouse in particular, such as the SINE B2 Mm2.

In summary, 15,996 (66.4%) of unmapped reads contained vector sequences or sequences of low complexity, low quality or short length (Table 2.4). The remaining 8,099 reads were searched against the Ensembl Trace Server using SSAHA2 with seeds 5. 5.1% had matches in the archive, of which 90.5% had matches to sequences of mouse origin. All of the non-mouse matches had an identity of less than 91%, except one, which matched with 100% identity to 2 sequences, with trace names rtn1ut06.g and rtn1yp83.g, from *Rattus norvegicus*. As rat and mouse are closely related, it is possible that this read does contain DNA from the mouse genome, but that it does not align to the mouse genome because of a genome assembly error. 245 reads mapped to mouse sequences in the Ensembl Trace Server with greater than 90% identity. The 8,099 uncharacterised reads were also searched against NCBI m36 using NCBI BLASTN. 2,901 (35.8%) had BLAST hits, but most had very low scores, just above the score threshold (half had a score of less than 33, 90% had a score of less than 59; Figure 2.4). The mapping algorithms of BLASTN and SSAHA2 therefore show small differences in output that may not significantly affect the final set of reliable mappings. Of the reads with BLAST hits, 76 also had hits to mouse sequences in the Ensembl Trace Server. However, there were also 295 reads that had hits to mouse sequences in the Trace Server but no BLAST hits. Again, these potentially represent sequences that have been incorrectly omitted from the mouse build.

Of the 5,198 (20.9%) remaining non-mapping reads, 4,363 were from tumours, 62 were from non-infected mice and 773 were from short infection time mice. Reads from control samples were highly over-represented (P=6.62x10⁻¹⁷²). There was also a highly

A		В			
Repeat Element	P-value	Repeat Element	P-value	Repeat Element	P-value
A-rich	0	4.5SRNA	7.90E-04	LTR/ERVL	1.21E-1
AT_rich	0	B1F	3.74E-22	LTR/MaLR	1.21E-8
C-rich	7.64E-241	B1F1	6.41E-08	Lx8	2.67E-1
CT-rich	1.94E-09	B1F2	1.03E-14	Lx9	9.04E-0
G-rich	0	B1_Mur1	1.07E-11	MIR	3.08E-2
GA-rich	0	B1_Mur2	1.05E-18	MIR3	8.14E-0
GC_rich	0	B1_Mur3	5.32E-05	MIRb	8.48E-2
GSAT_MM	3.76E-04	B1_Mur4	1.09E-07	MTD	4.17E-1
HAL1	2.76E-05	B1_Mus1	5.17E-05	MTE-int	2.38E-0
L1MEd	2.02E-03	B1_Mus2	5.55E-09	MTE2a	2.02E-0
LTR/ERV1	9.60E-21	B2_Mm2	4.07E-03	MTE2b	1.30E-0
MuLV-int	1.87E-26	B3	5.49E-49	MTEa	8.20E-0
MuRRS-int	2.23E-08	B3A	3.50E-25	ORR1D2	3.79E-1
RLTR4_MM-int	5.83E-45	B4	1.93E-20	ORR1E	1.21E-0
RLTR6-int	8.18E-03	B4A	7.06E-45	Other	2.10E-0
Simple_repeat	0	BC1_Mm	8.00E-03	PB1	2.48E-1
T-rich	8.15E-301	DNA/MER1_type	1.44E-27	PB1D10	1.99E-3
polypurine	4.87E-47	DNA/MER2 type	5.23E-04	PB1D9	5.99E-0
polypyrimidine	4.47E-07	ID	8.22E-03	RMER15	2.11E-0
		ID4	3.79E-10	RMER30	1.26E-0
		ID4_	2.25E-10	RSINE1	7.00E-6
		ID_B1	5.29E-69	SINE/Alu	9.88E-17
		L1M	4.16E-04	SINE/B2	3.31E-7
		L1M2	4.90E-03	SINE/B4	1.45E-18
		L1MC3	8.00E-03	SINE/ID	9.77E-2
		L1_Rod	3.19E-05	SINE/MIR	3.24E-4
		L2	4.73E-07	THER1_MD	3.16E-0
		LINE/L1	1.16E-65	URR1A	5.15E-0
		LINE/L2	5.59E-13	URR1B	1.09E-0
		LTR/ERVK	6.24E-07	scRNA	1.99E-0

Table 2.3. Repeat elements that are over-represented (A) and under-represented (B) among unmapped reads compared with unambiguously mapped reads. Over-represented elements include low-complexity regions and retrovirus-related elements, while under-represented elements include many that are frequently found in mouse genomic DNA. *P*-values were calculated using the 2-tailed Fisher Exact Test.

		Unambiguous
	reads (%)	mappings (%)
MMLV vector sequence	19.37	0.32
pBR322	4.84	0.08
low complexity	13.63	6.19
<=25 bp in length	26.62	0
>50% Ns	0.73	0.11
Other repeats	1.19	0.18
Total	66.38	6.88

Table 2.4. Summary of the proportions of unmapped and unambiguously mapping reads that contain vector sequences, or sequences of low complexity, low quality or short length. ">50% Ns" refers to sequences where the identity of more than 50% of bases is unknown. "Other repeats" refers to sequences containing repeat regions other than low complexity regions that were identified using RepeatMasker.

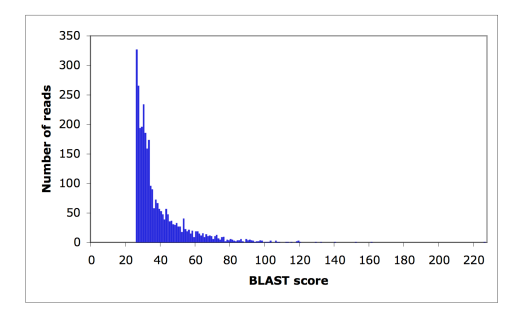


Figure 2.4. BLAST scores for uncharacterised unmapped reads. The majority of sequences that do not map with SSAHA2 but map with BLASTN have a low BLAST score.

significant under-representation (P=0) of reads containing both an LTR and an adapter sequence (1,972 reads) compared with those containing no LTR (1,065 reads) or no adapter (1,031 reads). These findings suggest a high presence of erroneous, contaminating reads. In addition, many reads were very short (Figure 2.3C) and may have failed to map due to the presence of a small number of differences from the reference genome sequence. Such differences may correspond to polymorphisms between the mouse strain *FVB*, from which the reads are derived, and strain *C57BL/6J*, upon which the mouse reference genome is based. 17.7% of reads were greater than 800 bp in length. The quality of reads rapidly deteriorates after ~700-900 bases of sequencing, which suggests that these are mostly of very poor quality or are chimeric sequences (discussed in Section 2.5).

There was also a highly significant over-representation of non-mapping reads without linker sequences ($P=1.61 \times 10^{-96}$) in the *Sleeping Beauty* dataset. Most of the non-mapping sequences flanked by an IR/DR and linker were short, with 50.2% being shorter than the 25 bp threshold for SSAHA2. As with the retroviral reads, there was a higher proportion of low-complexity sequences among unmapped *Sleeping Beauty* reads greater than 25 bp in length (3.1%) than among those that mapped unambiguously (2.4%). There was also a significant over-representation of GC-rich elements ($P=1.44 \times 10^{-4}$), and an underrepresentation of the LINE L1M2 (P=0.00265) and the rodent-specific LTR MTD ($P=2.85 \times 10^{-4}$) and SINEs B3 (P=0.00348), B3A (P=0.00265), PBID10 ($P=2.65 \times 10^{-3}$) and RSINE1 ($P=2.74 \times 10^{-4}$).

2.5 Filtering the mapped reads

During PCR amplification, unrelated sequences can hybridise to one another, resulting in clones comprising chimeric sequences. It is important that retroviral reads contain the LTR sequence since, if the part of the read that maps to the genome is directly adjacent to the LTR, the location of the mapped DNA is likely to be the true location of the retroviral insertion. For reads that contain an LTR and an adapter, these sequences should directly flank the genomic DNA. Therefore, for each read, the coordinates of the LTR and adapter sequences identified by cross_match were compared to the coordinates of the region that mapped to the mouse genome using SSAHA2. If the gap between these regions was within 5 bp, the read was accepted. Since the junction between the LTR and the genomic DNA is most important, reads were also accepted if the DNA that mapped to

the genome was within 5 bp of the LTR but there was a gap between the genomic DNA and the adapter, or if the read did not contain an adapter sequence. Base miscalling in low quality reads may result in a SSAHA2 alignment that does not extend right up to the LTR sequence even though the LTR and genomic DNA are directly adjacent. Therefore, up to a distance of 30 bp, reads were accepted if the sequence between the LTR and the aligning genomic DNA did not contain any restriction sites for *Tsp509I* (i.e. 5'-AATT-3') or *Sau3AI* (i.e. 5'-GATC-3'), depending on which had been used in the PCR. If a restriction site intercepts the LTR and genomic DNA, it is possible that the genomic DNA that immediately flanks the LTR, and represents the true location of the virus in the genome, may not have been mapped because it is too small or of poor quality but that it has ligated to a contaminating DNA fragment that has been mapped.

The components within the read should be in the configuration LTR-genome-adapter or adapter-genome-LTR. Therefore, any reads that had a different configuration were discarded. For example, the configuration LTR-adapter-genome suggests that a contaminating fragment of genomic DNA has ligated to the end of the adapter, and that the true flanking region of the LTR could not be mapped because it is too short or of poor quality. Reads containing multiple LTR or adapter sequences were subjected to the same filtering criteria, whereby reads were discarded if the sequence for one LTR did not directly abut the genomic sequence or the adapters intercepted the LTR and genomic sequence. Reads with no LTR were rejected unless an LTR identified by reducing the minimum score for cross_match to 5 followed the rules outlined above for stronger LTR matches.

81,846 reads (90.3%) were retained after filtering. Both accepted and rejected reads with gaps of greater than 5 bp were subjected to further analysis. If the average quality (Phred) score of the gap region was less than 30, the read was accepted as the gap may contain miscalled bases, causing SSAHA2 to prematurely terminate extension of the alignment across the full length of the genomic DNA within the read. Reads were also accepted if they mapped to the same location as other reads from the same tumour that did not contain a gap. The final set of accepted reads totalled 81,910 (90.3%). The filtering procedure is summarised in Figure 2.5. There were significantly more reads of greater than 800 bp in length among removed reads (39.5%) than retained reads (6.7%, P=0) and removed reads mapped to the genome with a lower percentage identity (92.6% \pm 5.9) than retained reads (99.0% \pm 2.3).

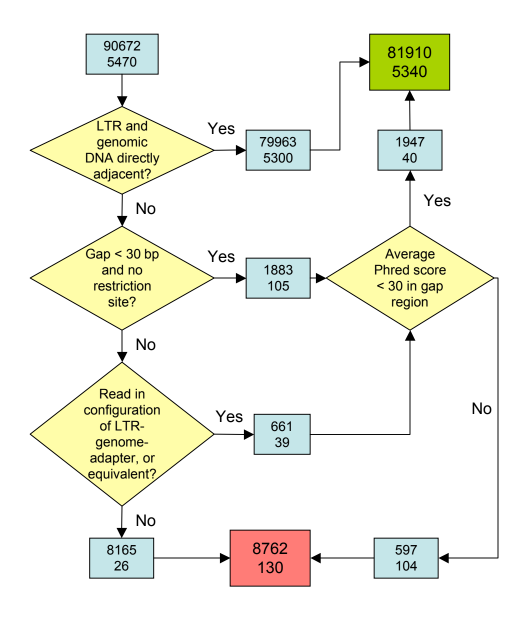


Figure 2.5. The filtering process for single mapping reads. Blue boxes contain the counts for accepted or rejected reads at each stage, where the top number in each box refers to the count for retroviral reads and the bottom number refers to the count for transposon reads. Final counts for the accepted and rejected reads are shown in the green and red box, respectively.

29,769 reads mapped to multiple locations in the mouse genome. These may be chimeric or low quality reads, or they may represent retroviruses that have inserted into duplicated or repetitive regions of the genome. 15,036 reads were identified where at least one of the mappings matched the criteria used for filtering single mapping reads. For 8,429 of these reads, only one mapping matched the criteria, and this was retained in the dataset while other mappings were discarded. Among the remaining 6,607 reads, there were 465 where only one mapping had an alignment of 100% identity. These mappings were retained and all others were discarded. In total, 8,894 (29.9%) of reads that mapped ambiguously were retained. As with the unambiguous mappings, there was a significant over-representation of reads greater than 800 bp in length in the removed reads (13.4%) compared to the retained reads (8.9%, $P=3.40 \times 10^{-28}$). The retained reads were pooled together with the retained single mapping reads, giving a total of 90,804 reads.

Transposon insertions were filtered using the same criteria, except that gaps between IR/DRs and genomic DNA were scanned for *NlaIII* (5'-CATG-3') or *BfaI* (5'-CTAG-3') restriction sites, depending on whether the IR/DR was from the left or right end of the transposon. 5,340 (97.6%) of reads mapping to a single genomic location and 941 (50.6%) of those mapping ambiguously were accepted. The filtering of reads that mapped unambiguously is summarised in Figure 2.5.

2.6 Identification and filtering of insertion sites

As 96 reads were sequenced for each PCR, there may be multiple reads that correspond to the same insertion site. The exact genomic coordinates and orientation of the retroviral or transposon insertion represented by each read were determined using the coordinates and orientation of the genomic DNA, resolved by SSAHA2. The methods are summarised in Figure 2.6. Reads from a single PCR mapping to within 2 kb were then clustered into a single insertion site, resulting in 29,553 retroviral insertion sites and 2,821 transposon insertion sites across all PCRs.

It is possible that endogenous LTR sequences within the mouse genome could be the target of non-specific PCR amplification in the retroviral screen. NCBI BLASTN, adjusted to search for short sequences (Word size 7, E value 10,000, filter OFF), was therefore used to identify sequences in NCBI m36 that resembled the MuLV LTR. In a preliminary analysis on NCBI m34, all 15 bp fragments of the LTR sequence

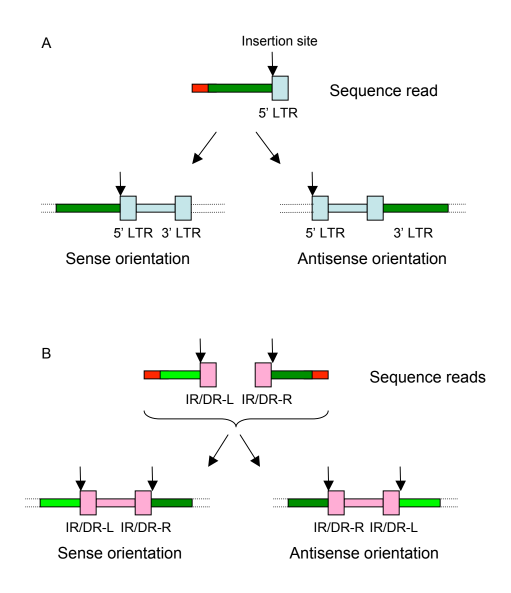


Figure 2.6. Determining the exact insertion site and orientation of retroviral (A) and transposon (B) insertions in the mouse genome. Adapter sequences are shown in red; genomic DNA is shown in green. A. The point of insertion is the genomic nucleotide adjacent to the 5' LTR of the MuLV retrovirus (shown in blue) in the sequence read. Alignment to the forward strand of the mouse genome indicates that the retrovirus has inserted in the 5'-3' orientation and the insertion site corresponds to the last nucleotide in the reported alignment. Alignment to the reverse strand indicates that the retrovirus has inserted in the 3'-5' orientation and the insertion site corresponds to the first nucleotide in the alignment. B. As for retroviral insertions, except that there are two sets of reads, containing a left or right IR/DR sequence. The T2/Onc transposon is shown in pink.

5'-GCTAGCTTGCCAAACCTACAGGTGGGGTCTTC-3' were used query as sequences, but 99% of the insertions near LTR-like sequences in short infection time mice were identified using LTR fragments 5'-GCTTGCCAAACCTAC-3' and 5'-CTTGCCAAACTACA-3', and therefore only these fragments were used in the current analysis. All of the apparent insertions in the uninfected control samples should be PCR artefacts, while short infection time DNA is expected to contain a higher proportion of PCR artefacts than tumour DNA. Among the 1,399 reads mapping to LTR-like sites, there were significantly more from uninfected samples and from short infection time samples than expected by chance $(P=3.17 \times 10^{-26} \text{ and } P=0, \text{ respectively})$. These findings support the theory that reads mapping to sites that resemble the retroviral LTR are the result of non-specific PCR amplification and do not represent real insertion sites. For example, 174 samples contain an insertion in the aminoadipate-semialdehyde synthase (Aass) gene, but the insertions are adjacent to a 14 bp sequence that precisely matches the MuLV LTR and are therefore likely to be false positives. Figure 2.7 shows these insertions displayed in Ensembl. The Distributed Annotation System (DAS) server ProServer was used to display both the retroviral and the transposon insertion sites in the context of the mouse genome in Ensembl contigview. Ensembl is a DAS client that can integrate genome annotation information from multiple servers, enabling users to view and compare annotations from multiple sources in a single display. All 1,399 reads at 675 LTR-like sites were removed from the dataset.

Apparent insertions in non-infection and short infection time samples were removed from the dataset, but a decision was made not to remove tumour insertions that mapped to the same locations. A preliminary analysis, in which the reads were mapped to mouse build NCBI m34, showed that many of the reads from non-infection and short infection time samples mapped to cancer genes that are known targets of retroviral insertional mutagenesis. Insertions within 5 kb of *Myc* were identified in 41.7% of non-infection samples, 26.2% of short infection time samples and 30.4% of tumour samples (see Figure 2.8). Similarly, the proportions of insertions from non-infection and short infection time samples in and around *Mycn* were 12.5% and 35.9%, respectively, but just 8.9% in tumour samples. Findings for the short infection time dataset could indicate that *Myc* and *Mycn* are insertion hotspots, or that selection for *Myc* and *Mycn* insertions occurs at an early time point. However, these explanations do not justify the presence of such insertions in non-infection samples. As all non-infection insertions map to only 142 distinct coordinates, it seems an unlikely coincidence that *Myc* and *Mycn* are targeted by

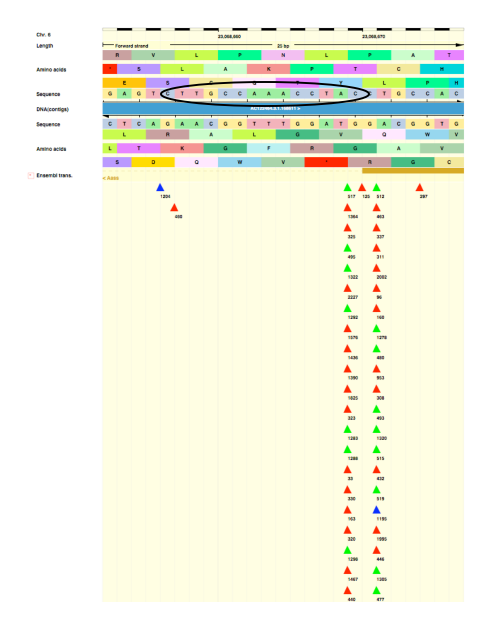


Figure 2.7. Insertions in the mouse aminoadipate-semialdehyde synthase (*Aass*) gene are PCR artefacts that map to an LTR-like sequence in the mouse genome. 174 samples contain an insertion in this region (46 are shown here as triangles). Insertions from tumours, short infection time samples and uninfected samples are shown as red, green and blue triangles, respectively. The LTR-like sequence is circled.

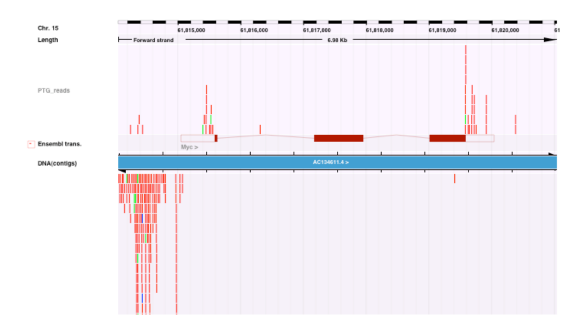


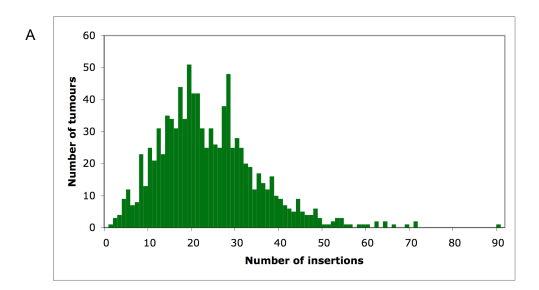
Figure 2.8. A high proportion of insertions in control samples map to the *Myc* gene. This figure shows some of the insertions in and around the *Myc* gene. Insertions from tumours, short infection time samples and uninfected samples are shown as red, green and blue rectangles, respectively.

non-specific primer binding, and there are no LTR-like sequences near these genes. The insertions may result from contamination during PCR or, even more worryingly, unintended infection of mice in the animal facility. The control samples are useful for picking out possible contaminants, like those described above that map to LTR-like sequences, but discarding all tumour insertions that map to the same sites as control insertions would most likely result in the removal of a considerable number of real insertions.

The insertion sites identified in individual PCRs were clustered into 22,579 retroviral insertion sites from 997 tumours. The average number of inserts per tumour was 23.49 ± 11.42 (Figure 2.9A). There were, on average, 3.72 ± 6.21 reads per insert (Figure 2.9B). The 2,821 transposon insertion sites identified in individual PCRs were clustered into 2,643 insertion sites from 73 tumours. There was an average of 36.21 ± 18.55 inserts per tumour, and 2.38 ± 4.08 reads per insert.

2.7 Estimating the coverage of the mutagenesis screens

Measuring the overlap of insertion sites between PCRs for an individual tumour gives some indication of the proportion of insertions that were identified in the screens. There were 616 tumours for which retroviral insertions had been identified from one PCR using Sau3A1 and one using Tsp5091. These contained 10,733 and 8,580 insertions identified using Sau3A1 and Tsp509I, respectively, of which 2,968 were identified using both enzymes. The overlap between PCR experiments was therefore 18.2%, rising to 32.9% if insertions represented by a single read were omitted. More than one enzyme is required because individual enzymes do not cut the genomic DNA sufficiently close to all insertions to enable PCR amplification of the intervening sequences. Since the overlap between PCRs is low, it seems likely that even two enzymes do not give sufficient coverage. However, the difference between the 2 PCRs may also result from insufficient sequencing, such that genomic DNA flanking an insertion is amplified but is not sequenced. This may explain why a high proportion of insertion sites represented by a single read are not identified by both PCRs, since they are more likely to be rare insertions that have a low representation in the PCR mixture and are less likely to be sequenced.



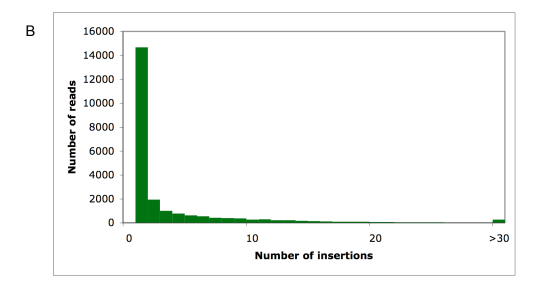


Figure 2.9. The number of insertions per tumour (A) and reads per insertion (B).

For 3 tumours, genomic DNA was also digested using BstYI, and 384 reads were sequenced. Reads were mapped to NCBI m34 and were compared to Sau3A1 and Tsp509I reads from the same tumours, also mapped to NCBI m34. There was a 39.0% overlap between insertion sites identified from PCRs using Sau3A1 and BstY1, and a 23.4% overlap between those identified from PCRs using Tsp509I and BstYI. A higher overlap is expected between Sau3A1 and BstYI because the BstYI target site (5'-RGATCY-3') contains the target sequence for Sau3A1. BstYI cuts less frequently than Sau3A1 and Tsp5091. For reads generated using Sau3A1 or Tsp5091, the average distance between the LTR and the restriction site at which the DNA was cut was 308.41 bp, but for reads generated using BstYI, the average distance was 386.52 bp. It is therefore difficult to directly compare the PCRs because fragments of BstYI-digested DNA will be longer, on average, and there is likely to be a higher proportion that cannot be amplified by PCR. For insertion sites that were identified using Sau3A1 or Tsp5091 but not using BstYI, the genomic DNA within the corresponding reads was scanned for BstYI target sites. Likewise, for insertion sites that were uniquely identified using BstYI, the genomic DNA was scanned for Sau3A1 and Tsp509I target sites. If the sequencing depth of 96 reads was sufficient, insertion sites should only be uniquely identified using *BstYI* if there are no Sau3A1 and Tsp509I target sites close enough to the insertion site for successful PCR. A BstYI target site was identified at a distance equal to, or closer than, the Sau3A1 or Tsp5091 site in reads corresponding to 2 out of 15 unique Sau3A1 insertion sites and 4 out of 20 unique Tsp509I insertion sites. However, for Sau3A1 and Tsp509I, a target site was identified at a distance equal to, or closer than, the BstYI site for 21/21 and 14/29 unique BstYI insertions, respectively. This suggests that more insertion sites could be obtained by increasing the sequencing depth to 384 reads per PCR, and that an even greater depth may be required to saturate the screen. However, as only 3 tumours were used in this analysis, and different enzymes were used to generate the digested DNA for 96-read and 384-read sequencing, it is difficult to reach any firm conclusions about the number of enzymes and the sequencing depth required for maximum coverage.

For the Sleeping Beauty screen, there were 60 tumours for which 2 PCRs were performed using restriction enzymes *BfaI* and *NlaIII*. Only 159 insertions (6.9%) were shared from 1,161 insertion sites identified using *BfaI* and 1,310 identified using *NlaIII*.

2.8 Analysis of the distribution of insertions around mouse genes

The long-range effects of MuLV enhancer mutations can complicate the identification of mutated genes. Analysing the distribution of insertions around mouse genes, and in particular, around the mouse orthologues of known cancer genes, can help to define rules for predicting which gene is being mutated by an insertion. The genomic coordinates and orientation of all mouse protein-coding and miRNA genes were extracted from Ensembl using the Perl API version 45 36f, and insertions were counted in 100 bp intervals up to 20 kb upstream and downstream of each gene. The gene orientation was used to determine the orientation of insertions with respect to each gene. Figures 2.10A-D show the number of genes that contain insertions in each 100 bp interval upstream and downstream in the sense and antisense orientation with respect to each gene. In the full set of genes, the number of sense and antisense insertions peak at around 500-600 bp upstream, and a similar pattern is observed around the mouse orthologues of known cancer genes. These sense and antisense insertions are likely to represent promoter and enhancer mutations, respectively (see Section 1.4.2.1.1), with the peak representing the optimal distance for mutation. Downstream insertions show a relatively uniform distribution with similar proportions of insertions in the sense and antisense orientation. This may indicate that most are randomly occurring non-oncogenic insertions, or that there is no optimum distance for an enhancer mutation that acts downstream of a gene. It is also likely that some of these insertions are affecting adjacent genes, and variation in the distance between genes may contribute to the observed distribution. There is also no obvious pattern in the downstream counts of cancer genes with insertions. The plots in Figures 2.10A-D show the counts of genes with insertions up to 20 kb upstream or downstream, regardless of whether adjacent genes intercept the 20 kb region. However, counting only as far as the adjacent gene gives a similar distribution, with peaks at 500-600 bp upstream in both orientations, and an essentially uniform distribution downstream. Counting actual insertions, rather than the number of genes containing insertions, skews the distribution towards genes containing larger numbers of insertions. For example, Myc contains many enhancer mutations, and the highest peak might represent the optimal distance for an enhancer mutation of Myc, rather than for all genes. However, once again the highest peak is at 500-600 bp upstream. A similar distribution is also obtained by counting only the genes that contain insertions represented by more than one read.

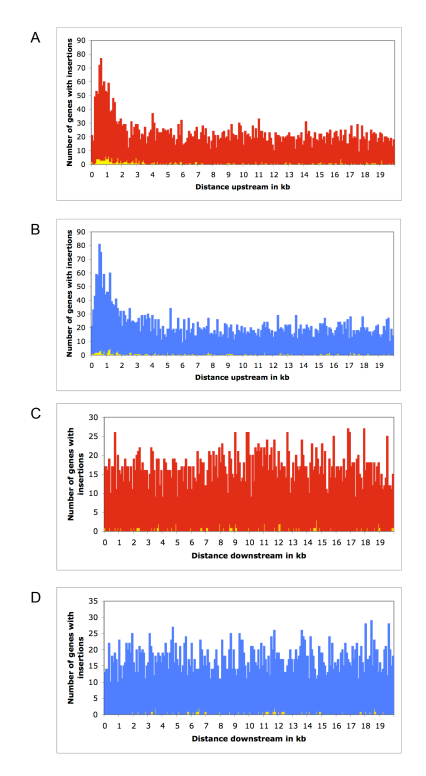


Figure 2.10. The number of genes with insertions in 100 bp intervals up to 20 kb upstream in the sense (A) and antisense (B) orientation and downstream in the sense (C) and antisense (D) orientation with respect to the gene. Counts of cancer genes with insertions in each interval are shown in yellow.

Known oncogenes and tumour suppressor genes with intergenic insertions up to 20 kb upstream and/or downstream are shown in Tables 2.5A and 2.5B, respectively. Tumour suppressor genes are expected to contain intragenic insertions that result in truncated, inactivated, transcripts. Of the 12 tumour suppressor genes flanked by intergenic insertions, only 1 has insertions represented by more than one read. This suggests that "singleton" insertions, i.e. insertions represented by a single read, are less likely to contribute to oncogenesis. They may be rare insertions that are not in the dominant tumour lineage or have integrated into a single lineage late on in tumour development, or they may be PCR artefacts. 8 known oncogenes had insertions within 2 kb upstream in the sense orientation, and 22 had insertions within 20 kb. These numbers fell to 5 and 9, respectively, if singleton insertions were removed. Likewise, there were 9 oncogenes with antisense insertions within 2 kb upstream, and 29 with insertions within 20 kb, but only 5 and 13, respectively, without singletons. As well as representing rare insertions, singleton insertions may result from limitations in PCR and sequencing depth. Therefore, in order to maximise the number of candidates that could be identified, singleton insertions were retained in the analysis since, if they are not important in tumourigenesis, they should not form statistically significant CISs (see Section 2.10).

Insertions around the *Pim1* oncogene (Figure 2.11A) suggest that downstream sense and antisense insertions can contribute to tumourigenesis. Downstream sense insertions also appear to affect the *Kit* oncogene (Figure 2.11B). However, there are fewer oncogenes with downstream sense insertions than upstream insertions, and even fewer with downstream antisense insertions. For some of the genes, e.g. *Gata1* (Figure 2.11C), it does appear that the downstream insertions are in fact mutating an adjacent gene. These observations concur with prior work, in suggesting that upstream antisense and sense insertions, corresponding to enhancer and promoter mutations, respectively, are the most common forms of mutation. Downstream insertions, while less common, appear to be more frequent in the sense orientation, which is the proposed orientation for downstream enhancer mutations (see Section 1.4.2.1.1).

Of the 22 oncogenes that had sense insertions within 20 kb upstream, 20 were still identified when the upstream limit was set to the 3' end of the upstream gene. All 9 genes without singleton insertions were similarly identified. Likewise, 23 out of 29 genes, including all 13 genes without singletons, that had antisense insertions within 20 kb upstream were still identified. 9 out of 14 genes containing downstream sense

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А Insertior Gene 20 kb no 2 kb no Within Orientation Mouse Ensembl ID 20 kb all singletons Name Elf4 singleton limits ENSMUSG0000018654 Ik7f1 22 14 0 0 22 ENSMUSG0000062312 Erbb2 13 n ñ ň ENSMUSG0000022346 Myc 12 11 8 7 11 8 ENSMUSG0000006362 Chfa2t3 0 ENSMUSG00000026923 ENSMUSG0000000409 Notch1 0 Lck Cbl 0 0 ENSMUSG0000034342 Pim1 Rhoh ENSMUSG0000024014 6 5 6 ENSMUSG00000029204 ENSMUSG0000032688 Malt1 Upstream 3 0 0 0 ENSMUSG0000036986 Pml 3 Ó 0 0 ENSMUSG00000025408 ENSMUSG00000037169 Ddit3 0 0 0 Mycn ENSMUSG0000000184 Ccnd2 2 ENSMUSG00000059248 ENSMUSG00000020893 Sept9 Per1 0 0 0 0 0 ENSMUSG0000021377 Dek 2 0 0 0 ENSMUSG0000021356 Irf4 0 0 0 ENSMUSG0000027829 ō Ccnl1 ENSMUSG0000066306 Numa1 2 0 0 0 ENSMUSG0000003282 Plag1 Mvc 0 ENSMUSC000002234 388 303 203 299 ENSMUSG0000026923 Notch1 19 16 19 16 10 11 0 0 15 10 ENSMUSG0000024014 Pim1 ENSMUSG00000070348 ENSMUSG00000018654 Ccnd1 14 14 13 13 10 0 0 Ikzf1 Ccnd2 ENSMUSG0000000184 4 ENSMUSG0000006362 Cbfa2t3 8 0 4 0 ENSMUSG0000006389 ENSMUSG00000022952 Mpl . Runx1 0 8 0 Plag1 Sept9 Flt3 ENSMUSG0000003282 n 0 0 8 6 5 ENSMUSG00000059248 ENSMUSG00000042817 0 0 ENSMUSG0000031103 Elf4 4 0 0 ENSMUSG0000043962 Akt3 Ó 3 0 3 Upstream ENSMUSG00000048251 Bcl11b antisense ENSMUSG0000030745 Il21r 3 0 0 0 ENSMUSG0000034342 Chl 0 0 0 ENSMUSG00000025958 ENSMUSG00000020453 Creb1 2 Patz1 0 0 0 ENSMUSG0000021457 Svk 2 0 0 ENSMUSG0000021356 ENSMUSG00000056234 Irf4 0 2 0 0 0 Ncoa4 0 ENSMUSG0000022797 Tfrc 2 0 0 0 ENSMUSG0000032698 1 mo2 2 Ó 0 0 ENSMUSG0000002028 ENSMUSG00000025408 MII1 2 Ddit3 0 0 ENSMUSG0000041358 Nut Lck 0 0 2 0 ENSMUSG0000000409 ENSMUSG0000029438 ENSMUSG00000024014 Bcl7a 17 Pim 1 ENSMUSG00000038227 ENSMUSG00000022346 Hoxa9 0 0 6 5 0 Myc , Fstl3 ENSMUSG0000020325 0 0 4 Cars Ddx6 ENSMUSG0000010755 3 n 0 ENSMUSG00000032097 Downstream ENSMUSG00000034041 Lyl1 0 0 0 ENSMUSG0000057329 Bcl2 2 0 0 0 ENSMUSG00000069305 Hist4h4 ENSMUSG0000029204 Rhoh 0 0 0 ENSMUSG0000005672 Kit 2 0 0 0 ENSMUSG00000034165 ENSMUSG00000004895 Ccnd3 Prcc ENSMUSG0000028718 Stil ENSMUSG0000031162 Gata1 Pim1 ENSMUSG00000024014 ENSMUSG00000030745 Il21r 0 0 ENSMUSG0000069305 Hist4h4 3 0 0 0 Downstream ENSMUSG0000020453 Patz1 ň ñ ň ENSMUSG0000068860 Gm128 antisense 0 0 0 ENSMUSG0000070002 FII 0 0 ENSMUSG00000026656 ENSMUSG00000020167 Fcgr2b Tcfe2a 0 0 0 ENSMUSG000003404 vl1 В Insertion Gene 20 kh no 2 kh no Within orientation Mouse Ensembl ID ENSMUSG0000003068 name Stk11 20 kb all 2 kb all singletons limits singletons Upstream ENSMUSG0000009863 Sdhb sense ENSMUSG0000036712 ENSMUSG00000009863 Cyld Sdhb ENSMUSG0000003068 Stk11 0 0 0 Upstream ENSMUSG0000013663 Pten 0 0 ENSMUSG0000026526 Fh1 antisense ENSMUSG0000028687 Mutvh 0 0 0 ENSMUSG0000034023 Fancd2 Blm ENSMUSC000003052 ENSMUSG0000024947 Downstream Men1 0 0 0 sense ENSMUSG0000025231 Sufu 0 ENSMUSG0000044702 ENSMUSG00000040084 Downstream Bub1t antisense ENSMUSG0000024947 Men1

Table 2.5. Number of intergenic insertions up to 20 kb upstream and downstream of known oncogenes (A) and tumour suppressor genes (B) from the Cancer Gene **Census.** "20 kb all" and "2 kb all" give the total number of insertions up to 20 kb and 2 kb upstream/downstream. "2 kb no singletons" and "20 kb no singletons" give the number of insertions represented by more than 1 read. "Within limits" gives the number of insertions up to the adjacent upstream or downstream gene.

Chapter 2

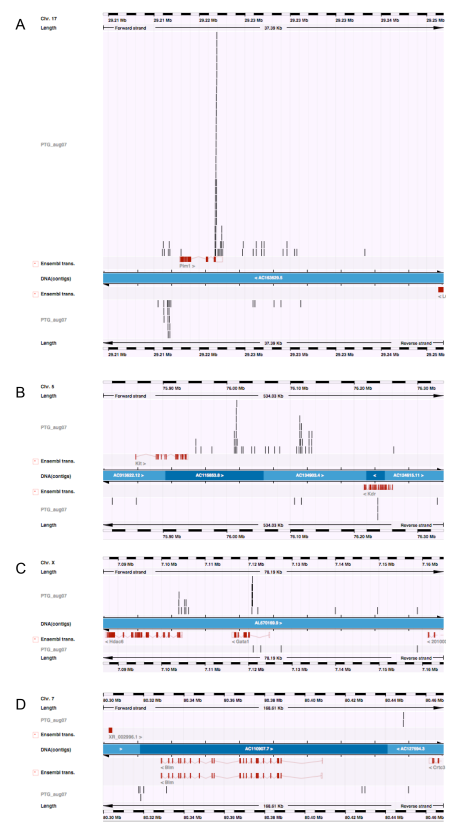


Figure 2.11. Insertions around known cancer genes *Pim1* (A), *Kit* (B), *Gata1* (C) and *Blm* (D). Insertions are shown as black bars in the context of Ensembl genes, shown in red. Insertions above and below the blue bar are in the sense and antisense orientation, respectively.

insertions, and 5 out of 10 genes containing downstream antisense insertions, were still identified when the downstream limit was set to the 5' end of downstream gene. The lower proportion for genes with downstream sense insertions, and lower still for downstream antisense insertions, most likely reflect the fact that these insertions are less likely to contribute to oncogenesis. The same applies to tumour suppressor genes, of which only 4 out of 12 were still identified when the upstream and downstream limits were set to adjacent genes. Therefore, based on these results, it seems reasonable to assign insertions to a gene only if they are within the limits of adjacent genes.

As indicated by the high proportion of singleton insertions and the presence of insertions beyond the boundaries of adjacent genes, it is likely that most of the tumour suppressor genes listed in Table 2.5 are not mutational targets. However, the *Blm* gene contains an intragenic insertion, as well downstream sense insertions (Figure 2.11D). It is possible that the intergenic insertions are not oncogenic, or that they are affecting a nearby gene, or there may be an error in the *Blm* gene prediction in Ensembl, such that the insertions appear to be intergenic but are in fact intragenic. Alternatively, the insertions could be disrupting a downstream regulatory element, resulting in reduced transcription or gene inactivation.

There is no obvious pattern in the distribution of transposon insertions upstream or downstream of genes. This is not surprising for upstream antisense and downstream insertions, since the *Sleeping Beauty* transposon T2/Onc has low enhancer activity. However, insertions in the upstream sense orientation might be expected to follow a similar distribution to those in the retroviral screen. The T2/Onc promoter is perhaps not as strong as the MuLV promoter and mostly mutates by producing truncated transcripts, rather than by increasing levels of the wildtype protein. Alternatively, some of the apparent promoter mutations in the retroviral screen may in fact be enhancer mutations, or a high background of non-oncogenic T2/Onc insertions may be masking the true pattern of oncogenic mutations. There is only one oncogene (*Irf4*) and no tumour suppressor genes with sense or antisense insertions up to 20 kb upstream or downstream. While this may in part reflect the smaller size of the dataset, it also suggests that oncogenic T2/Onc insertions are usually intragenic.

Chapter 2

2.9 Assigning insertions to genes

The coordinates and orientation of the longest transcript of all protein-coding and miRNA genes in the mouse genome were extracted from Ensembl version 45 36f using the API. Genes nestled within other genes were removed from the analysis, since these complicate the specification of gene boundaries for assigning intergenic insertions to genes. Intragenic retroviral insertions were assigned to the genes within which they resided. For intergenic insertions, the flanking genes were identified. If an insertion was upstream of the first gene or downstream of the last gene on a chromosome, it was assigned to the first or last gene, respectively. If only one of the flanking genes was within 100 kb of the insertion, that gene was assigned the insertion. If one of the flanking genes contained intragenic insertions, the intergenic insertions were also assigned to that gene. Based on the observations of insertions around known cancer genes outlined in Section 2.8, if an insertion was in the downstream antisense orientation relative to one gene, but in a different orientation relative to the other gene, it was assigned to the other gene, and other intergenic insertions were also assigned to that gene. Finally, for the remaining unassigned intergenic insertions, the nearest insertion to each gene was identified, and all insertions were assigned to the gene that had the nearest insertion. Sleeping Beauty T2/Onc insertions were processed in a similar way, except that if an intergenic insertion was in the upstream sense orientation with respect to one gene, but in a different orientation with respect to another gene, it was assigned to the former gene.

2.10 Identifying statistically significant common insertion sites

Oncogenic insertions must be distinguished from a background of non-oncogenic insertions. Insertions from different tumours that reside in the same genomic region, defined as common insertion sites (CISs), are more likely to contribute to tumourigenesis, but statistical approaches are required to determine whether a CIS is significantly different to the random, background distribution of insertions. Monte Carlo simulations, and a more recent method that uses a kernel convolution-based statistical framework, have been applied to the retroviral and *Sleeping Beauty* datasets, and the results compared.

2.10.1 Monte Carlo simulations

This method is based on the procedures described in Suzuki et al. (2002) and Mikkers et al. (2002). The 26,144 retroviral insertions were randomised across the mouse genome (golden path length 2,661,205,088 bp, mouse build NCBI m36). A wide range of window sizes were used, and the number of windows containing at least M insertions were counted, where M was a number of 2 or more (up to 14 for large window sizes). The randomised insertions were ordered across the genome $(X_{[1]} \text{ to } X_{[26,144]})$, and windows were taken as the interval from $X_{[i]}$ to $X_{[i+M-1]}$ (see Suzuki *et al.*, 2002). If the distance between an insertion and the next M-1 insertions on the chromosome was less than the window size $(X_{[i+M-1]} - X_{[i]})$, it was counted as a CIS. The next window was positioned at i+M. 100,000 iterations were performed, and mean counts and the 0.99 upper quantile were calculated for each number (M) of insertions. This gives the number of CISs of M insertions that one would expect to find by chance in each window size, and the maximum number for P=0.01. As in Mikkers et al. (2002), fractions (represented as Efr) of 0.001, 0.005 and 0.01 of the total number of insertion sites expected to be random CIS clusters were calculated. These are 26.144, 130.72 and 261.44, respectively, for retroviral insertions, and 2.64, 13.22 and 26.43, respectively, for transposon insertions. Maximum window sizes for significant CISs for varying values of M can then be calculated by finding the window size at which the upper quantile of the random distribution is less than the expected number of false CISs (Table 2.6).

For each gene to which insertions had been assigned, the number of insertions was counted and the distance between insertions was calculated. If any of the insertions fell within a window size that met the criteria for a significant CIS, the gene was accepted as a candidate cancer gene. For an *Efr* of 0.001, 0.005 and 0.01, the number of identified candidates in the retroviral screen was 1,404, 1,677 and 1,829, respectively. For the *Sleeping Beauty* screen, the number of candidates was 62, 91 and 115, respectively. This approach differs from the method in Suzuki *et al.* (2002) in that insertions were considered in the context of each gene, and a consistent approach was used to identify all candidates. In Suzuki *et al.* (2002), CISs were identified independently of genes, and then assigned to genes, but further genes were selected as candidates if they contained multiple insertions that were not in significant CISs. In addition, the method in Suzuki *et al.* (2002) uses 3 fixed window sizes to define CISs, which, particularly in a screen of this

Λ	_						Numbe	r of inse	ertions					
A	Efr	2	3	4	5	6	7	8	9	10	11	12	13	14
	0.001	0.1	5.0	19.5	45.0	80.0	120.0	168.0	220.0	275.0	333.0	391.5	455.0	521.0
	0.005	0.5	10.0	35.0	75.0	120.0	175.0	235.0	299.5	366.0	437.5	510.0	586.0	663.0
	0.01	1.0	14.4	45.7	95.0	150.0	210.0	280.0	351.5	425.5	505.0	587.5	671.0	757.0

D			N	umber	of inse	ertions		
5	Efr	2	3	4	5	6	7	8
	0.001	1	45	193	450	800	1200	1650
	0.005	5	101	345	750	1200	1750	N/A
	0.01	10	142	452	950	1500	2100	N/A

Table 2.6. Maximum window sizes in kb for significant CISs for varying numbers of insertions in the retroviral (A) and *Sleeping Beauty* (B) screens. Window sizes are given for *Efr* (fraction of the number of insertion sites expected to be random CIS clusters) of 0.001, 0.005 and 0.01, for which the corresponding numbers of false CISs are 26.144, 130.72 and 261.44 for retroviral insertions, and 2.64, 13.22 and 26.43 for transposon insertions. N/A is given where the window size is larger than any gene plus 100 kb of flanking sequence, and is therefore not relevant to the analysis.

	Number of	Number of non-			
Method	cancer genes	cancer genes	Accuracy	Coverage	мсс
КС	42	487	0.0794	0.1193	0.1433
MC <i>Efr</i> =0.001	66	1144	0.0545	0.1875	0.1010
MC <i>Efr</i> =0.01	80	1500	0.0506	0.2273	0.0944
All	175	5483	0.0309	0.4972	0.0562

р		Number of	Number of non-			
D	Method	cancer genes	cancer genes	Accuracy	Coverage	МСС
	KC	6	21	0.2222	0.0170	0.3115
	MC <i>Efr</i> =0.001	10	45	0.1818	0.0284	0.2708
	MC <i>Efr</i> =0.01	11	90	0.1089	0.0313	0.1836
	All	59	1279	0.0441	0.1676	0.0767

Table 2.7. Comparison of the methods used to generate candidate cancer genes lists from the retroviral (A) and *Sleeping Beauty* (B) screens. The accuracy, coverage and Matthew's correlation coefficient (MCC) are based on the number of known cancer genes in the candidate gene lists. KC = kernel convolution-based framework, MC *Efr*=0.001 and MC *Efr*=0.01 refer to Monte Carlo simulations using *Efr* (fraction of the number of insertion sites expected to be random CIS clusters) of 0.001 and 0.01, All = all genes to which insertions were assigned, regardless of whether they were statistically significant.

size, could result in some CISs being missed. Therefore, this method uses the approach in Mikkers *et al.* (2002) to define maximum window sizes for all values of M.

2.10.2 Kernel convolution

As discussed in Section 1.4.2.1.2, the Monte Carlo (MC) method may not be suitable for very large datasets. Significant CISs were therefore also identified using the kernel convolution (KC)-based statistical framework (de Ridder *et al.*, 2006). A list of insertions was supplied to the Netherlands Cancer Institute, where Jeroen de Ridder produced and returned a list of genomic coordinates corresponding to CISs generated using the KC method. In this method, a kernel function is placed at every insertion in the dataset and the number of insertions at any genomic position can be estimated by summing all the kernel functions. Insertions in close proximity to one another will produce a higher peak in the estimated number of insertions (de Ridder *et al.*, 2006, also discussed in Section 1.4.2.1.2).

867 retroviral cross-scale CISs were identified using the KC-based framework with P=0.05. These are all the CISs identified using a range of kernel widths (0.05, 0.1, 0.25, 0.5, 1, 2.5, 5, 10, 30, 50, 100 and 150 kb). The kernel width controls the smoothness of the estimated number of insertions (de Ridder et al., 2006). In other words, it controls the size of the genomic region in which neighbouring insertions affect the estimate of the number of insertions at the observed insertion. For each CIS, the flanking genes were identified using the Ensembl API version 45 36f and were compared to the genes identified using MC simulations. Among the 867 KC CISs, there were 765 where the nearest or further gene was represented in the Efr=0.01 MC list of candidates. Genes flanking the remaining 102 KC CISs may be missing from the MC list because insertions have been misassigned or because of differences between the two statistical approaches. As described in Section 1.4.2.1.2, for large screens, the statistically significant window size in the MC method may be so small that it is less than the width of biologically relevant CISs, causing these to be missed. Many of the CISs unique to the MC analysis are likely to be false positives since, at an Efr of 0.01, 261.44 randomly occurring CISs are expected.

652 CISs identified using a kernel width of 30 kb (P=0.05) were chosen for further analysis since this width, which was also used in Uren *et al.* (2008), should capture a high

proportion of biologically relevant CISs without splitting independent CISs or merging CISs that represent different types of mutation within a gene. For example, in genes that are mutated by multiple mechanisms, upstream enhancer mutations may form one CIS, while intragenic or downstream enhancer mutations may form another. For intragenic CISs, the gene containing the CIS was defined as the candidate cancer gene. For intergenic CISs, the flanking genes were compared to the list of candidates generated using Monte Carlo (MC) simulations. Where one of the flanking genes was within the MC list, this was chosen as the candidate gene. Where both nearest genes were within the MC list, both were initially included in the KC list because it is possible that a CIS could be mutating multiple nearby genes. Where neither nearest gene was in the MC list, the nearest genes were compared to a list of all genes to which insertions had been assigned, rather than just those to which significant CISs had been assigned using MC simulations. Genes could not be identified for 26 CISs, and these were assigned to genes manually, by observing insertions in the context of genes using the Ensembl DAS track (see Section 2.6). 102 CISs were assigned to more than 1 gene, and these were also assessed manually to determine whether one gene could be removed from the list. 14 CISs were removed where all insertions mapped to the same genomic coordinates, as these are likely to be artefacts. The final dataset comprised 630 CISs assigned to 608 genes. 30 CISs were associated with more than 1 gene, and 37 genes contained more than 1 CIS.

The lists of genes generated by the KC and MC methods were compared to the list of mouse orthologues of known cancer genes (see Section 2.2.3) and the Matthew's correlation coefficient (MCC) was calculated.

$$MCC = \frac{TP \times TN - FP \times FN}{\sqrt{((TP + FP)(TP + FN)(TN + FP)(TN + FN))}}$$

TP is the number of cancer genes in the candidate cancer gene list (true positives), FP is the number of non-cancer genes in the list (false positives), TN is the number of noncancer genes not in the list (true negatives), and FN is the number of cancer genes not in the list (false negatives). Genes that were not in the list were calculated as all 18,017 mouse genes with human orthologues, as identified in Ensembl version 48, minus those in the list. MCC is used in machine learning as a measure of the quality of a prediction and it takes into account the counts of true and false positives and negatives to generate a single number that can be compared across predictions. The candidate cancer gene lists generated from the retroviral screen are expected to contain known cancer genes, and these can therefore be used as a measure of the quality of the list. MCC is a more useful measure than accuracy or coverage alone, especially when comparing lists of different lengths. For example, a short list may have high accuracy but low coverage, and for a longer list, the reverse may be true. MCC returns a value between -1 and +1, where +1 is a perfect prediction (i.e. in this case, the list contains all known cancer genes and no noncancer genes), 0 is a random prediction, and -1 is an inverse prediction. The MCC score, plus accuracy and coverage, for the KC list and each MC list are shown in Table 2.7A (page 98). All MCCs generated in this analysis are positive but are very small because many of the genes are not known cancer genes. The KC list had the lowest coverage but the highest accuracy, and achieved the highest MCC score. As expected, the MC list generated using a *Efr* of 0.001 achieved a higher MCC score than the *Efr*=0.01 list since there should be 10-fold reduction in the number of randomly occurring CISs, and the higher accuracy more than compensated for the lower coverage. The list containing all genes that were assigned to insertions, rather than just those with statistically significant insertions, achieved the highest coverage but performed worst overall. Despite the fact that the list of known cancer genes is incomplete, measurement of the MCC score enables direct comparison of the gene lists and is likely to be meaningful. In light of these findings, the KC list was judged to be most accurate and was chosen for the cross-species comparative analyses performed in Chapter 5.

In order to gain an impression of whether the correct genes had been chosen for the KC CISs, known oncogenes were identified within the list of genes flanking each CIS. 37 oncogenes had been chosen, while 16 had not. Of the unselected oncogenes, 3 were genes nearest to the CIS, and 13 were further away. The insertions around these genes were analysed in the context of the mouse genome in Ensembl contigview. Only one oncogene nearest the CIS and one further away appeared to have been wrongly assigned, and for one additional nearest oncogene, it appeared that both this gene, and the correctly assigned gene might be mutational targets. The list of CIS genes was modified to include these three genes (*Rhoh*, *Cbl* and *Ccnd2*), but the results of the MCC comparison and analysis of the distribution of insertions suggest that, by and large, the most likely candidate gene has been selected.

Of the 39 cross-scale *Sleeping Beauty* CISs identified by the KC method, 36 were also present in the *Efr*=0.01 MC list. The remaining 3 were a long way from the nearest gene,

further than the 100 kb limit used in the MC simulations. As *Sleeping Beauty* has low enhancer activity, these are likely to be non-oncogenic insertions that have preferentially inserted into a particular genomic region, or are mutating a gene that has not been identified in the Ensembl gene build. 79 genes from the *Efr*=0.01 MC list were not identified by the KC method. 5 KC genes were missing from the MC list generated using an *Efr* of 0.001, and in all cases the CIS was greater than 100 kb from the gene, and 27 of the *Efr*=0.001 MC genes were not identified by KC. The KC method is designed primarily for large datasets and may therefore miss a significant proportion of biologically relevant CISs in the *Sleeping Beauty* dataset. However, the MCC score is highest for the candidate gene list generated using the KC method, and other lists follow the same pattern as the corresponding lists generated from the retroviral dataset (Table 2.7B, page 98).

21 Sleeping Beauty CISs were identified using the KC-based framework with a kernel width of 30 kb and P=0.05 (Appendix B1), but 5 were situated close to the transposon array on chromosome 1, and 4 were situated close to the array on chromosome 15. These were removed from the list because they are likely to result from "local hopping" of the transposon (see Section 1.4.2.2.1). The T2/Onc splice acceptor and splice donor sequences are derived from exon 2 of the *En2* gene and exon 1 of the *Foxf2* gene, respectively (Collier *et al.*, 2005). Statistically significant CISs were identified in both these genes, and the insertions were found to cluster around the splice junctions used to construct T2/Onc (Figure 2.12). These CISs most likely represent artefacts resulting from the mapping of T2/Onc sequences, rather than flanking genomic sequences, to the mouse genome, and they were removed from the dataset. This leaves just 10 CISs and so, for the purposes of comparison with the retroviral dataset, discussed in Chapter 3, the more inclusive MC lists of candidate cancer genes were also used.

2.10.3 Final set of candidate genes

Following a survey of the candidate genes identified from the retroviral screen by the kernel convolution-based method, it became clear that some insertions mapped to exactly the same coordinates. This is unlikely to occur by chance, except where mutation of a very localised region of a gene is required for oncogenesis. There were 26 animals from which 2 tumours had been collected, 70 from which 3 tumours had been collected, and 3 from which 4 tumours had been collected. Where a tumour has spread to a different site,

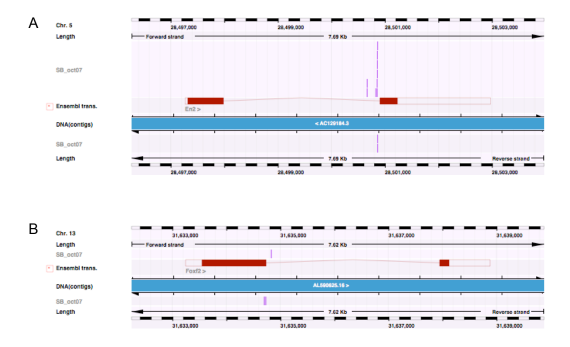


Figure 2.12. Insertions in En2 (A) and Foxf2 (B) are located at the splice junctions used to construct the T2/Onc transposon and are contaminating sequences. Insertions are shown as pink lines in the context of the Ensembl gene, shown in red. Insertions above and below the blue line are in the sense and antisense orientation, respectively.

a high proportion of insertions may be shared by both the original tumour and the secondary tumour, and this will influence the identification of significant CISs. Therefore, where 2 or more insertions from different tumours in the same animal occurred within 50 base pairs of one another, all but 1 of the insertions were removed from the dataset. A distance of 50 bp was chosen by counting the number of insertions that cooccurred within varying distances, and taking the distance at which the number levelled off. This reduced the dataset to 22,180 insertions. In addition, there is the possibility that insertions may map to the same position because of contamination during PCR. Therefore, if there were 2 or more sites in the genome where insertions from 2 tumours co-occurred within 10 bp, 1 of the co-occurring insertions was removed from the dataset at each location. A 10 bp window was used since it allows for a small amount of variation in the alignment of sequences using SSAHA2 (see Section 2.5), without significantly risking the removal of insertions that happen to fall into dense CISs. If the co-occurrence has resulted from aerosol contamination, it is assumed to be more likely that the insertion represented by the fewest number of reads is the contaminant and, therefore, in each case, this insertion was removed. The kernel convolution-based approach was applied to the final dataset of 20,114 insertions, and this resulted in 439 candidate cancer genes, of which 416 had a single CIS, 18 had 2 CISs, 2 had 3 CISs, 2 had 4 CISs and 1 had 5 CISs. The total number of CISs was 447, of which 24 were assigned to 2 genes. The CISs and associated genes are shown in Appendix B2.

2.11 Discussion

The aim of this chapter was to generate a reliable list of candidate cancer genes from insertional mutagenesis screens performed using the retrovirus MuLV and the *Sleeping Beauty* transposon T2/Onc. In order to maximise the number of insertions that could be identified within tumours, SSAHA2 was optimised to enable the mapping of as many reads as possible. The high number of unmapped reads was found to result from a high proportion of very short reads, especially in the *Sleeping Beauty* dataset, as well as reads containing genomic DNA of low complexity or low quality and reads that contained contaminating vector sequences. A small proportion may also result from errors in the mouse genome assembly. There did not seem to be any significant advantage in using BLASTN to map the reads, and as SSAHA2 is a faster algorithm, it is a good choice for mapping large numbers of reads. However, a possible alternative to SSAHA2 for future screens is the BLAST-Like Alignment Tool, BLAT (Kent, 2002). The UCSC Genome

Browser website (http://genome.ucsc.edu/, Kent *et al.*, 2002) uses BLAT to map users' sequences to the genome, and, because of its high speed and accuracy, BLAT has recently replaced BLAST as the default DNA search algorithm on the Ensembl website. Nevertheless, given the modest differences between SSAHA2 and BLAST (Altschul *et al.*, 1990), it is likely that BLAT would also perform similarly since the short, repetitive and low quality non-mapping reads can only be mapped at the expense of accuracy.

The reads were filtered to remove those in which the genomic DNA did not appear to represent the true location of the insertion. A gap between the genomic and retroviral DNA can result from low quality sequencing or the presence of unrelated DNA fragments within the clone, and efforts were made to retain low quality reads, whilst removing contaminating chimeric sequences. Comparisons between PCRs performed on the same tumours suggested that using more restriction enzymes and increasing the sequencing depth should increase the number of insertions that can be identified. Advances in sequencing technologies, such as 454 sequencing (see Section 1.4.2.1.2), will enable the use of more restriction enzymes and a greater depth of sequencing at a lower cost per read, thereby facilitating the identification of a higher proportion of insertions.

Identifying the genes that are most likely to have been mutated by insertions is hampered by the presence of enhancer insertions that can act at long range. Analysis of the distribution of insertions around mouse genes, and in particular, known cancer genes, suggested that the optimal distance is around 500-600 bp upstream, although the distance can be much greater, e.g. enhancer mutations can act as far as 270 kb downstream of the *Myc* promoter (Lazo *et al.*, 1990). It appears that downstream insertions are less likely to be oncogenic, although those in the sense orientation with respect to upstream genes may be more likely to contribute to oncogenesis. Enhancers can act over large distances via chromatin loop interactions, and they may therefore affect the activity of multiple genes (Uren *et al.*, 2005). However, analysis of the distribution of insertions around cancer genes suggests that, in general, enhancer insertions affect the promoters of the nearest, flanking genes.

Two approaches, Monte Carlo simulations (Suzuki *et al.*, 2002) and a kernel convolutionbased statistical framework (de Ridder *et al.*, 2006), have been used to identify statistically significant CISs in the retroviral and transposon screens. Known cancer genes can be used as a partial set of true positives to evaluate candidate cancer genes in the vicinity of CISs, and Matthew's Correlation Coefficient was used to show that the kernel convolution-based framework gives the most reliable set of candidate cancer genes. The final set of candidates generated from the *Sleeping Beauty* screen comprises just 10 genes, reflecting the small size of the initial dataset and problems in mapping the reads. 439 candidates were identified from the MuLV screen, thereby supporting the theory that many genes contribute to tumourigenesis. The candidate cancer genes are analysed and characterised in Chapter 3.

Chapter 3 Analysis of mouse candidate cancer genes identified by insertional mutagenesis

3.1 Introduction

This chapter describes methods used to characterise the mouse candidate cancer genes identified by retroviral and transposon-mediated insertional mutagenesis. The integration of other cancer-associated datasets provides a means of filtering the genes to identify the strongest candidates for a role in tumourigenesis (see Section 1.5). Importantly, human cancer-associated datasets can be used to assess the relevance of insertional mutagenesis to human cancer. Analysis of Gene Ontology terms and gene pathways, as well as the identification of genes with binding sites for transcription factors relevant to cancer, can help to define the cancer pathways in which candidate genes may act. Comparative analyses between the mouse candidate genes and other cancer-related datasets are described in Section 3.2. The mutational profile varies between insertional mutagens, and is affected by insertional bias and the mechanisms by which the mutagen disrupts genes (see Section 1.4.2.1.1). Genes that are identified by multiple mutagens are strong candidates for a role in tumourigenesis. The candidate genes identified using MuLV and the *Sleeping Beauty* (SB) transposon T2/Onc are compared in Section 3.3.

The distribution of insertions in and around candidate cancer genes gives an indication of the likely mechanisms of mutagenesis (see Section 1.4.2.1.1) and therefore provides an insight into the structure and function of mutant oncoproteins. In Section 3.4.1, the distribution of intragenic insertions within candidates from the MuLV screen is explored, and genes are classified according to their predicted mutation type. The co-occurrence of both retroviral and transposon insertions within a localised region of a gene provides a strong indication that mutation within that region contributes to tumourigenesis. Therefore, in Section 3.4.2, the distribution of insertions in genes identified by both screens is used to predict the likely mechanisms of mutation. While it is clear that genes are frequently mutated by enhancer or promoter mutation or by premature termination of gene transcription, it is unclear whether the disruption of regulatory elements is a nalysis of insertions within regulatory features extracted from the Ensembl database.

Retroviral insertional mutagenesis identifies mainly oncogenes but it is also possible to identify tumour suppressor genes, and candidates are presented in Section 3.4.4. Finally, expression data for 18 MuLV-induced tumours is analysed in Section 3.4.5 in an attempt to confirm the deregulation of candidate genes.

Tumourigenesis is a multi-step process involving the co-operation of multiple cancer genes and pathways (see Section 1.2.3). Section 3.5 describes approaches for identifying co-operative cancer genes and presents a number of strong collaborations between genes identified in the retroviral screen. The work described in this chapter demonstrates the relevance of insertional mutagenesis to the study of human cancer, and identifies candidate cancer genes that warrant further investigation.

3.2 Comparative analyses between the insertional mutagenesis data and other cancer-related datasets

3.2.1 Description of the datasets

This section describes the datasets used for comparison with the candidate cancer genes identified by retroviral and transposon-mediated insertional mutagenesis. For all datasets where it was necessary to convert gene names to Ensembl identifiers, Ensembl BioMart (http://www.ensembl.org/biomart/index.html) was used. BioMart is a data mining tool that can be used to extract specific information from Ensembl for multiple genes simultaneously via a simple web interface. For all human datasets, mouse genes with human orthologues were also identified using Ensembl BioMart (version 48). The dataset of known cancer genes from the Cancer Gene Census (Futreal *et al.*, 2004) is described in Section 2.2.3.

3.2.1.1 The Retrovirus Tagged Cancer Gene Database (RTCGD)

mentioned in Section 1.4.2.1.2, As RTCGD (Akagi al., 2004; et http://rtcgd.abcc.ncifcrf.gov/) is a database that manages data from retroviral and transposon-mediated insertional mutagenesis screens. All candidate cancer genes in the database were obtained from the website on 01/11/07. In total, the database contained with MGI 537 unique (Mouse Genome Informatics, genes http://www.informatics.jax.org/) symbols identified from 512 retroviral CISs (25 CISs had been assigned to 2 genes). The MGI symbols were used to identify 544 mouse Ensembl genes. 16 genes had 2 Ensembl identifiers (e.g. *Akap13*, which, according to Ensembl, is duplicated in 2 adjacent copies) and 9 could not be identified. 55 genes were identified from 52 transposon CISs (3 had been assigned to 2 genes) and all but 2 had Ensembl gene identifiers.

3.2.1.2 The Catalogue of Somatic Mutations in Cancer (COSMIC)

COSMIC stores and displays somatic mutation information relating to human cancers that has been curated from published scientific literature (Forbes *et al.*, 2006, see also Section 1.3.1). The complete set of mutations in COSMIC version 35 (dated 04/02/08) was downloaded from the website ftp://ftp.sanger.ac.uk/pub/CGP/cosmic/data_export. The list comprised 52,678 mutations in 1,550 genes from 45,743 tumours of 38 cancer types. Unique HGNC (HUGO Gene Nomenclature Committee; http://www.genenames.org/) human gene symbols were converted to human Ensembl gene identifiers. 1,521 mouse Emsembl genes were identified as having a human orthologue with somatic mutations in COSMIC. Individual genes were also searched against the COSMIC database via the website http://www.sanger.ac.uk/genetics/CGP/cosmic/.

3.2.1.3 Human candidate cancer genes from Sjöblom et al. (2006)

This dataset, which is described in Section 1.3.1, comprises 121 candidate breast cancer genes and 69 candidate colon cancer genes. HGNC symbols were used to extract Ensembl identifiers for all genes, and 181 mouse orthologues were identified.

3.2.1.4 Transcription factor binding sites

Mouse genes with Nanog and Oct4 binding sites were extracted from Loh *et al.* (2006), while human genes with p53 binding sites were extracted from Wei *et al.* (2006). Both datasets are described in Section 1.3.5. Ensembl gene IDs were identified from MGI symbols and/or Entrez Gene (http://www.ncbi.nlm.nih.gov/sites/entrez?db=gene) accession numbers for genes in the Nanog and Oct4 datasets, and from HGNC symbols and/or RefSeq (http://www.ncbi.nlm.nih.gov/RefSeq/) accession numbers for genes in the S,006 Nanog binding sites, 2,408 were assigned to 1,923 mouse

Ensembl genes (i.e. some genes contained multiple binding sites), of which 1,889 encoded proteins or miRNAs. 817 mouse Ensembl genes, including 797 encoding proteins or miRNAs, were identified for 902 of the 1,083 Oct4 binding sites. 1,725 and 732 genes had human orthologues with Nanog and Oct4 binding sites, respectively. The p53 dataset contained 474 binding loci associated with human genes, of which 423 had Ensembl identifiers, resulting in 409 unique human Ensembl genes. 388 mouse Ensembl genes had a human orthologue with at least one binding site for p53.

3.2.1.5 Amplicons and deletions in paediatric acute lymphoblastic leukaemias

Regions of copy number change affecting more than one case of acute lymphoblastic leukaemia (ALL) were extracted from Mullighan *et al.* (2007; discussed in Section 1.3.3.3). In the publication, genomic coordinates were mapped to the human genome assembly NCBI 35, and these were therefore mapped across to NCBI 36 using the UCSC LiftOver tool (http://genome.ucsc.edu/cgi-bin/hgLiftOver). Overlapping amplicons and overlapping deletions were merged into single regions of copy number gain and loss, respectively. 1,905 mouse Ensembl genes contained a human orthologue within one of 8 non-overlapping amplicons, while 1,514 contained a human orthologue within one of 52 non-overlapping deletions.

3.2.1.6 Gene ontology (GO) terms

The GO project (http://www.geneontology.org) provides controlled vocabularies for describing genes and gene products in terms of their molecular function, their role in biological processes and their localisation to cellular components. The annotations assigned to genes and their products are known as GO terms. The public webserver g:Profiler (Reimand *et al.*, 2007, http://biit.cs.ut.ee/gprofiler/) was used to identify over-represented GO terms among the mouse candidate cancer genes, which were submitted as a list of mouse Ensembl gene identifiers. g:Profiler also identifies over-represented KEGG (http://www.genome.jp/kegg/) and REACTOME (http://www.reactome.org/) pathways, over-represented TRANSFAC (http://www.gene-regulation.com/) regulatory motifs, and miRNAs in miRBase (http://microrna.sanger.ac.uk/) for which target genes are over-represented among the candidate gene list.

3.2.2 Comparison with insertional mutagenesis data

The 1-tailed Fisher Exact Test was used to determine whether there was a significant overlap between candidate cancer genes identified by the MuLV and SB screens and those in other cancer-associated datasets. For comparison with the human datasets, the number of mouse orthologues of human genes was counted, whereas for comparison with the mouse Nanog and Oct4 binding site and RTCGD datasets, all mouse genes encoding proteins and miRNAs were counted. In total, there were 18,017 mouse Ensembl genes with human orthologues and 24,374 mouse genes that encoded proteins or miRNAs. All 439 of the candidate cancer genes identified in the MuLV screen using the kernel convolution (KC)-based approach (de Ridder *et al.*, 2006; see Section 2.10) encoded a protein or miRNA, and 384 had a human orthologue in Ensembl v48. All 10 of the *Sleeping Beauty* candidate genes were protein-coding, and 9 had human orthologues. For each significant CIS gene identified by the MuLV and SB screens, other cancer-associated datasets in which they also occurred are listed in Appendices C1 and C2, respectively.

Of the 439 CIS genes identified using MuLV, 118 (26.9%) were found among genes from other retroviral screens in the RTCGD database, and 6 (1.4%) were found among genes identified by transposon-mediated mutagenesis. This corresponds to a coverage of 21.7% and 11.3% of genes in the RTCGD database identified by retroviral and transposonmediated mutagenesis, respectively. The larger overlap with candidates identified in retroviral screens suggests that retroviruses and transposons have different mutational profiles (see also Section 3.3) but may also reflect the fact that there is more retroviral data available. However, a high proportion of candidates are unique either to this dataset or to the RTCGD database. This may in part reflect limitations in insertion site identification, such as the number of restriction enzymes used in linker-mediated PCR and the depth of sequencing. Because of the variety of methods used to detect significant CISs, there may also be a high number of false detections in the RTCGD database, since 53% of CISs did not reach the significance threshold when the KC-based approach was applied to the data in RTCGD (de Ridder et al., 2006). In addition, the RTCGD database contains genes that were identified by insertional mutagenesis on a range of genetic backgrounds and using a range of retroviral mutagens, and each mutagen and background may generate a different spectrum of candidates (see Section 3.5.1 for details on the identification of genotype-specific candidates). For example, Sox4 and Fgf3 are the 3rd

and 6^{th} most frequently mutated genes in RTCGD and yet they were not among the candidates identified in the MuLV screen. Almost all of the *Sox2* insertions were identified in mice with an AKxD or BXH2 strain background, while all of the *Fgf3* insertions were from a screen that used mouse mammary tumour virus (MMTV), rather than MuLV, as the retroviral mutagen.

Human orthologues of mouse candidate genes were enriched among oncogenes in the Cancer Gene Census (36 oncogenes, $P=7.88 \times 10^{-18}$) and the COSMIC database (69 genes, $P=1.36 \times 10^{-9}$). There were no recessive cancer genes among the candidates (P=1), demonstrating that the screen identifies predominantly oncogenes. Surprisingly, there were just 3 genes (*Lrrfip1*, *Nup214* and *Bcl11a*; P=0.74) that overlapped between the candidates of the retroviral screen and the candidates from Sjöblom *et al* (2006). This may reflect the fact that the Sjöblom dataset was an exon resequencing study of breast and bowel tumours exclusively, and it may be biased against genes mutated in lymphomas.

The 36 orthologues of mouse candidate cancer genes in the Cancer Gene Census were enriched for genes that are mutated in lymphoid tumours (31 genes, $P=2.66 \times 10^{-4}$). This suggests that the retroviral screen mainly identifies genes that are important in the development of lymphoid malignancies. There was also a slight enrichment of genes that are mutated by chromosomal translocation, although this was not significant (31 genes, P=0.067). A more significant association might be expected because translocation is a common mechanism of mutation in lymphoid cancers, and a number of genes that are frequently targeted by insertional mutagenesis are involved in translocations in human tumours. In addition, MuLV mutagenesis may have a similar effect to translocations, since it often changes the regulatory environment of a gene and/or produces truncated Chromosomal translocations, and leukaemias, lymphomas and oncoproteins. mesenchymal tumours, all of which frequently harbour translocations, are overrepresented in the Cancer Gene Census. This is partly because both translocation partners feature in the list of cancer genes, but also because, traditionally, cancer gene identification has been more frequently performed in these cancer types (Futreal, 2007). This may explain why the candidate cancer genes identified in the retroviral screen contain an over-representation of genes in the Cancer Gene Census, but that translocations are not over-represented among these candidates in the Census. Finally, there was an over-representation of known cancer genes that bear somatic mutations in human cancer (36 genes, P=0.0130). These findings demonstrate the efficacy of the MuLV retrovirus as a somatic mutagen that can be used to model the clonal evolution of human cancers, particularly those of lymphoid origin.

Human orthologues of mouse candidate cancer genes were significantly enriched among genes with p53 binding sites (14 genes, P=0.0394). The p53 pathway is important in tumourigenesis (see Section 1.2.6), and the identification of genes that act in this pathway provides further evidence that the screen has identified promising candidates for a role in cancer. It has been proposed that the CIS genes *Ptpre* and *Notch1* are upregulated by p53, while Nedd4l is downregulated (Wei et al., 2006). Ptpre is required for p53-induced differentiation of IW32 erythroleukaemia cells (Tang and Wang, 2000), while upregulation of Notch1 by p53 in human cancer cell lines contributes to cell fate determination (Alimirah et al., 2007). Nedd4l is overexpressed in human prostate cancer cells (Qi et al., 2003) and in the rare cutaneous T-cell lymphoma associated with Sézary Syndrome (Booken et al., 2008), suggesting that suppression by p53 inhibits cancer growth. There was also a significant enrichment of mouse candidate cancer genes among genes with Nanog (53 genes, $P=5.86 \times 10^{-4}$) and Oct4 (32 genes, $P=1.64 \times 10^{-5}$) binding sites. Nanog and Oct4 regulate self-renewal, pluripotency and differentiation of ES cells (see Section 1.3.5). 9 CIS genes have binding sites for both Nanog and Oct4 and these include Mycn, Il6st and Chd1, which are upregulated in human ES cells, mesenchymal stem cells and haematopoietic stem/progenitor cells, respectively (Kim et al., 2006a). *Il6st* (also known as gp130) is a key component of the signalling pathway required for the maintenance of embryonic stem cell pluripotency (Yoshida et al., 1994) and mouse haematopoietic stem cell function (Audet *et al.*, 2001). These results suggest that a significant proportion of candidates may be involved in tumour cell self-renewal, therefore providing support for the cancer stem cell hypothesis, described in Section 1.2.3.2.

Mouse candidate genes with human orthologues were also over-represented in regions of copy number gain (54 genes, P=0.0180) and copy number loss (47 genes, $P=5.82 \times 10^{-3}$) in paediatric ALL. CIS genes that were deleted in ALL included *Lef1*, *Ikzf1*, *Ikzf3*, *Etv6*, *Elf1* and *Erg*, while those that were amplified included *Runx1*, *Myb* and *Ahi1*. This suggests that genes that are mutated by insertional mutagenesis, and contribute to mouse tumourigenesis, may also be mutated by copy number changes in human cancers. However, the overlapping genes are implicated in B-cell development and differentiation,

which are disrupted in human B-progenitor ALL and in MuLV-induced murine lymphomagenesis. It therefore remains to be seen whether CIS genes significantly overlap with regions of copy number change in other human cancers, and this is addressed in Chapters 4 and 5. The CIS genes may help to narrow down the candidates in regions of copy number change in the ALL dataset. For example, the deleted region on human chromosome 16q22.1 contains 11 genes, but the mouse orthologue of only 1 of these genes (FAM65A) is targeted by MuLV in insertional mutagenesis and therefore represents a putative target for deletion in ALL. Table 3.1 provides a list of the regions that are amplified and deleted in ALL and the CIS genes within these regions.

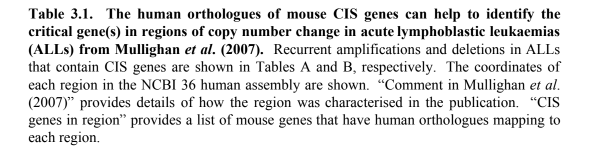
The candidate cancer genes were over-represented among genes in the KEGG pathways associated with acute and chronic myeloid leukaemia ($P=2.14\times10^{-13}$ and $P=1.75\times10^{-7}$, respectively) and Jak-STAT signalling, and in the T cell receptor signalling KEGG and REACTOME pathways ($P=1.35\times10^{-5}$ and $P=1.96\times10^{-6}$, respectively). This is encouraging, since the genes are candidates for a role in lymphomagenesis. However, genes were also over-represented in the endometrial cancer KEGG pathway ($P=7.14\times10^{-4}$), demonstrating that some of the candidates (including *Pik3cd*, *Pik3r5*, *Akt1*, *Lef1*, *Myc*, *Ccnd1* and *Tcf7*) also contribute to other cancer types. Over-represented GO terms are listed in Table 3.2. These include terms related to the development, differentiation and proliferation of B- and T-cells, reflecting the lymphoid origin of the mouse tumours, and terms specifically related to cancer, such as cell proliferation, apoptosis, angiogenesis, cell motility and kinase activity.

Four transcription factor binding sites from the TRANSFAC database were also overrepresented among the candidate genes. The most significant was the MAZ (Mycassociated zinc finger protein) binding matrix (TF:M00649, $P=1.49\times10^{-8}$), which binds the MAZ transcription factor. MAZ interacts with MYC and histone deacetylases, and *MAZ* overexpression drives expression of the oncogene *PPAR* γl in human breast cancer cells (Wang *et al.*, 2008). It is also overexpressed in acute myeloid leukaemia (Greiner *et al.*, 2000) and in the terminal phase of chronic myeloid leukaemia (Daheron *et al.*, 1998). The second most significant binding matrix was TF:M01104 ($P=2.51\times10^{-6}$), which binds the mouse Movo-b zinc finger protein. This protein is highly expressed in the mouse testis (Unezaki *et al.*, 2004), and has no known role in tumourigenesis, but has been shown to be involved in vascular angiogenesis in the developing embryo (Unezaki *et al.*, 2007). Finally, binding matrices for transcription factors LRF (leukaemia/lymphoma

			Comment in	
Chromosome	Start (bp)	End (bp)	Mullighan et al.	CIS genes in region
1	127000000	247249719	719 genes telomeric of	Mef2d, Nid1, Slamf6, Cd48, Anp32e,
			PBX1	Lyst, Btg2, Ptprc, Mixl1, Ccdc19, Sell,
				Ppp2r5a, Zbtb7b, Rorc, Slamf7, Mcl1,
				Itpkb, AI848100, Rcsd1, 5730559C18Rik,
				Irf2bp2
2	1	31987853	235 genes	D12Ertd553e, Mycn
6	1	26216000	190 genes	Irf4, Exoc2, Rreb1, Sox19
6	135556000	135714000	MYB, MIRN548A2,	Myb, Ahi1
			AHI1	
9	6000000	140273252	155 genes telomeric of	Nup214, Phyhd1, Sema4d, Gadd45g,
			ABL1	Ccrk, Eng, Gfi1b, Ak1, Egfl7, Notch1,
				Coro2a, A2AN91_MOUSE, Akna,
				A130092J06Rik
10	1	40290000	All 10p	Cugbp2, Map3k8, Il2ra, Zfp438
21	32896000	35199000	33 genes including	Runx1, Ifnar1
			Runx1	
22	1	21888000	277 genes telomeric of	Bid, Tuba8, BC030863, Cecr5, Vpreb2
			BCR	

В

			Comment in	
Chromosome	Start (bp)	End (bp)	Mullighan et al.	CIS genes in region
2	232347739	242951149	124 genes	Lrrfip1
4	109254845	109303845	LEF1	Lef1
5	163535000	180857866	172 genes	C330016O10Rik, Mgat1
7	1	58058273		Stard3nl, Mafk, Ikzf1, Mad1l1, Lfng, Hibadh, Hoxa7, Sdk1, 3110082117Rik,
				Jazf1
9	1	50600000		Cd72, Anxa2, Dock8
11	117882000	118379000	16 genes distal to MLL	Treh, Bcl9l
12	11694055	11939588	ETV6	Etv6
13	40453000	40484000	ELF1	Elf1
13	47885000	47968000	RB1	Rcbtb2
16	66116000	66423000	FAM65A, CTCF, RLTPR,	2310066E14Rik (FAM65A)
			ACD, PARD6A, C16orf48, LOC388284, GFOD2, RANBP10, TSNAXIP1, CENPT	
17	1	18837000	383 genes	Lgals9, AA536749, Prr6, Pik3r5, Ntn1,
	05405000			Slc43a2, Ovca2, Smg6, Rtn4rl1
17	35185000	35230000		Ikzf3
19	229000	1531000	63 genes telomeric to <i>TCF3</i>	Ptbp1, Arid3a, Midn
20	27000000	62435964	All 20q	Bcl2l1, Serinc3, Stk4, Ndrg3, Sla2, Ncoa3, Ppp1r16b, Prkcbp1, Zfp217
21	38706000	38729000	ERG	Erg



P-value 8.90E-14 4.89E-14 3.29E-14	CIS			
4.89E-14	genes 167	GO ID GO:0065007	Ontology BP	GO term GO term
2 20E-14	155	GO:0050789	BP	regulation of biological process
	64	GO:0048518	BP	positive regulation of biological process
3.98E-14	148	GO:0050794	BP	regulation of cellular process
2.07E-12 1.01E-07	57 19	GO:0048522 GO:0008284	BP BP	positive regulation of cellular process positive regulation of cell proliferation
1.48E-06	27	GO:00031325	BP	positive regulation of cellular metabolic process
2.18E-06	10	GO:0050867	BP	positive regulation of cell activation
1.51E-05	6	GO:0045787	BP	positive regulation of cell cycle
1.52E-05	79	GO:0019219 GO:0051252	BP BP	regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolic process
5.73E-06 3.83E-05	76 76	GO:0051252 GO:0045449	BP	regulation of RNA metabolic process regulation of transcription
2.77E-05	73	GO:0006355	BP	regulation of transcription, DNA-dependent
4.32E-07	26	GO:0006357	BP	regulation of transcription from RNA polymerase II promoter
1.46E-11	30	GO:0051726	BP	regulation of cell cycle
1.29E-10	54	GO:0048523	BP	negative regulation of cellular process
1.60E-08 2.47E-07	28 13	GO:0042127 GO:0050865	BP BP	regulation of cell proliferation regulation of cell activation
4.84E-07	30	GO:0009966	BP	regulation of signal transduction
2.71E-06	89	GO:0031323	BP	regulation of cellular metabolic process
1.51E-05	9	GO:0051270	BP	regulation of cell motility
4.54E-05	80	GO:0010468	BP	regulation of gene expression
8.52E-11 8.80E-08	57 44	GO:0048519 GO:0050793	BP BP	negative regulation of biological process regulation of developmental process
3.09E-10	28	GO:0051094	BP	positive regulation of developmental process
5.83E-08	30	GO:0043067	BP	regulation of programmed cell death
2.86E-08	20	GO:0043068	BP	positive regulation of programmed cell death
2.47E-08	20	GO:0043065	BP	positive regulation of apoptosis
2.38E-06	15	GO:0006917	BP	induction of apoptosis
2.38E-06 4.87E-08	15 30	GO:0012502 GO:0042981	BP BP	induction of programmed cell death regulation of apoptosis
6.43E-06	21	GO:0051093	BP	negative regulation of developmental process
2.18E-07	17	GO:0002682	BP	regulation of immune system process
2.47E-07	13	GO:0002694	BP	regulation of leukocyte activation
1.26E-07	13	GO:0051249	BP	regulation of lymphocyte activation
6.55E-09 1.24E-06	13 5	GO:0050863 GO:0050854	BP BP	regulation of T cell activation regulation of antigen receptor-mediated signaling pathway
1.24E-06 1.76E-05	4	GO:0050856	BP	regulation of T cell receptor signaling pathway
3.52E-06	8	GO:0002683	BP	negative regulation of immune system process
1.24E-05	13	GO:0002684	BP	positive regulation of immune system process
2.18E-06	10	GO:0002696	BP	positive regulation of leukocyte activation
1.69E-06 1.31E-07	10 10	GO:0051251 GO:0050870	BP BP	positive regulation of lymphocyte activation positive regulation of T cell activation
3.40E-05	6	GO:0050870 GO:0042102	BP	positive regulation of T cell activation positive regulation of T cell proliferation
2.43E-06	23	GO:0051239	BP	regulation of multicellular organismal process
9.53E-06	89	GO:0019222	BP	regulation of metabolic process
2.35E-06	27	GO:0009893	BP	positive regulation of metabolic process
2.86E-05	9	GO:0040012	BP BP	regulation of locomotion
1.29E-12 6.04E-12	27 25	GO:0001775 GO:0045321	BP	cell activation leukocyte activation
6.15E-12	24	GO:0046649	BP	lymphocyte activation
1.13E-11	19	GO:0042110	BP	T cell activation
3.85E-05	9	GO:0046651	BP	lymphocyte proliferation
2.29E-05	8	GO:0042098	BP	T cell proliferation
2.71E-11	47	GO:0002376	BP BP	immune system process
1.21E-05 3.60E-05	6 5	GO:0001776 GO:0002260	BP	leukocyte homeostasis lymphocyte homeostasis
2.89E-05	4	GO:0043029	BP	T cell homeostasis
8.96E-10	41	GO:0007049	BP	cell cycle
9.55E-10	61 75	GO:0007242	BP BP	intracellular signaling cascade
1.96E-09 1.96E-09	75	GO:0048869 GO:0030154	BP	cellular developmental process cell differentiation
1.15E-10	19	GO:0002521	BP	leukocyte differentiation
	15	GO:0030098	BP	lymphocyte differentiation
3.10E-09		00.0000000		T cell differentiation
3.10E-09 6.62E-06	9	GO:0030217	BP	
3.10E-09 6.62E-06 5.95E-09	103	GO:0030217 GO:0032502	BP	developmental process
3.10E-09 6.62E-06 5.95E-09 3.25E-09	103 80	GO:0030217 GO:0032502 GO:0048856	BP BP	developmental process anatomical structure development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08	103 80 39	GO:0030217 GO:0032502 GO:0048856 GO:0016265	BP BP BP	developmental process anatomical structure development death
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05	103 80	GO:0030217 GO:0032502 GO:0048856	BP BP	developmental process anatomical structure development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09	103 80 39 45 84 72	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731	BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11	103 80 39 45 84 72 27	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731 G0:0002520	BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11 4.63E-12	103 80 39 45 84 72 27 27	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731 G0:0002520 G0:0048534	BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoletic or lymphoid organ development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11	103 80 39 45 84 72 27	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731 G0:0002520	BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoiesis
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11	103 80 39 45 84 72 27 27 27 25	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731 G0:00048534 G0:0048534 G0:0030097	BP BP BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoletic or lymphoid organ development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.55E-08 1.97E-05	103 80 39 45 84 72 27 27 27 27 25 62 34 7	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0007275 G0:0048731 G0:0002520 G0:0048534 G0:003097 G0:0048534 G0:0008283 G0:00050673	BP BP BP BP BP BP BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopolesis organ development cell proliferation epithelial cell proliferation
3.10E-09 6.62E-06 5.95E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.97E-05 3.85E-05	103 80 39 45 84 72 27 27 27 25 62 34 7 9	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:0007275 G0:0048731 G0:0007250 G0:0048534 G0:003250 G0:0048513 G0:0008283 G0:0050673 G0:0032943	BP BP BP BP BP BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoiesis organ development cell proliferation epithelial cell proliferation monouclear cell proliferation
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.56E-08 1.97E-05 3.85E-05 1.59E-07	103 80 39 45 84 72 27 27 25 62 34 7 9 53	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:00048731 G0:0002520 G0:0048534 G0:00048534 G0:00048513 G0:00048513 G0:00048513 G0:00050673 G0:0032943 G0:0032943 G0:00348468	BP BP BP BP BP BP BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoiesis organ development cell proliferation epithelial cell proliferation mononuclear cell proliferation cell development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.55E-08 1.97E-05 3.85E-05 1.59E-07 4.15E-08	103 80 39 45 84 72 27 25 62 34 7 9 53 39	G::0030217 G::0032502 G::004856 G::0016265 G::0009653 G::0048731 G::0048731 G::0048531 G::0005283 G::0048513 G::0005283 G::0005283 G::0005283 G::0032943 G::0032943 G::0032943 G::0032943	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoietis organ development cell proliferation epithelial cell proliferation cell development cell development
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 2.88E-09 2.88E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-08 1.97E-05 3.85E-05 1.59E-07 4.15E-08 1.67E-08	103 80 39 45 84 72 27 27 25 62 34 7 9 53	G0:0030217 G0:0032502 G0:0048856 G0:0016265 G0:0009653 G0:00048731 G0:0002520 G0:0048534 G0:00048534 G0:00048513 G0:00048513 G0:00048513 G0:00050673 G0:0032943 G0:0032943 G0:00348468	BP BP BP BP BP BP BP BP BP BP BP BP BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoiesis organ development cell proliferation epithelial cell proliferation monouclear cell proliferation cell development cell death programmed cell death
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 2.88E-09 2.88E-09 2.88E-09 2.88E-09 1.80E-11 5.95E-09 1.56E-08 1.59E-07 4.15E-08 1.24E-08 1.24E-08	103 80 39 45 84 72 27 25 62 34 7 9 53 39 39 39 39 39	G::0030217 G:0032502 G:0048856 G::0016265 G::0009653 G:0007275 G::0048534 G::0002520 G:0048534 G::003097 G::0048513 G::003097 G::0048513 G::0050673 G::0050673 G::0050673 G::0050673 G::0050673 G::0048468 G::0008219 G::0012501 G::0006915 G::0006915	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoiesis organ development cell proliferation epithelial cell proliferation monouclear cell proliferation cell development cell development angiogenesis
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.97E-05 3.85E-05 3.85E-05 1.55E-08 1.45E-08 1.67E-08 1.46E-05 1.54E-05	103 80 39 45 84 72 27 25 62 34 7 9 53 39 39 39 39 39 31 31 78	G::0030217 G::0032502 G::0048856 G::0016265 G::009653 G::0048731 G::002520 G::0048534 G::0048534 G::0048533 G::0048513 G::0048513 G::0008219 G::0008218 G::0008219G::0008219 G::0008219G::0008219 G:	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopolesic or lymphoid organ development hemopolesis organ development cell proliferation epithelial cell proliferation cell development cell development cell death programmed cell death apoptosis angiogenesis macromolecule metabolic process
3.10E-09 6.62E-06 5.95E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 1.80E-11 5.95E-09 1.80E-11 5.95E-09 1.56E-08 1.97E-05 3.85E-05 1.55E-07 4.15E-08 1.67E-08 1.24E-08 1.24E-08 1.24E-08 1.24E-08 1.24E-08	103 80 39 45 84 72 27 27 27 27 27 27 27 27 27 27 27 27	G::0030217 G:0032502 G:0048856 G:0016265 G:0009653 G:0048731 G:0002520 G:0048513 G:0048513 G:0030873 G:0048513 G:0005073 G:0048513 G:0005073 G:0048513 G:0005073 G:0005073 G:0005073 G:0005073 G:0004816 G:0006915 G:0001525 G:00043170 G:0043283	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoietis organ development cell proliferation epithelial cell proliferation cell development cell death programmed cell death apoptosis angiogenesis macromolecule metabolic process biopolymer metabolic process
3.10E-09 6.62E-06 6.52E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.56E-08 1.56E-08 1.57E-08 1.24E-08 1.24E-08 1.24E-08 1.24E-05 3.08E-05 1.57E-05	103 80 39 45 84 72 27 27 27 25 62 34 7 9 53 39 39 39 39 39 13 178 63	G::0030217 G:0032502 G:0048856 G:0016265 G:009653 G:0009275 G:0048531 G:0002520 G:0048531 G:0048533 G:0048533 G:0048533 G:0032943 G:00032943 G:00032943 G:0008219 G:0008219 G:0005255 G:0004525 G:0004525 G:0043170 G:0043283 G:0043283	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoietic or lymphoid organ development hemopoiesis organ development cell opoliferation cell opoliferation cell development cell death apoptosis angiogenesis macromolecule metabolic process biopolymer metabolic process biopolymer modification
3.10E-09 6.62E-06 5.95E-09 4.15E-08 2.83E-05 6.62E+09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.59E-09 1.59E-09 1.59E-07 4.15E-08 1.67E-08 1.24E-08 1.24E-08 3.08E-09 1.57E-05 7.715E-05	103 80 39 45 84 72 27 27 27 26 62 34 7 9 53 39 39 39 39 39 31 31 78 153 62	G::0030217 G:0032502 G:0048856 G:0016265 G:0016265 G:0048731 G:0002520 G:0048534 G:0002520 G:0048513 G:0008283 G:0005073 G:0048453 G:0005073 G:00048513 G:0005073 G:0005073 G:00048463 G:0005075 G:00048463 G:0001525 G:0001525 G:0001525 G:00043170 G:00043170 G:00043170 G:00043170 G:00043170	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoletic or lymphoid organ development hemopoletic or lymphoid organ development hemopoletic or lymphoid organ development epithelial cell proliferation cell development cell development cell development cell development apoptosis angiogenesis macromolecule metabolic process biopolymer metabolic process biopolymer modification protein modification process
3.10E-09 6.62E-06 6.52E-09 3.25E-09 4.15E-08 2.83E-05 6.25E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.56E-08 1.56E-08 1.57E-08 1.24E-08 1.24E-08 1.24E-08 1.24E-05 3.08E-05 1.57E-05	103 80 39 45 84 72 27 27 27 25 62 34 7 9 53 39 39 39 39 39 13 178 63	G::0030217 G:0032502 G:0048856 G:0016265 G:009653 G:0009275 G:0048531 G:0002520 G:0048531 G:0048533 G:0048533 G:0048533 G:0032943 G:00032943 G:00032943 G:0008219 G:0008219 G:0005255 G:0004525 G:0004525 G:0043170 G:0043283 G:0043283	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoiesic or lymphoid organ development hemopoiesis organ development cell opoliferation cell opoliferation cell development cell death apoptosis angiogenesis macromolecule metabolic process biopolymer medabolic process biopolymer modification
3.10E-09 6.62E-06 5.95E-09 4.15E-08 2.83E-09 2.83E-05 6.62E-09 2.88E-09 1.80E-11 4.63E-12 2.02E-11 5.95E-09 1.56E-08 1.97E-05 3.85E-05 1.54E-05 1.54E-05 3.08E-09 1.57E-05 3.08E-09 1.57E-05 3.48E-05 3.4	103 80 39 45 84 72 27 25 62 34 7 9 39 39 39 39 39 39 133 178 153 62 54 153 62 54	G::0030217 G::0032502 G::0048856 G::0016265 G::0009653 G::0007275 G::0048513 G::0002520 G::0048513 G::00048513 G::00048513 G::00048513 G::00048513 G::0005073 G::0004816 G::0005073 G::00048468 G::000691525 G::00043120 G::00043163 G::0004367 G::00043687 G::000430887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::00043887 G::0004388 G::0004388 G::0004388 G::0004388 G::0004388 G::00043887 G::0004888 G::0004888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::00048888 G::000488888 G::000488888 G::000488888 G::000	BP	developmental process anatomical structure development death anatomical structure morphogenesis multicellular organismal development system development immune system development hemopoletic or lymphoid organ development hemopoletics organ development cell proliferation epithelial cell proliferation cell development cell development apoptosis angiogenesis macromolecule metabolic process biopolymer metabolic process biopolymer metabolic process biopolymer metabolic process biopolymer modification protein modification of transcription from RNA polymerase II promoter B cell differentiation positive regulation of transcription from RNA polymerase II promoter b cell forentiation
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Table 3.2. Over-represented GO terms among CIS genes identified using MuLV. "CIS genes" is the number of CIS genes annotated for each term. The ontologies shown are biological process (BP) and molecular function (MF). Terms are staggered to show GO term hierarchies, with terms of equivalent hierarchy being listed in order of decreasing significance. Terms associated with T- and B-cells are shown in blue.

related factor; TF:M01100) and VDR (vitamin D receptor; TF:M00444) were also significantly over-represented among candidate genes ($P=1.38\times10^{-5}$ and $P=1.47\times10-5$, respectively). LRF is a master regulator of oncogenesis that directly represses transcription of the tumour suppressor gene $p19^{ARF}$ (Maeda *et al.*, 2005) and plays an essential role in determining B- versus T-cell fate (Maeda *et al.*, 2007). VDR is also widely implicated in human tumourigenesis (for review, see Thorne and Campbell, 2008). Specific analysis of over-represented GO terms for the genes associated with each transcription factor showed that all were enriched for terms relating to the cell cycle and KEGG pathways for acute and chronic myeloid leukaemia. Only candidates associated with the Movo-b binding site were enriched for genes with protein serine/threonine kinase activity ($P=2.21\times10^{-5}$), suggesting that Movo-b may play an important role in regulating protein kinases. Likewise, only genes with Lrf binding sites were enriched for terms associated with apoptosis, suggesting that Lrf may also repress other tumour suppressor genes, including *Wwox* and *Trp53inp1*.

The output from g:Profiler also showed that candidate genes were over-represented among the predicted targets of 3 miRNAs: mmu-miR-449b (P=4.31x10⁻⁵), mmu-miR-449c ($P=9.69 \times 10^{-6}$) and hsa-miR-565 ($P=6.29 \times 10^{-5}$), which suggests that these miRNAs may play an important role in regulating genes involved in tumourigenesis. The list of candidates also included 5 genes that encode miRNAs: mmu-miR-142, mmu-miR-17, mmu-miR-802, mmu-miR-181c and mmu-miR-23a. miRNAs play an important role in haematopoiesis, and miRNA deregulation has been widely observed in leukaemias and lymphomas (for review, see Garzon and Croce, 2008). The human orthologues of mmumiR-142 and mmu-miR-181 are both implicated in the regulation of mammalian haematopoiesis, since *hsa-miR-142* is at a translocation site within a case of aggressive Bcell leukaemia, while B-cell-specific hsa-miR-181 promotes B cell differentiation (Chen and Lodish, 2005). Deregulated miRNAs also contribute to cancer in other cancer types. hsa-miR-23a, the human orthologue of mmu-miR-23a, is upregulated in human hepatocellular carcinomas (Kutay et al., 2006), while the human orthologue of mmu-miR-17 is implicated as an oncogene in a range of cancers, and is discussed further in Section 4.5.2.1.2.

Of the 10 candidates genes in the *Sleeping Beauty* dataset, 5 had been previously identified by retroviral insertional mutagenesis in RTCGD and were also identified in the MuLV screen described herein (see Section 3.3), while 3 had been previously identified

by transposon-mediated mutagenesis. Therefore, a higher percentage (42.9%) of these candidates than those in the MuLV screen (1.4%) overlapped with other transposon screens, again highlighting the different mutational profiles of the two mutagens. The Sleeping Beauty dataset is small, and none of the genes had Nanog or Oct4 binding sites (P=1 for both tests), while 1 had a p53 binding site (P=0.178) and 1 overlapped with the Sjöblom et al. (2006) dataset (P=0.087). However, 6 candidates were identified in the Cancer Gene Census and 5 had mutations in COSMIC. This is significantly greater than the number expected by chance $(P=4.26 \times 10^{-9} \text{ and } P=4.02 \times 10^{-4}, \text{ respectively})$. 5 genes were dominant cancer genes in the Cancer Gene Census ($P=1.16 \times 10^{-7}$), while 1 (*PTEN*) was recessive (P=0.032). Candidate genes were also enriched in regions of copy number loss (3 genes, P=0.0338) but not in regions of gain (2 genes, P=0.2450) in the Mullighan et al. (2007) dataset. There was an over-representation of genes (AKT2 and PTEN) in the melanoma, endometrial cancer and glioma KEGG pathways (P=1.68x10⁻³, P=9.26x10⁻⁴ and $P=1.36 \times 10^{-3}$, respectively) and in the REACTOME pathway associated with negative regulation of the PI3K/AKT network ($P=8.11 \times 10^{-5}$). Candidate genes were also overrepresented in the B-cell receptor signalling pathway (AKT2 and PPP3CA; $P=1.40 \times 10^{-3}$). Only 2 GO terms, "regulation of biological process" (8 genes, $P=6.10 \times 10^{-5}$) and "transcription factor activity" (Notch1, Fli1, Myb, Ikzf1 and Erg; P=3.39x10⁻⁵) were overrepresented, but the test was limited by the small size of the dataset.

The enrichment of candidate genes from the MuLV and *Sleeping Beauty* screens within human cancer-associated datasets demonstrates the efficacy of insertional mutagenesis as a tool for discovering human cancer genes, as well as those in mice. In addition, overlaying other cancer-associated datasets on to the insertional mutagenesis data helps to characterise the candidate genes and facilitates the identification of novel cancer genes. However, the approach is biased towards the identification of genes involved in the development of cancers of lymphoid origin. Candidate genes were positively associated with the Mullighan *et al.* (2007) dataset, which was generated using ALLs, and the Cancer Gene Census and COSMIC database, in which genes implicated in haematopoietic and lymphoid tumourigenesis are over-represented. Many of the over-represented GO terms were also directly related to the differentiation and activation of B- and T-cells. Conversely, candidates showed no significant association with the Sjöblom *et al.* (2006) dataset of colon and breast cancer genes. This highlights the importance of developing insertional mutagenesis screens that can induce other types of tumour, e.g. by integrating tissue-specific promoters into transposons or by spatial and temporal

regulation of transposase expression (see Section 1.4.2.2.1). The datasets discussed in this section are further referred to in the proceeding sections and chapters in relation to individual cancer gene candidates.

3.3 Comparison of candidate cancer genes in the MuLV and Sleeping Beauty datasets

There was a significant overlap between the lists of candidate cancer genes obtained using the retroviral and *Sleeping Beauty* (SB) screens. There was an overlap of 5 genes $(P=9.64 \times 10^{-31})$ when both lists generated using the kernel convolution (KC)-based approach were compared. Comparing the KC list of candidates from the retroviral screen to the *Sleeping Beauty* candidates generated using Monte Carlo (MC) simulations (Efr=0.001) produced an overlap of 10 genes $(P=1.95 \times 10^{-8})$. The KC lists therefore yielded the most significant overlap, which is consistent with the work described in Section 2.10.2, where the KC method was proposed to generate the most reliable set of candidate genes.

The distributions of MuLV and T2/Onc insertions across the mouse genome are shown in Figures 3.1A and 3.1B, respectively. The figures also show the location of candidate cancer genes that were identified by both screens, as well as all other *Sleeping Beauty* candidates identified using the KC method and a subset of the most frequently disrupted candidates from the MuLV screen. The most frequently mutated genes in MuLV-induced tumours were *Gfi1/Evi5*, *Myc/Pvt1* and *Ccnd3*. These genes had insertion densities of 427.28, 314.19 and 172.09, respectively, using the KC method with kernel width 30 kb. Remarkably, none of the SB-induced tumours contained insertions in or around these genes. While these genes are known to contribute to tumourigenesis, the frequency of insertions may reflect the bias of retroviruses to insert into particular sites in the genome (see Section 1.4.2.1.1). In addition, many of the MuLV insertions in these genes appear to be enhancer activity. Therefore, the frequency of insertions may reflect the choice of mutagen and does not imply that a gene would contribute to a similar proportion of spontaneous tumours in the mouse.

Conversely, a significant CIS comprising insertions in 6 SB-induced tumours was identified in the tumour suppressor gene *Pten*, but none of the MuLV-induced tumours

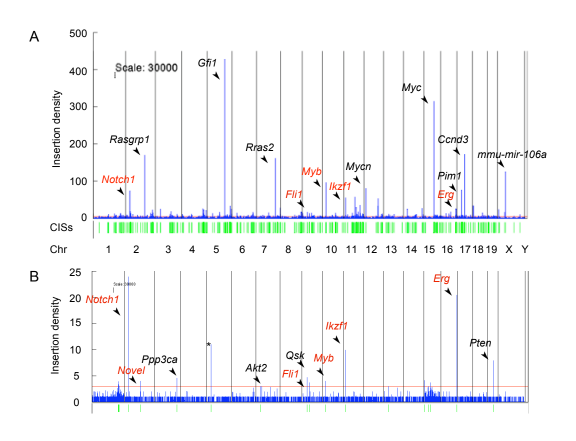


Figure 3.1. MuLV (A) and T2/Onc (B) insertions across the mouse genome. The plots show the density of insertions calculated using the kernel convolution-based method (de Ridder *et al.*, 2006) with a kernel width of 30 kb. Common insertion sites (CISs) are shown in green. The red line represents the threshold above which insertions form significant CISs (P<0.05). Gene names shown in red contain significant CISs in both screens. Gene names shown in black contain significant common insertion sites that are unique to one screen. * marks artefacts in *En2*.

contained an insertion within this gene. None of the T2/Onc insertions were in *Bloom*deficient tumours, which have an increased propensity for insertions within tumour suppressor genes (see Section 1.4.2.1.1). For 3 tumours, multiple insertions were identified, suggesting that the gene may be inactivated by insertions affecting both copies, rather than by one insertion in a region of loss of heterozygosity (LOH) (Figure 3.2A). The lack of MuLV insertions may reflect the fact that MuLV prefers to insert near to transcriptional start sites (Section 1.4.2.1.2) and is therefore more biased towards the identification of oncogenes than is the T2/Onc transposon. These observations suggest that MuLV and the T2/Onc transposon are unique mutagens with complementary mutagenic profiles, and that performing screens with both these mutagens can identify more candidate cancer genes than with either alone.

As well as the distinct differences between the mutagenic profiles of MuLV and T2/Onc, a number of known and implicated cancer genes (*Notch1*, *Erg*, *Ikzf1*, *Myb* and *Fli1*) contained significant CISs in both screens. The co-occurring MuLV and T2/Onc insertions within these genes are discussed in Section 3.4.2. After *Myc/Pvt1*, *Gfi1/Evi5* and *Ccnd3*, the most highly mutated genes in the MuLV screen were *Rasgrp1* (insertion density 169.59) and *Rras2* (insertion density 161.49). Although significant *Sleeping Beauty* CISs were not identified in these genes using the KC method, *Rras2* did contain 1 T2/Onc insertion, and *Rasgrp1* contained 3 T2/Onc insertions, which was significant using the Monte Carlo method with *Efr*=0.005. Likewise, *AA536749*, *Zmiz1*, and known oncogenes *Irf4* and *Etv6*, contained significant MuLV CISs identified using the KC method and T2/Onc CISs that were significant using the MC method with *Efr*=0.001.

The human orthologue of AA536749 is myosin phosphatase Rho-interacting protein (*p116Rip* or *M-RIP*). p116Rip is a filamentous actin-binding protein that is capable of disassembling the actomyosin-based cytoskeleton and acts downstream of RhoA (Mulder *et al.*, 2003). The actin cytoskeleton plays a role in many cancer-related functions such as cell motility, cell differentiation, cell survival and cell division. The LIM kinases (LIMK1 and LIMK2) are regulators of actin dynamics that also act downstream of Rho GTPase and play an important role in tumour invasion and metastasis (Scott and Olson, 2007). The identification of insertions in tumours generated by both mutagens suggests that *p116Rip* may also play an important role in tumourigenesis.

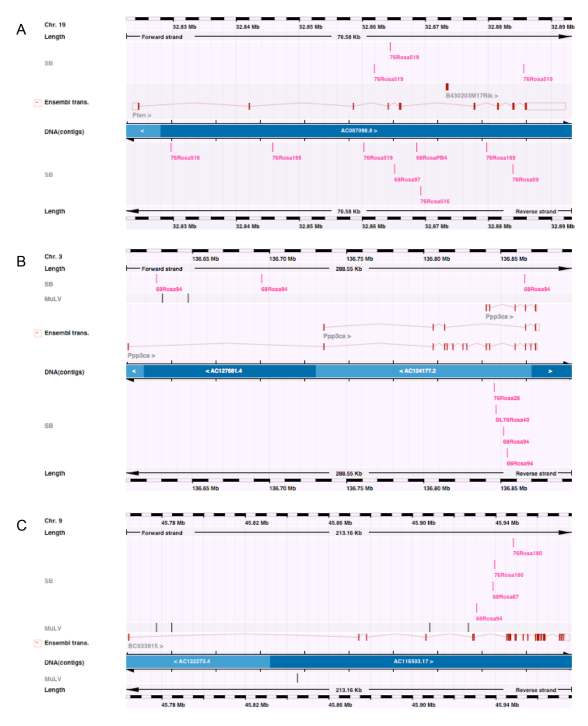


Figure 3.2. Known and putative tumour suppressor genes identified in the *Sleeping Beauty* (SB) screen. *Pten* (A) did not contain any MuLV insertions. *Ppp3ca* (B) and *BC033915* (C) contained MuLV insertions but not in a statistically significant CIS. For all genes, there was at least 1 SB tumour that contained more than 1 insertion, suggesting that inactivation of both genes may be required for tumourigenesis. The tumour in which each T2/Onc insertion was identified is provided as a label under the insertion, which is shown in pink. Ensembl genes are shown in red and, where applicable, MuLV insertions are shown as black vertical lines. Insertions above and below the blue line are in the forward and reverse orientation, respectively.

Zmiz1 enhances p53 (Lee *et al.*, 2007) and Smad transcriptional activity (Li *et al.*, 2006b), suggesting a tumour suppressive role. However, it is also required for vasculogenesis (Beliakoff *et al.*, 2008) and activates transcription of the androgen receptor (Beliakoff and Sun, 2006; Sharma *et al.*, 2003), which contributes to the formation and progression of human prostate cancer (for review, see Nieto *et al.*, 2007). In addition, a fusion between *ZMIZ1* and *ABL1* was recently identified in a human B-cell acute lymphoblastic leukaemia (Soler *et al.*, 2008). There are 4 other known fusion partners (*BCR, ETV6, NUP214* and *EML1*) for *ABL* in human haematological malignancies, and one putative partner, *RCSD1* (De Braekeleer *et al.*, 2007). Remarkably, *Zmiz1, Etv6, Nup214* and *Rcsd1* all contained statistically significant CISs in the retroviral screen and, although not significant, *Bcr* contained 2 MuLV insertions, while *Eml1* contained 1 MuLV and 1 T2/Onc insertion. This reflects the fact that mutagenesis by MuLV often resembles the effects of translocation, as mentioned in Section 3.3.2.

The remaining candidates containing significant *Sleeping Beauty* CISs identified using the KC method also contained retroviral insertions, although not significant CISs. The known oncogene *Akt2* and the serine/threonine protein phosphatase *Ppp3ca* contained 1 and 2 MuLV insertions, respectively. One of the 3 SB-induced tumours in which *Ppp3ca* was disrupted contained 4 insertion sites, suggesting that this gene encodes a tumour suppressor (Figure 3.2B). This is supported by research showing that *Ppp3ca* can dephosphorylate cyclin dependent kinases (CDKs), therefore potentially inhibiting cell cycle progression (Cheng *et al.*, 1999, see Section 1.2.6 for more on CDKs). In addition, *Ppp3ca* overexpression increases the levels of p53 and inhibits cell growth (Ofek *et al.*, 2003), and expression is reduced in androgen-independent prostate cancer cells (Singh *et al.*, 2008).

The gene encoding serine/threonine protein kinase BC033915 (known as QSK in humans) was also identified in both screens, and although the MuLV CIS was not significant, it did contain 5 retroviral insertions (Figure 3.2C). In the COSMIC database, 2 of the 296 human tumour samples that have been tested for mutations in the *QSK* gene contain missense mutations. One heterozygous S882C substitution was identified in the primary renal cell carcinoma PD1583a, which contains just one other small intragenic mutation in 519 genes examined, and a P836S substitution (zygosity unknown) was identified in the non-small cell lung cancer cell line NCI-H1770, which contains 201 small intragenic mutations in 4,688 genes examined. 1 silent mutation (heterozygous

substitution R476R) was also identified in the malignant melanoma cell line MZ7-mel, but this line contains 428 mutations in 4,668 genes examined and therefore appears to have a hypermutable phenotype. Both missense mutations are located in a glutamine-rich region (Prosite profile PS50322). All of the retroviral and transposon insertions in *QSK* precede this region and may therefore produce truncated gene transcripts in which the region is missing. 2 heterozygous, missense mutations were also identified in the resequencing study by Sjöblom et al (2006). QSK and 11 other kinases related to AMPactivated protein kinase (AMPK) are known to be activated by the tumour suppressor kinase LKB1 (Lizcano et al., 2004). Activation of one of these kinases (MARK1) by LKB1 has been shown to regulate microtubule dynamics by phosphorylating the microtubule-associated protein Tau, thereby reducing the affinity of Tau for microtubules and inhibiting tubulin polymerisation (Kojima *et al.*, 2007). QSK may play a similar role, since RNAi-mediated knockdown of the Drosophila orthologue of OSK resulted in spindle and chromosome alignment defects (Bettencourt-Dias et al., 2004). One of the SB-induced lymphomas contained 2 insertion sites in Osk, and this, coupled with the observations described above, suggests that *Qsk* may be a tumour suppressor gene.

Finally, a novel gene, *ENSMUSG0000075015*, contained a significant T2/Onc CIS and 1 MuLV insertion. 2 T2/Onc insertions were in the antisense orientation with respect to the gene, suggesting that the gene might encode a tumour suppressor, but functional analysis is required to determine the role of this gene in tumourigenesis.

The *Sleeping Beauty* dataset is relatively small and few candidate cancer genes have been identified. However, comparison with the MuLV screen demonstrates the potential benefits of using multiple mutagens to increase the spectrum of candidate cancer genes, but also to identify strong candidates that are independently mutated by both screens and are therefore unlikely to result solely from insertional bias. 3 of the 10 genes identified in the *Sleeping Beauty* screen are known or putative tumour suppressor genes, i.e. *Pten*, *Ppp3ca* and *Qsk*, suggesting that the T2/Onc mutagen is an effective tool for identifying recessive cancer genes. Scaling up the screen to identify further candidates would provide a valuable dataset to complement the retroviral insertional mutagenesis data.

3.4 Determining the mechanisms of MuLV insertional mutagenesis

3.4.1 Analysing the distribution of intragenic insertions

Analysis of the distribution of insertions within and around genes can help to determine the likely mechanisms of mutation. Oncogenic insertions in intergenic regions are likely to be promoter or enhancer mutations that result in increased levels of the wildtype protein. However, the effect of intragenic insertions, of which there are 8,447 (42.0% of the total), may be less obvious. The Ensembl API version 45 36f was used to identify the genomic coordinates of untranslated regions (UTRs), exons and introns in the longest transcript of each candidate cancer gene. From these, coding and non-coding exons, and introns within coding regions or UTRs, were distinguished. The "gene regions" were defined as 5' UTR, intron in 5' UTR, coding exon, intron flanked by coding exons, intron in 3' UTR, and 3' UTR. For each candidate, the number of insertions in each gene region was counted, and the orientation of each insertion with respect to the disrupted gene was determined. The total number of insertions in each gene region is shown in Figure 3.3A. The collective length of each gene region across all candidate genes was also calculated and, for each region, the insertion count was divided by the length in base pairs to give an indication of the proportion of insertions given the region size (Figure 3.3B). For insertions within introns or exons of the coding region, the identity, i.e. number, of the exon or intron containing the insertion was determined. This is helpful in determining the mechanism of mutation since, if a specific oncogenic gene product is formed, multiple insertions would be expected to localise to the same region of the gene.

Within genes, introns were the most common site of insertion, but for their size, they were the least commonly hit region. Intronic insertions may result in the formation of Nor C-terminal truncations. There are polyadenylation sites in both orientations of the retroviral provirus but promoters are only found in the forward orientation of the retroviral LTRs (see Section 1.4.2.1.1). Therefore, while antisense insertions can only form C-terminal truncations, sense insertions can form both N-terminal and C-terminal truncations. The distribution and orientation of intronic insertions will vary in different oncogenes depending on how an oncogenic mutant is created, and tumour suppressor genes can be inactivated by any distribution of insertions that results in non-functional Nor C-terminal truncations. There were therefore roughly equal numbers of sense and antisense insertions in introns. Genes with the highest numbers of intronic insertions were *Ikzf1* (56 insertions) and *Notch1* (52 insertions). The insertions in *Ikzf1*, a tumour

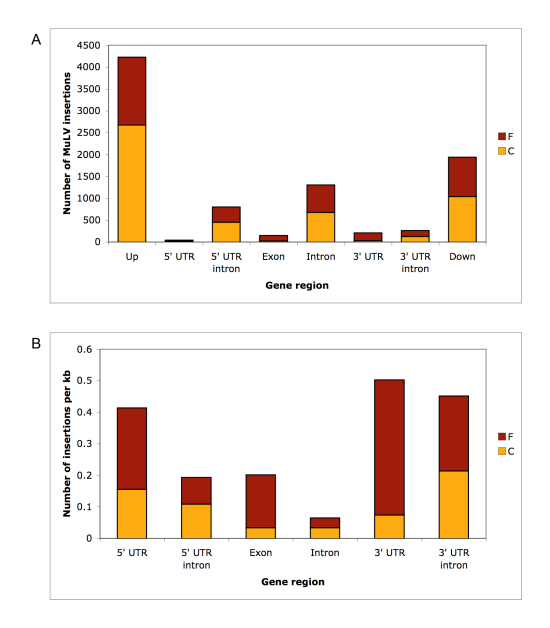


Figure 3.3. The distribution of MuLV insertions within candidate cancer genes. (A) The total number of insertions in each gene region. (B) The number of insertions as a proportion of the total length of the gene region across all candidate genes. The number of insertions in the sense orientation (F) with respect to genes is shown in red, while the number in the antisense orientation (C) is shown in yellow.

suppressor gene (see Section 3.4.4), are likely to cause premature termination of gene transcription, resulting in gene inactivation, while those in *Notch1* are likely to produce distinct, truncated mutant proteins that are implicated in tumourigenesis (see Section 3.4.2 for further details). Flt3 contained 40 intronic insertions that were all in the sense orientation, suggesting that the gene is most likely disrupted by the formation of Nterminal truncations (Figure 3.4A). Most insertions were within intron 9 and are predicted to result in the production of proteins lacking the extracellular, ligand-binding, Ig-like domain. FLT3 is mutated in around one third of human acute myeloid leukaemias, yet it is mutated by internal tandem duplications or by point mutations that produce a constitutively active protein (Small, 2006). 4,170 out of the 20,259 haematopoietic and lymphoid cancer samples tested in COSMIC have a mutation in *FLT3*, of which 185 have a missense mutation at amino acid 835 in the protein kinase core domain. Most of the remaining samples have internal tandem duplications that are represented in COSMIC by complex mutations and indels. This suggests that retroviral insertional mutagenesis may not always accurately recapitulate the mutations contributing to human cancers. Antisense insertions occurring in introns close to the 5' end of a gene could be inactivating mutations affecting tumour suppressor genes, or may result in the production of a truncated transcript from a cryptic transcription start site further downstream within the gene. It is also possible that they are acting as enhancer mutations.

The second most frequently hit regions were introns in the 5' UTR, i.e. introns that are flanked on each side by exons of the 5' UTR. Again, these collectively form a larger region than coding and non-coding exons. There were 28.8% more insertions in the antisense orientation than in the sense orientation. Sense insertions are most likely to be promoter mutations, which result in increased production of the full-length cellular Antisense insertions may be prematurely terminating gene transcription, protein. resulting in the complete absence of the gene product, as might be expected for tumour suppressor genes, or they may result in the production of a truncated transcript from a cryptic transcription start site. They could also be intragenic enhancer mutations or, as the longest gene transcript has been selected for this analysis, it is possible that some are enhancer mutations that are upstream of alternative gene transcripts, and are therefore producing full-length, wildtype proteins at increased levels. Cyclin D3 (Ccnd3) contained the highest number of insertions (204) within 5' UTR introns, with 85% occurring in the antisense orientation (Figure 3.4B). Since Ccnd3 is an oncogene, and contains no known cryptic transcription start sites or alternative transcripts, it is likely that

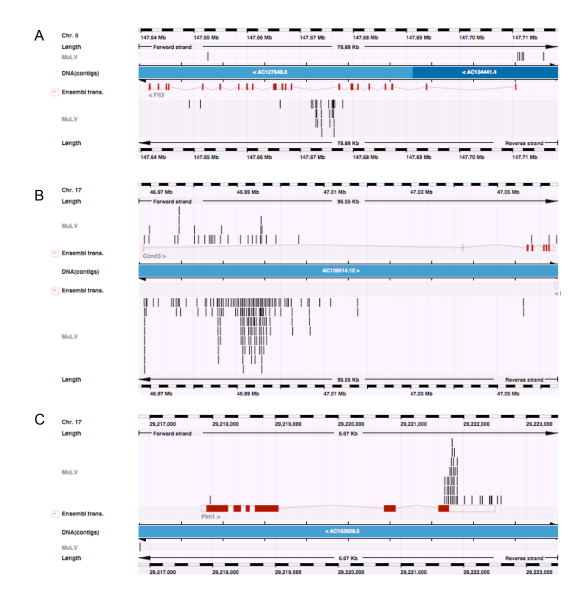


Figure 3.4. Intragenic MuLV insertions in candidate cancer genes. (A) Intronic insertions in *Flt3* are predicted to generate N-terminally truncated gene products. (B) Antisense insertions in the 5' UTR of *Ccnd3* are likely enhancer mutations. (C) Insertions in the final coding exon and 3' UTR of *Pim1* may cause premature termination of gene transcription that leads to a more stable gene product. Insertions are shown in black. Genes are shown in red. Insertions above and below the blue line are in the forward and reverse orientation, respectively.

the insertions are enhancer mutations. *Lck* contained 18 sense insertions but no antisense insertions in intronic regions of the 5' UTR, and a further 3 sense insertions in the 5' UTR and first coding exon, suggesting that all of the insertions are involved in the formation of chimeric transcripts in which the retroviral promoter drives increased expression of the cellular gene.

The 5' UTRs of candidate genes contained an over-representation of sense insertions, as expected for promoter insertions. *Myc* contained the most sense insertions, totalling eight. Antisense insertions in the 5' UTR may interfere with gene transcription, preventing protein production or resulting in transcription from a cryptic or alternative promoter.

Exons and 3' UTRs showed a strong bias towards insertions in the sense orientation. 80% of sense insertions in coding exons were in *Notch1*, *Mycn*, *Map3k8*, *Ccr7*, *Pim1* and *Jundm2*, and in all cases, insertions were at the 3' end of the gene, close to the 3' UTR. In the case of *Mycn* and *Pim1*, which also contained a large number of 3' UTR insertions, these insertions cause premature termination of gene transcription that result in the removal of mRNA-destabilising motifs and, therefore, the generation of a more stable gene transcript (Cuypers *et al.*, 1984; Selten *et al.*, 1985; van Lohuizen *et al.*, 1989). Insertions within *Pim1* are shown in Figure 3.4C. It is possible that *Ccr7* and *Jundm2* are disrupted by the same mechanism, since both contained sense insertions in the final coding exon and the 3' UTR. The near-exclusivity of sense insertions in these genes suggests that the polyadenylation site in the forward orientation. In summary, the density of insertions in exons and UTRs was higher than for introns, suggesting the importance of promoter insertions and "stabilising" insertions as mechanisms of mutagenesis.

For each candidate cancer gene, the distribution of insertions was used to predict the likely mechanisms of mutagenesis and, therefore, the likely structures of mutated gene products. Sense insertions that were upstream of the gene, within the 5' UTR or in an intron flanked by exons of the 5' UTR were classified as promoter mutations. Upstream insertions in the antisense orientation were classified as enhancer mutations. Insertions in the 3' UTR were classified as "stability" mutations, i.e. insertions that may result in the removal of mRNA-destabilising motifs, while sense and antisense insertions in exons or

introns in the coding region were classified as C- or N-terminally and C-terminally truncating mutations, respectively. Finally, antisense insertions in introns within the 5' UTR remained unclassified, since these have a number of possible effects (see above). This yielded 360 genes with enhancer insertions, 309 with promoter insertions, 45 with stability mutations, 183 with C-terminally truncating insertions, 202 with C- or Nterminally truncating mutations, and 92 with antisense insertions in introns within the 5' UTR. Most genes are associated with multiple types of insertion (see Table 3.3). Genes were further classified according to the predicted protein generated by the mutations. Genes containing any combination of promoter, enhancer and stability mutations should generate the wildtype protein at increased levels compared with the endogenous gene. It was assumed that where both sense and antisense insertions occurred in the same exon and intron, the gene was C-terminally truncated, whereas if only sense insertions occurred, the gene was N-terminally truncated. Sense insertions in the last intron were classified as C-terminally truncating, as commonly observed, for example, in Pim1 and This generated 7 types of mutant – upregulated wildtype (201 genes), C-Mvcn. terminally truncated (30 genes), N-terminally truncated (2 genes), C- and N-terminally truncated (4 genes), and upregulated wildtype plus C-terminally truncated (122 genes), Nterminally truncated (56 genes) or C- and N-terminally truncated (24 genes). This suggests that a high proportion of genes contribute to tumourigenesis by increased production of the wildtype protein. C-terminally truncating mutations appear to be more common than N-terminally truncating mutations, but this may reflect the fact that insertions in the sense orientation were assumed to be C-terminally truncating if antisense insertions were also present, and therefore some may have been misclassified. The genes associated with each mutation type are presented in Table 3.3.

Elucidation of the mechanisms of mutagenesis is complicated by insertional bias and the ability of MuLV to disrupt a gene in multiple ways. Therefore, predictions must be experimentally validated, e.g. by measuring the length of transcripts generated by insertion-containing genes and by analysis of gene expression in MuLV-induced tumours (see Section 3.4.5). Reducing the number of ways in which an insertional mutagen can disrupt a gene would facilitate the analysis of insertions within genes. For example, by using a transposon engineered with a splice acceptor site and polyadenylation site on one strand only, it would be possible to distinguish C- and N-terminal truncations with a high degree of certainty.

Mutation type	Mechanisms of mutagenesis	Number of genes	Gene names
wildtype	P, E		Cxcr4, Med13, mmu-mir-23a, Btg2, Art2b, Fgfr3, Fut8, Mknk2, Lyst, Fli1, Scyl1, Actr3, Tceb3,
	.,.		Cach, Yinchi J., Mind Yim, Zab, Gag, Miczi, Girlo, Yun, Kinka, Song, Har, Scy, Ti, Haor, Rpli1, Kir, Zfp438, 1700122011Rik, Ier2, Acot11, Lta, Metrnl, Mpl, Cerc5, Pegf5, Cd47, Haao, Rpli1, Mixil, Hrbl, Q8BP09_MOUSE, Stmn1, C330024D12Rik, Ccnd2, Hspa9, Ttl10, Stat5a, Egfl7, Bex6, Irf2bp2, Cstad, Tpd52, Hibadh, Lfng, Lgals9, Psma1, ENSMUSG00000074256, Cd72, 1700019017Rik, Anp32e, C130026L21Rik, Rcbtb2, Jph4, Ccrk, Brd2, AA536749, Frmd8, Mafk, Vpreb2, Hnrpf, A13005007Rik, Lmo2, Ccdc19, AI848100, Aars2, Akt1, ENSMUSG00000073531, mmu-mir-181c, Rreb1, mmu-mir-17, Spn, 6430598A04Rik, Ppp2r5a, Ina, SIC39a13, mmu-mir- 802, mmu-mir-142, Rcsd1, Lat, 2310016C08Rik, Trp53inp1, Chd2, Tc77, Gadd45g, Rpl24, Cbl, Appl2, Ifnar1, Park7, Vdac1, ENSMUSG00000058984, ENSMUSG000000731320, ENSMUSG00000059313, ENSMUSG00000013756, ENSMUSG0000063435, ENSMUSG00000059882, ENSMUSG00000034596, ENSMUSG0000063488,
wildtype	P, E, CT	41	LIGHDSG00000012358, Rara, Bci9, Kli3, Mobkla, Tmem90a, Evil, AS30013C23Rik, Mgat1, Ldha, 4831426119Rik, Tcof1, Sic38a1, 2310066E14Rik, Arhgdib, Plekha2, Gimap4, A630001012Rik, Zfp217, Il6st, Rasgrp2, B3gnt2, A930002121Rik, Fgr, Pag1, Sema4d, Ptp4a2, Rhoh, Zeb2, Mef2d, Ggta1, Stat5b, Pik3r5, Arid1a, Csk, Thra, Lcp1, Parvg, Hmga1, Gpr132,
wildtype	E		Ddr1, Zdhhc19, Chd7, Bid, Zfp36l2, Edg1, Dad1, Gfi1b, Mcl1, Fgd2, ENSMUSG0000074788, Rps14, Chst3, Trim47, Fgfr2, Aqp4, ENSMUSG0000074675, Frat2, Kihl25, Chchd7, Hhex, ENSMUSG0000072756, ENSMUSG0000061115, ENSMUSG0000046809, ENSMUSG00000072757, ENSMUSG0000052894
wildtype	P, CT		4932417H02Rik, Phyhd1, Cd53, Rhbdf2, Gpr56, Ptp4a3, Pitpnm2, Olfr56, 6330548G22Rik, Evi2b
wildtype	P, E, S		Hoxa7, 4632428N05Rik, BC027057, Sox19, OTTMUSG00000016805, Gadd45b, Ccnd1, 3930402G23Rik, Pim1
wildtype	E, S		Evi5, Gpr152, Clec2d, Scd1, BC030863, Scube1
wildtype	Р		Ppp1r10, Pscd4
wildtype	P, E, S, CT		Pvt1, Rassf2
wildtype	P, S		Orai2, Mycn
wildtype	E, CT		Lrrc8c
wildtype	P, S, CT		Zbtb7b
wildtype & C-trunc	P, E, C, C/N	30	Nsmcel, Gsel, Ahil, PrrG, Ski, Ptprc, Itpr2, Map3kl, Abcb9, Foxp1, Exoc2, Cd97, Srgn, Tmem131, Vamp8, AB041803, Etv6, Prkch, Ssbp3, Supt3h, Akna, Gm525, Rasgrp1, Zfp608, Tcfap4, Arid3a, Ntn1, 2010107G12Rik, Sema4b, Ets1
wildtype & C-trunc	E, C, C/N	26	Akap13, Cd2, Dym, Vps13d, Tmprss3, Pxn, Pygm, Tgfbr3, Zc3h12a, Bcl11b, Mad1l1, Spsb4, Gimap6, H2-D1, Nup214, Lims1, Znrf1, Kcnn4, Tmem49, Kctd2, Sept9, Slamf6, Dopey2, Itpkb,
wildtype & C-trunc	P, E, C		Exoc6, Colq, Hdac7a, Cybasc3, Lef1, Mylc2pl, Tmem173, ENSMUSG00000074787, Nfe2, B3gnt1, Treh, Slamf7, Aqp9, Cd27, 5730559C18Rik, EG433384
wildtype & C-trunc	P, C, C/N		Cldn10a, Spata13, Ahnak, Vil2, Stard10, Ubac2
wildtype & C-trunc	E, S, C, C/N		Cugbp2, Arhgap26, Arhgef3, 1110036003Rik, Rab37, Psmb8
wildtype & C-trunc	P, C, CT		Asb2, Elf1, Adrbk1, Arpp21, Ptpre
wildtype & C-trunc	P, E, C, CT		Ncoa3, Emp3, Usp52, Il2ra, Wasf2
wildtype & C-trunc	Е, С		A130092J06Rik, Cdkl3, Stra8, Gna15, Kit
wildtype & C-trunc	P, E, C, C/N, CT		Zmiz1, 4932422M17Rik, Ubxd5, Ikzf1
wildtype & C-trunc	P, E, S, C, C/N		Padi2, Ksr1, Pigv
wildtype & C-trunc	P, C, C/N, CT		1600014C10Rik, Anxa2, D18Ertd653e
wildtype & C-trunc	P, C		BC008155, Epha6
wildtype & C-trunc	C, C/N, CT		Bcl2l1, Ppp1r16b Arhgef10l, Il21r
wildtype & C-trunc wildtype & C-trunc	E, C, C/N, CT P, S, C, C/N		Nfkb1
wildtype & C-trunc	P, E, S, C, C/N, CT		Jundm2
wildtype & C-trunc	S, C, N		Rnf43
wildtype & C-trunc	S, C		Ovca2
wildtype & C-trunc	P, E, S, C		Trpm1
wildtype & C-trunc	E, S, C		C330016O10Rik
wildtype & N-trunc	P, E, C/N	31	Runx1, Set, Cd3e, Cd48, 2410014A08Rik, Pctk2, Fchsd2, Paics, Thy1, Msh5, OTTMUSG00000005737, Jup, Sdk1, Prkcbp1, Mbd2, Ardc5, Coro2a, Hipk1, Erg, D12Ertd553e, Rras2, Nedd4l, Myb, Eng, Plac8, Tspan2, 1190002H23Rik, Tap2, Ptbp1, Mns1, Ube11, Chd1
wildtype & N-trunc	E, C/N	Q	Sic36a3, Nfkbil1, Rtn4rl1, Bcl11a, Ak1, Ndrg3, NP_001074704.1, Map3k8
wildtype & N-trunc	P, E, C/N, CT		Tcte3, Sla2, Sla, Grap2, Pik3cd, Tbxa2r
wildtype & N-trunc	P, C/N, CT		Zfp710, Tspan14, Dnahc8
wildtype & N-trunc	P, E, S, C/N		Irf4, Midn
wildtype & N-trunc	E, S, C/N		Stard3nl, 1700081D17Rik
wildtype & N-trunc	P, E, S, C/N, CT		E230001N04Rik
wildtype & N-trunc	P, E, N, CT		Ccnd3
wildtype & N-trunc	P, E, S, N, CT		Мус
C-trunc	C, C/N	29	Nfix, Sic43a2, 2010106G01Rik, Nup210, Smg6, Hvcn1, Stk4, Lrrfip1, Dpp4, Rorc, Ramp1, Myo18a, Clec16a, Jazf1, Sh3bp5, Pixnd1, Prdm16, 3110082117Rik, Serinc3, Fyb, B320120H23Rik, St6galnac5, Scotin, II16, E2f2, Usp7, Recql5, Sirt2, Abcg1
C-trunc	С		Fmnl1
N-trunc	C/N		1300007F04Rik, Exosc5
C-trunc & N-trunc	C, N	4	Xrcc6, Rnf166, Sh3d19, Iqch
wildtype & C-trunc & N-trunc	P, E, C, N	10	Lck, Slc1a3, Dhx40, A2AN91_MOUSE, Rnf157, Katnal1, Mgat4a, Dock8, Cyb5, Cbfa2t3
wildtype & C-trunc & N-trunc	E, C, N	6	Wwox, Nid1, Capsl, Tcf25, Flt3, Tbc1d1
wildtype & C-trunc & N-trunc			Ubash3a, Pml
wildtype & C-trunc & N-trunc			Runx3, Pecam1
wildtype & C-trunc & N-trunc			Mrvi1
wildtype & C-trunc & N-trunc	P, E, S, C, N	1	Ccr7, Notch1, Gfi1, Ikzf3

Table 3.3. The predicted mutation types and mechanisms of mutagenesis based on the distribution of MuLV insertions within and around candidate cancer genes. C-trunc = C-terminally truncated, N-trunc = N-terminally truncated, P = promoter insertion, E = enhancer insertion, C = antisense intragenic insertion, C/N = sense intragenic insertion, S = stabilising insertion, CT = antisense insertion 5' of first coding exon (i.e. truncating, leading to inactivation or use of cryptic transcription start site, or enhancer mutation).

3.4.2 Analysing co-occurring insertions in candidate genes disrupted by MuLV and T2/Onc

The co-occurrence of MuLV and T2/Onc insertions in distinct regions of genes provides strong evidence that the insertions do not result from insertional bias and that they play an important role in oncogenesis. Such insertions can provide important clues about the mechanism of mutation and, therefore, about the structure and function of genes and oncoproteins involved in cancer. In this section, the distributions of MuLV and T2/Onc insertions are compared within known and implicated cancer genes that overlap between the MuLV kernel convolution (KC)-based list of candidates and *Sleeping Beauty* (SB) candidates from the KC list, i.e. *Notch1*, *Myb*, *Fli1*, *Erg* and *Ikzf1*, and from the Monte Carlo (*Efr=*0.005) list, i.e. *Rasgrp1* and *Etv6*.

In Notch1, MuLV and T2/Onc insertions co-occurred in the same orientation in 3 distinct regions of the gene (Figure 3.5A). Antisense MuLV and T2/Onc insertions were identified in the second intron. The retroviral insertions could be assumed to be enhancer mutations, yet T2/Onc has low enhancer activity. Therefore, these are more likely to be truncating mutations, and this is consistent with the observation that radiation-induced deletions in the 5' region of Notch1 result in truncated proteins that lead to the development of mouse thymic lymphomas (Tsuji et al., 2003). The authors showed that deletion of, or MuLV insertion into, the juxtamembrane extracellular region encoded by exons 1 and 2 results in transcription from cryptic transcription start sites further downstream in Notch1 and leads to the production of an active protein lacking most of the extracellular domain. Co-occurring sense insertions were also identified in the 28th and 29th introns. Based on their orientation, these insertions are expected to produce Nterminally truncated proteins containing only the intracellular domain of Notch1. This form of Notch1, called Notch1IC, is constitutively active and is associated with leukaemogenesis (for review, see Aster et al., 2008). Finally, there were co-occurring insertions, again mostly in the sense orientation, within the final coding exon. These insertions were upstream of the PEST domain, which regulates turnover of Notch1IC (Aster et al., 2008). Deletion of the PEST domain, by MuLV insertion in T-cell lymphomas and by radiation in the study by Tsuji et al. (2003), is believed to contribute to tumourigenesis in collaboration with other activated oncogenes (Feldman et al., 2000; Hoemann et al., 2000; Tsuji et al., 2003). 184 human tumour samples out of 1,909 tested contain NOTCH1 mutations in the COSMIC database. Of these, 180 are in

Chapter 3

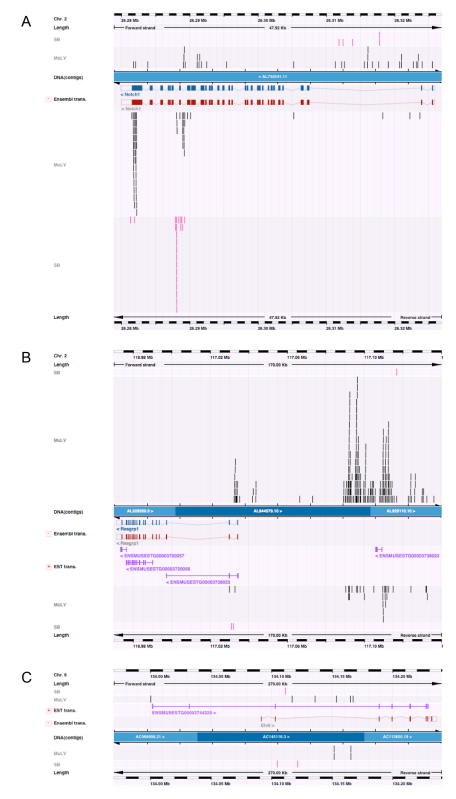


Figure 3.5. Co-occurring MuLV and T2/Onc insertions help to identify the mechanism of mutagenesis of genes *Notch1* (A), *Rasgrp1* (B) and *Etv6* (C). MuLV insertions are shown in black, T2/Onc insertions are shown in pink. Ensembl gene transcripts are shown in red and blue. ESTs are shown in purple. Insertions above and below the blue bar labelled DNA(contigs) are in the forward and reverse orientation, respectively.

haematopoietic and lymphoid tissue, which corresponds to 24% of all samples of this cancer type tested. *NOTCH1* is therefore specifically, and significantly, associated with cancers of this type.

Co-occurring MuLV and T2/Onc insertions were also identified in the first intron, preceding the first coding exon, of Rasgrp1 (Figure 3.5B). These insertions, which are in the sense orientation with respect to the gene, are likely to be promoter mutations that result in overexpression of the full-length transcript. MuLV enhancer mutations were also found upstream in the antisense orientation but, unsurprisingly, these were not identified in the SB screen since T2/Onc has low enhancer activity. There were no intragenic insertions beyond the first intron, suggesting that only the full-length gene contributes to oncogenesis. This is supported by the observation that, among 273 tumour samples tested, there are none with somatic mutations in RASGRP1 in the COSMIC database. Deregulated expression of full-length murine Rasgrp1 contributes to the development of T lymphocytic leukaemias (Klinger et al., 2005), and to the progression of skin carcinogenesis through the activation of the Ras oncogene (Luke et al., 2007). Interestingly, in previous screens, insertions that are \sim 60-100 kb upstream have been assigned to Rasgrp1 (Hansen et al., 2000; Hwang et al., 2002; Kim et al., 2003a; Mikkers et al., 2002; Stewart et al., 2007; Suzuki et al., 2006; Suzuki et al., 2002). However, analysis of the insertions in the context of Ensembl shows that they are flanking an Ensembl EST gene for which there is no associated Ensembl gene transcript (Figure 3.5B). Expression analysis of *Rasgrp1* in the affected tumours is required to determine whether it is indeed disrupted by these insertions, but this observation suggests that the analysis of insertion sites in the context of the mouse genome could potentially help in the identification of "new" mouse transcripts.

All of the MuLV and T2/Onc insertions identified in gene Etv6 were in the second intron, in both orientations (Figure 3.5C). Since sense insertions can form both N-terminal and C-terminal truncations but antisense insertions can form only C-terminal truncations, it is likely that where insertions occur in both orientations in the same intron of an oncogene, they are causing premature termination of gene transcription that results in C-terminally truncated gene products. In the case of Etv6, this would result in the production of polypeptide lacking both of its functional domains. Etv6 is a transcriptional repressor that is essential for haematopoietic stem cell function. The N-terminal sterile alpha motif/pointed (SAM PNT) domain (IPR0003118) is responsible for hetero- and homodimerisation with other ETV6 and Ets (erythroblast transformation specific)-family Deletion of this domain decreases the inhibition of macrophage colony proteins. stimulating factor receptor (MCSFR) promoter activation by CBFA2B and C/EBPa, but does not completely abrogate it (Fears et al., 1997). However, the SAM PNT domain is necessary for interaction with, and inhibition of, the FLI1 oncogene (Kwiatkowski et al., The winged helix DNA-binding Ets domain (IPR000418) is essential for 1998). inhibiting the activation of MCSFR (Fears et al., 1997). Therefore, deletion of both these domains in the mouse most likely produces a non-functional protein, resulting in the overexpression of Etv6 target genes, such as *Fli1*. As mentioned previously, *ETV6* forms a fusion with ABL, but also with many other genes, in human leukaemias (for review, see Bohlander, 2005). Interestingly, while most fusions with tyrosine kinase genes contain a breakpoint in intron 4 or 5 of ETV6, and, for example, fusions with RUNX1 contain a breakpoint in intron 5, there are also fusions with unique or rare recurrent gene partners in which ETV6 has a breakpoint in intron 2. It has been suggested that promoters in the latter ETV6 truncation upregulate nearby oncogenes (Jalali et al., 2008; Panagopoulos et al., 2006). However, the distribution of MuLV and T2/Onc insertions within ETV6 suggests that the nonfunctional truncation may also itself contribute to leukaemogenesis. The ETV6-RUNX1 fusion is consistently associated with deletion of the normal ETV6 allele, suggesting that normal ETV6 represses ETV6-RUNX1 by interaction via the SAM PNT domain (Hart and Foroni, 2002; Raynaud et al., 1996). Homo- and heterodimerisation are believed to repress the activity of Ets proteins (Carrere et al., 1998; see discussion below in relation to the Erg gene) and therefore, it is possible that by deleting one allele, fewer heterodimers will be formed with other Ets proteins, resulting in increased activity of those proteins. Incidentally, 1 sense and 1 antisense MuLV insertion were identified 91.05 kb and 147.04 kb, respectively, upstream of the Etv6 gene. It could be assumed that these insertions are not oncogenic, since they are a considerable distance from the gene. However, the sense insertion is just 1.22 kb upstream of an Ensembl EST gene for which there is no associated Ensembl gene transcript, which suggests that there may be an unannotated alternative transcript of *Etv6* (see Figure 3.5C).

The rest of the genes that were disrupted by both MuLV and SB showed variation in the distribution of insertions. In some cases, this reflects differences in the mutational mechanisms of the two mutagens. For example, 1 retroviral insertion and 1 transposon insertion were found to co-occur just upstream of Myb (Figure 3.6A) in the sense orientation, where they are likely to be causing promoter mutation, but the vast majority

of retroviral insertions were putative enhancer mutations, occurring upstream in the antisense orientation, or downstream, predominantly in the sense orientation with respect to *Myb*. The presence of MuLV sense and antisense insertions, and 1 SB sense insertion, just upstream of an Ensembl EST gene suggests that, as for *Etv6*, there is an additional *Myb* transcript that has not been annotated as an Ensembl gene transcript. The remaining T2/Onc insertions were intragenic. 3 were in the last (13^{th}) intron in the antisense orientation with respect to *Myb*, while 1 was in the 10^{th} intron in the sense orientation. A C-terminal truncation caused by the latter would truncate the C-terminal Myb domain (IPR015395), which is known to bind the inhibitor Cyp-40 (Leverson and Ness, 1998). Oncogenic *v*-*Myb* contains a mutated binding site that prevents binding of Cyp-40 and so prevents negative regulation (Leverson and Ness, 1998). The contribution of insertions within the last intron is unclear since a C-terminally truncated protein would contain an intact binding domain. It is possible that the last exon of *Myb* encodes a protein sequence with a hitherto uncharacterised role in oncogenesis.

Variation in the patterns of MuLV and T2/Onc insertions in *Fli1* also reflect differences in mutational mechanism. All of the MuLV insertions were upstream in the sense and antisense orientation, acting as promoter and enhancer mutations, respectively. Again, some of the upstream MuLV insertions were a considerable distance from *Fli1* but 3 sense insertions were within the first exon of an Ensembl EST gene for which there is no associated Ensembl gene transcript, suggesting the presence of an additional, unannotated *Fli1* gene transcript (Figure 3.6B). While 2 of the SB insertions were also upstream in the sense orientation, the remaining 3 were in the first intron in the sense orientation, most likely producing an overexpressed, N-terminally truncated transcript in which none of the functional domains are deleted.

Similarly, both mutagens were found upstream of *Erg* (Figure 3.6C) in the sense orientation, and MuLV insertions also occurred upstream in the antisense orientation. However, 19 intragenic T2/Onc insertions were found in the sense orientation within a 1,531 bp region in the 1st intron, while 3 MuLV sense insertions were found in the 2nd intron. Like *Etv6*, *Erg* encodes a SAM_PNT and an Ets domain and, assuming that the insertions are producing N-terminally truncated transcripts, the T2/Onc insertions may give rise to truncated proteins containing both domains, while the MuLV insertions would give rise to proteins with a disrupted SAM_PNT domain but full-length Ets domain. The



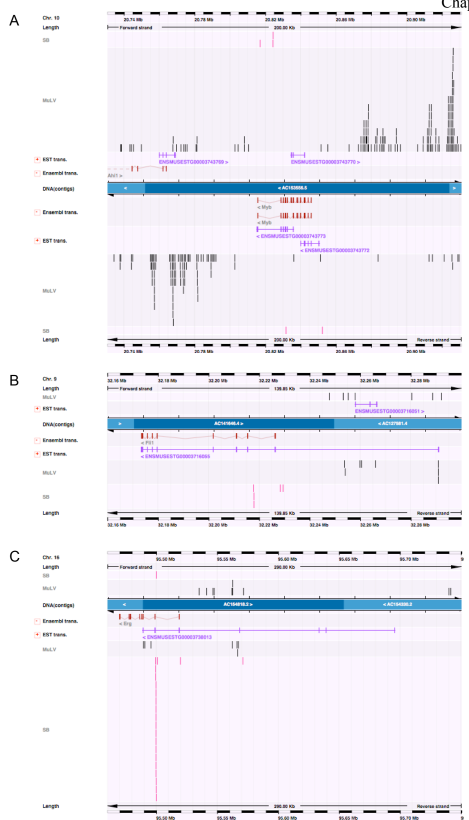


Figure 3.6. Variation in the distribution of MuLV and T2/Onc insertions in *Myb* (A), *Fli1* (B) and *Erg* (C) may reflect differences in the mechanisms of mutagenesis. MuLV insertions are shown in black, T2/Onc insertions are shown in pink. Ensembl gene transcripts are shown in red and blue. ESTs are shown in purple. Insertions above and below the blue bar labelled DNA(contigs) are in the forward and reverse orientation, respectively.

apparent presence of functional domains in the Erg truncations, but not in the Etv6 truncations, may reflect the fact that Erg is a transcriptional activator (Duterque-Coquillaud et al., 1993), whereas Etv6 is a transcriptional repressor (see above). The closely aligned T2/Onc insertions in the first intron of Erg could be contaminants, but it is also possible that, in the absence of enhancer activity, the production of an overexpressed, N-terminally truncated protein is the most effective way to mutate the gene. The SAM PNT domain is involved in the formation of heterodimers with other Ets proteins. Erg/Ets-2 dimer formation prevents Ets-2 from acting as a transcriptional activator of *Mmp3* (Basuyaux *et al.*, 1997; Buttice *et al.*, 1996) and dimerisation may prevent Ets proteins from binding to genomic DNA target sites (Carrere et al., 1998). Therefore, it is possible that the 3 MuLV insertions in the 2nd intron that appear to disrupt the SAM PNT domain prevent dimerisation and so cause an increase in the transcriptional activity of Erg and other Ets proteins that bind to Erg. The high proportion of MuLV promoter and enhancer insertions suggests that this may be a less efficient way of upregulating the gene, although it could also reflect the tendency of MuLV to insert close to transcription start sites. All of the MuLV insertions, and one of the T2/Onc insertions, that were identified upstream in the sense orientation were greater than 40 kb upstream of the Erg gene, which is a considerable distance for promoter mutation. However, the insertions resided within an Ensembl EST gene that overlaps with the Erg gene, suggesting that there may be an additional, unannotated, Erg transcript that is targeted by insertional mutagenesis.

In summary, differences in the distribution of MuLV and T2/Onc insertions may help to distinguish oncogenes and tumour suppressor genes. Intragenic insertions in oncogenes are more likely to be localised, since specific mutations, such as those described in *Notch1*, may be required for oncogenesis. However, it is more likely that tumour suppressor genes can be inactivated in multiple ways, and the distribution of insertions may be less defined, as demonstrated in *Ikzf1*, where MuLV and T2/Onc insertions were scattered throughout the gene (see also Section 3.4.3, below).

3.4.3 Identification of tumour suppressor genes inactivated by MuLV

Although retroviral insertional mutagenesis identifies predominantly oncogenes, tumour suppressor genes also featured in the list of candidate cancer genes. The most prevalent, with 93 insertions, was *Ikaros (Ikzf1)*. *Ikaros* encodes a haematopoietic-specific zinc

finger DNA-binding domain protein that regulates B- and T-cell differentiation (Georgopoulos *et al.*, 1997). Mice with reduced *Ikaros* expression develop leukaemias and lymphomas with complete penetrance (Winandy *et al.*, 1995). 31 insertions were also identified in *Aiolos (Ikzf3)*, which is also a member of the Ikaros family. Ikaros and Aiolos appear to play dual roles in T cell development since they can regulate the activation or repression of lineage-specific genes through the formation of chromatin remodelling complexes in lymphocytes (Georgopoulos, 2002; Kim *et al.*, 1999). However, the importance of *Ikaros* and *Aiolos* as tumour suppressor genes is demonstrated by their frequent deletion in paediatric acute lymphoblastic leukaemia (ALL) (Mullighan *et al.*, 2007), and *Ikaros* is deleted in 83.7% of ALLs containing the *BCR-ABL* translocation (Mullighan *et al.*, 2008).

Other implicated tumour suppressor genes in the candidate list included Wwox (22) insertions), E2f2 (17 insertions), Mobkl2a (Mob1; 16 insertions), Xrcc6 (Ku70; 10 insertions), Ovca2 (9 insertions) and Adrbk1 (Grk2; 8 insertions). Wwox spans the human FRA16D fragile site and is frequently disrupted in human cancers (Bednarek et al., 2000). $Wwox^{+/-}$ mice develop significantly more ethyl nitrosurea (ENU)-induced lung tumours and lymphomas than wildtype mice, suggesting that *Wwox* can act as a haploinsufficient tumour suppressor gene (Aqeilan et al., 2007). Loss of E2f2 accelerates Myc-induced lymphomagenesis in mice (Opavsky et al., 2007), while Xrcc6-deficient mice develop thymic and disseminated T cell lymphomas (Li et al., 1998). MOB1 activates the tumour suppressor LATS1, which is inactivated in human sarcomas and ovarian and breast cancers (Hergovich et al., 2006), and inactivating insertions in Mob1 may therefore contribute to tumourigenesis by preventing the activation of *Lats1*. OVCA2 is one of two adjacent genes that are frequently deleted in human ovarian, brain, breast and lung tumours (Schultz et al., 1996). GRK2 acts in a negative feedback loop to control TGF β signal transduction, which is often dysregulated in cancer (Ho et al., 2005), and was shown to significantly reduce proliferation of thyroid cancer cell lines (Metaye et al., 2008).

There is no straightforward approach for identifying candidate tumour suppressor genes because, while they are likely to contain only intragenic insertions, some oncogenes, such as *Notch1* and *Pim1*, are also mutated predominantly by intragenic insertions. However, as mentioned in Section 3.4.2, insertions in oncogenes are more likely to form specific oncogenic mutants and may therefore be more localised within the gene. In addition,

tumour suppressor genes are likely to contain multiple insertion sites within the same tumour, as described for *Pten* and *Qsk* in Section 3.3, because both copies of the gene must be inactivated. This does not hold for haploinsufficient tumour suppressor genes, which require only one inactivating insertion for tumourigenesis and are therefore more likely to be identified by insertional mutagenesis than genes requiring an insertion in both genes. 14 tumours contained multiple insertions within *Ikzf1*, while 1 contained multiple insertions in *Ikzf3*. However, none of the other genes so far discussed in this section were mutated by multiple insertions. This suggests either that they are haploinsufficient tumour suppressor genes, as demonstrated for *Wwox*, or that the coverage of the screen was too low, such that multiple insertions occurred but were not identified. To further complicate matters, oncogenes may also contain multiple insertion sites within the same tumour, either because the gene is a preferential target site for the virus, or because upregulation of both gene copies provides an even greater growth advantage to the cell. However, taken together, the distribution of insertions and the number of insertion sites within each tumour can help to identify potential tumour suppressor candidates.

Smg6 contained 53 insertions and was mutated by multiple insertions in 4 tumours. All insertions were intragenic and were distributed throughout the gene in both orientations, although many were clustered within a single intron (Figure 3.7A). The human orthologue, EST1A/SMG6, has been shown to interact with telomerase and the human telomerase reverse transcriptase (hTERT) (Redon et al., 2007), and overexpression in kidney 293T cells leads to progressive telomere shortening (Snow et al., 2003). Early in tumourigenesis, telomere shortening contributes to chromosomal destabilisation and therefore promotes genomic instability and cancer progression (see Sections 1.2.3.3 and 1.3.3.1). A telomere maintenance mechanism is subsequently activated and is required for tumour progression and immortality (Stewart, 2005). Therefore, SMG6 could play an oncogenic or tumour suppressive role in this process. SMG6 is also an essential factor in the nonsense-mediated mRNA decay (NMD) pathway, which degrades mRNAs carrying premature stop codons and regulates the expression of naturally occurring transcripts, including those involved in cell cycle progression (Rehwinkel et al., 2005). SMG6 may therefore play a tumour suppressive role by negatively regulating oncogene expression via NMD. The presence of both sense and antisense insertions in the 9th intron suggests that they are involved in C-terminal truncation of the gene product. This would result in the removal of the PINc nucleotide binding domain (IPR006596), which is required for degradation of single-stranded RNA, and an inactivated domain has been shown to inhibit

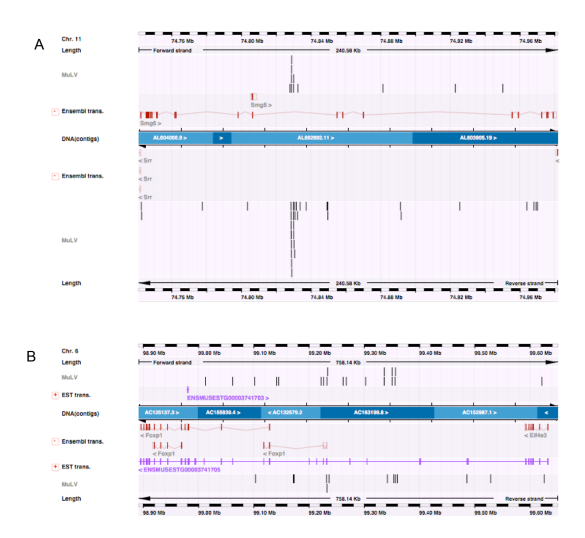


Figure 3.7. *Smg6* (A) and *Foxp1* (B) are putative tumour suppressor genes identified by MuLV insertional mutagenesis. MuLV insertions are shown in black. Ensembl gene transcripts are shown in red. ESTs are shown in purple. Insertions above and below the blue bar labelled DNA(contigs) are in the forward and reverse orientation, respectively.

NMD in *Drosophila* (Glavan *et al.*, 2006). This suggests that abrogation of NMD activity is the mechanism by which *Smg6* contributes to MuLV-induced lymphomagenesis. *SMG6* also resides within a deleted region containing 383 genes identified in 2.6% of human B-cell ALLs and 4.0% of T-cell ALLs in Mullighan *et al.* (2007).

Rassf2 contained 12 insertions, 2 of which were identified in a single tumour. Rassf2 is a negative regulator of Ras that is silenced by CpG island hypermethylation in a range of cancers, including gastric (Endoh *et al.*, 2005), liver (Nishida *et al.*, 2008), breast and lung (Cooper *et al.*, 2008; Kaira *et al.*, 2007). It has been shown to prevent cell transformation in primary colorectal cancers (Akino *et al.*, 2005).

Foxp1 contained 29 insertions, including 2 insertions in one tumour (Figure 3.7B). Overexpression of Foxp1 is associated with poor prognosis in lymphomas (Banham *et al.*, 2005), but loss of Foxp1 expression in breast cancer is also associated with poor prognosis (Fox *et al.*, 2004) and Foxp1 maps to a region on chromosome 3 (p14.1) that frequently shows loss of heterozygosity in a range of human cancers (Banham *et al.*, 2001). This suggests that Foxp1 can act as an oncogene or a tumour suppressor gene, depending on the tissue type (Koon *et al.*, 2007). The distribution of insertions in and around Foxp1 suggests that many are upstream, and therefore that the gene is being upregulated, which is consistent with the oncogenic role of Foxp1 in lymphomas. However, there is an Ensembl EST gene with no associated Ensembl gene transcript that spans the entire Foxp1 CIS, suggesting that the insertions could in fact be intragenic.

This section suggests that the MuLV screen can be helpful in identifying candidate tumour suppressor genes. However, computational analysis of insertions in and around genes can only provide an indication of whether a candidate cancer gene is likely to be oncogenic or tumour suppressive, and analysis of gene expression in MuLV-induced tumours, followed by functional validation, is essential for further confirmation.

3.4.4 Identifying retroviral insertions in regulatory features

The orientation and distribution of insertions around genes helps to identify promoter and enhancer mutations and insertions that prematurely terminate gene transcription. However, it is also possible that insertions could disrupt a gene by inserting into regulatory elements, thereby preventing the binding of transcriptional activators or repressors. In Ensembl version 45, regulatory features were available for the human, but not the mouse, genome. Features were built using 3 genome-wide anchor datasets: DNaseI hypersensitivity sites identified by ChIP-seq analysis (Boyle et al., 2008), CCCTC-binding factor (CTCF) binding sites identified by ChIP-Chip (Kim et al., 2007b), and histone 3 lysine 4 tri-methylation (H3K4me3) also identified by ChIP-chip. ChIP-chip, ChIP-seq and DNaseI hypersensitivity (a marker of open chromatin) are discussed in Section 1.3.5. The DNaseI hypersensitivity sites were identified in CD4+ T cells, but most were also found in CD8+ T cells and B cells and around 10% were lymphocyte-specific. This dataset is therefore particularly relevant to the MuLV screen, which generated predominantly lymphomas (see Section 2.2.1). CTCF is an insulator protein that prevents the spread of heterochromatin and prevents enhancers from activating unrelated promoters. CTCF binding sites were identified in primary human fibroblasts but were largely conserved across cell types (Kim *et al.*, 2007b). The histone modification H3K4me3 is associated with transcription start sites of active genes. 5 supporting ChIP-Chip datasets of histone modifications (H4K20me3, H3K27me3, H3K36me3, H3K79me3 and H3K9me3) were also used. In the Ensembl regulatory build, overlapping elements identified in each analysis were merged into a single element, and each element was classified based on the datasets in which it was identified. Elements associated with DNaseI hypersensitivity and H3K36me3 were classified as promoter-associated elements, while elements associated with DNaseI and either H3K4me3 or H3K79me3, or DNaseI and H3K4me3 and either CTCF or H3K36me3, were classified as gene-associated elements. It is worth noting that elements in the regulatory build define regions that are much larger than individual transcription factor binding sites and only define regions that are likely to be involved in regulation.

Since the regulatory build was only available for the human genome, elements and their classifications were downloaded from ftp://anonymous@ftp.ensembl.org/pub/release-45/homo_sapiens_45_36g/data/reg_build/ and were mapped to the NCBI m36 mouse genome assembly using UCSC LiftOver (http://genome.ucsc.edu/cgi-bin/hgLiftOver). Out of 113,230 features, 77,446 (68.4%) were successfully mapped to the mouse build. Of those that failed to convert, 30 were split (i.e. they mapped to 2 locations in the mouse genome due to breaks in synteny), 466 were duplicated, 1,285 were partially deleted and 34,003 were completely deleted. It is not surprising that such a large number of elements did not map, since only a fraction of the human and mouse genomes align confidently and

LiftOver uses these alignments to find corresponding positions between human and mouse.

It has been estimated that 32-40% of human transcription factor binding sites are not functional in rodents, suggesting a high evolutionary turnover of sites (Dermitzakis and Clark, 2002). However, highly conserved elements have also been identified, and are particularly prominent around vertebrate developmental genes (Woolfe *et al.*, 2005). The parameters of LiftOver were set such that the minimum ratio of bases that must remap was 0.1, and therefore it is possible that some of the mapped elements are poorly conserved. However, while this could suggest that the elements are not functional in the mouse, there is evidence to suggest that regulatory function can be conserved without sequence similarity, e.g. in the case of the RET locus in humans and zebrafish (Fisher *et al.*, 2006). Therefore, a decision was made to be inclusive, rather than to reject elements of low sequence similarity.

2,036 (10.1%) of the 20,114 insertion sites mapped to regulatory features, of which 21 (0.02%) were promoter-associated and 1,394 (68.5%) were gene-associated. This compared with 743 (9.9%) of the 7,518 insertion sites associated with the 439 candidate cancer genes, of which 11 (0.02%) were promoter-associated and 443 (59.6%) were gene-associated. 228 candidate genes had insertions within regulatory features. There was no significant difference (P=0.38) between the number of insertions assigned to candidate cancer genes in regulatory features and the number of other insertions in regulatory features. However, insertions assigned to candidate cancer genes were under-represented in gene-associated elements (P=7.48x10⁻¹¹). This result was surprising since it might be assumed that oncogenic insertions would be more likely to be associated with regulatory elements.

Since many insertions may map to the same regulatory region, therefore skewing the results, the number of regions containing insertions was also counted. Insertions were identified in 1,483 regulatory features, of which 14 (0.94%) were promoter-associated and 971 (65.5%) were gene-associated. Insertions associated with candidate cancer genes were identified in 343 regulatory features, of which 5 (1.5%) were promoter-associated and 160 (46.6%) were gene-associated. Once again, gene-associated elements were under-represented among insertions associated with candidate cancer genes (P=6.00x10⁻¹⁷). Counting the number of regulatory features of each type that contained insertions

revealed an under-representation of features associated with H3K36me3 ($P=2.04 \times 10^{-3}$), H3K4me3 ($P=3.38 \times 10^{-15}$), H3K79me3 ($P=1.65 \times 10^{-4}$) and DNaseI hypersensitivity sites (P=0.012). All significance tests were performed using the Chi-squared test for independence. Interestingly, DNaseI hypersensitivity and all of the histone modifications stated above are known to be associated with active genes, while those histone modifications that showed no significant difference (H3K27me3, H3K9me3 and H4K20me3) are associated with gene repression (Barski *et al.*, 2007). H3K4me3 and H3K27me3 have also been shown to be associated with active genes and silent genes, respectively, in human T cells (Roh *et al.*, 2006), which is of particular relevance to this MuLV dataset of lymphomas. Insertions that are not associated with candidate cancer genes are less likely to be oncogenic, and their over-representation in regulatory features associated with active genes may reflect the preference of MuLV for inserting within active genes. Since none of the regulatory features are over-represented among candidate genes, it appears that disruption of regulatory features may not be a common mechanism of mutagenesis of the MuLV retrovirus.

3.4.5 Expression analysis of MuLV-induced tumours

Computational approaches can be used to predict candidate cancer genes and the likely mechanisms of mutation, but these must be confirmed using experimental methods. Gene expression analysis is a useful tool towards validating candidates, since it is expected that genes that are disrupted by MuLV will be differentially expressed in insertion-containing tumours versus those that do not contain insertions. Although widespread expression analysis has not been performed on the MuLV-induced tumours, expression data was available for 18 tumours. The analysis was performed by David Adams using high density Nimblegen 5045 MM8 60mer expression arrays, where MM8 is the mouse build (the UCSC equivalent to NCBI m36) and 60mer is the length of the oligonucleotide probes on the array. The array covers 18,879 transcripts with unique RefSeq NM accession numbers, and 6,751 with RefSeq XM accession numbers. NM and XM refer to reported and predicted transcripts, respectively. Each NM transcript has three probes, while 1,861 XM transcripts have 3 probes and the rest have 2 probes. The normalised expression values across all probes in each transcript, as provided by Nimblegen, were used in this analysis.

81 candidate cancer genes from the MuLV screen contained MuLV insertions in at least 1 of the 18 tumours, and 20 contained insertions in at least 2 tumours. RefSeq accession numbers, Entrez Gene identifiers and MGI symbols were extracted from BioMart (version 49) for each of the 439 candidate cancer genes from the MuLV screen. All of the genes except *mmu-mir-17*, *ENSMUSG00000074675*, *Rnf157* and *Pvt1* were identified on the array. Genes directly flanking each candidate gene were identified using the coordinates of all genes in Ensembl version 45. For candidate genes with insertions in 2 or more tumours, a two-sided *t*-test was performed to determine whether the level of expression in tumours containing an insertion in the gene. The results are shown in Table 3.4. The *t*-test was also performed on genes flanking the candidate cancer genes, in order to ascertain whether the insertions had been assigned to the correct gene.

Only 1 of the candidates, *Trpm1*, showed significant differential expression in tumours containing an insertion compared to those that did not, and the insertions appeared to cause a decrease in gene expression. Loss of *Trpm1*, also known as melastatin, correlates with metastatic potential in human and mouse melanoma cells (Deeds et al., 2000). Interestingly, the insertions in tumours used in this analysis were 11.7 kb and 21.2 kb upstream of *Trpm1*, which suggests either that there is a longer transcript that is not annotated in Ensembl, or that the gene is disrupted by insertion into upstream regulatory elements, although the insertions did not overlap with regulatory features in the dataset described in Section 3.4.4. Although the difference was not significant, the mean expression level in insertion-containing tumours was at least 2-fold higher than in other tumours for genes Notch1, Rasgrp1, Pik3r5, Jundm2, Pim1 and Rras2. None of the genes flanking the candidate genes showed significant differential expression, but Spon1, Lrrc8b and Fos, which flank genes Rras2, Lrrc8c and Jundm2, respectively, had a mean expression level that was at least 2-fold higher in insertion-containing tumours. Due to their enhancer activity, MuLV insertions can have long-range effects, and therefore it is possible that Spon1 and Fos are also affected by insertions disrupting Rras2 and Jundm2, respectively. On the other hand, *Lrrc8b* showed a greater difference in expression than did *Lrrc8c*, and it is possible that *Lrrc8b* is the true candidate cancer gene in this region.

The scale of this analysis was too small to provide any definitive evidence that the correct candidate cancer gene has been selected. The results suggest that there may not be a strong association between insertion-containing genes and higher expression levels. It is

	Number of tumours with					
Gene name	insertions	Mean 1	SD 1	Mean 2	SD 2	P-value
Notch1	2	5815.239	6599.7549	1078.072	126.6176	0.1773
Mad1l1	2	4708.679	875.3291	5297.206	1253.8023	0.6712
Rasgrp1	2	14100.646	1935.705	7066.692	4240.0499	0.1195
Pik3r5	2	3813.572	2155.5715	800.739	2405.31807	0.1531
Jundm2	2	3250.289	3778.6357	579.06	571.0842	0.2024
Hnrpf	2	36521.059	7138.7373	37617.657	2704.6885	0.8494
Trpm1	2	153.399	27.9747	398.287	37.2045	0.0002
B3gnt2	2	2291.355	1024.3581	1947.636	857.7561	0.7632
Spn	2	1882.812	264.2865	1629.536	646.2697	0.7024
Hibadh	3	4449.752	1192.4941	5461.04	1742.7667	0.6032
Pim1	3	10856.253	13210.2491	1640.201	962.0191	0.4871
Myb	4	11458.869	5636.9595	8207.643	2707.9078	0.5177
Lrrc8c	4	1939.271	247.5314	1891.16	488.6951	0.927
Evi5	4	867.045	875.3993	797.081	706.3382	0.9485
Ccnd3	6	8402.866	4375.7706	7656.238	2977.1405	0.8832
Rras2	7	21102.81	4947.9028	6690.916	8678.9839	0.16
Мус	7	21190.356	7534.8977	19642.481	8734.1825	0.8881
Gfi1	8	7495.101	2592.9975	4365.898	2676.8333	0.3846

Table 3.4. Gene expression values for candidate cancer genes in insertion-containing tumours compared with tumours that do not contain insertions. Mean 1 and SD 1 are the mean and standard deviation of expression levels for genes in tumours containing insertions, and Mean 2 and SD 2 are the mean and standard deviation for genes in tumours that do not contain insertions. *P*-values were calculated using the *t*-test.

possible that, over time, insertions may lose their ability to disrupt cellular genes by promoter or enhancer mutation, e.g. because the retroviral LTRs are silenced by hypermethylation. Alternatively, since tumours are heterogeneous, and the tumour samples may also contain stromal cells, the effect of the insertion may be diluted by the presence of wildtype gene expression in contaminating cells. An analysis of gene expression across all tumours, with replicates, is required to substantiate these suggestions.

3.5 Identification of co-operating cancer genes in the MuLV dataset

As discussed in Section 1.4.2.1.3, there are two main approaches for identifying collaborating cancer genes using insertional mutagenesis. By conducting the screen in genetically engineered mice in which oncogenes are overexpressed or tumour suppressor genes are inactivated, it is possible to identify genes that collaborate with the gain or loss, respectively, of those cancer genes in oncogenesis. An alternative approach involves identifying co-occurring CIS genes in individual tumours. Both approaches have been employed to analyse the MuLV dataset of 439 statistically significant CIS genes.

3.5.1 Genotype-specific cancer genes

The retroviral screen described in this thesis was performed on mice deficient in a range of tumour suppressor genes (see Section 2.2.1). For each gene identified using the kernel convolution-based method for determining significant CISs, the number of insertions assigned to the gene and the number of insertions of each genotype were counted. See Section 2.9 for a description of the methods used to assign insertions to genes. The 2-tailed Fisher Exact Test was used to determine whether there was any significant difference between the number of insertions of a particular genotype within each gene and the number of the genome, and also between the number of insertions of a particular genotype and the number of wildtype insertions within a gene compared to the proportions in the rest of the genome:

a	b
С	d

a = Number of insertions of a given genotype assigned to the gene

b = Number of insertions of a given genotype not assigned to the gene

c = Number of insertions of other genotypes assigned to the gene, or number of wildtype insertions assigned to the gene

d = Number of insertions of other genotypes not assigned to the gene, or number of wildtype insertions not assigned to the gene

Significance tests were performed for each genotype, and also for groups of genotypes to increase the power of the analyses. For example, in order to test for a significant bias towards any p21-deficient background, all insertions on a p21-null homozygous or heterozygous background, and all insertions on a homozygous or heterozygous p21-null and p27-null double mutant background, were counted. In order to account for multiple testing, the R package QVALUE (Storey and Tibshirani, 2003) was used to generate a qvalue for each test. The q-value is a measure of the minimum false discovery rate incurred if the test is called significant, where the false discovery rate is the number of false positives divided by the number of significant tests. This differs from the *P*-value, which is a measure of the minimum false positive rate incurred when the test is called significant, where the false positive rate is the number of false positives divided by the number of true null tests. Using a P-value of 0.05, 5% of tests will be called significant when they are in fact null, which would result in a very high number of false positives if a large number of tests were performed. On the other hand, using a q-value of 0.05, 5% of the tests that have been called significant will be false positives, which is more manageable for large numbers of tests. The QVALUE bootstrap method was used to estimate the overall proportion of true null hypotheses, since this method is deemed most appropriate for situations in which the distribution of null *P*-values is skewed towards a value of 1, as is the case in this analysis. The shape of the distribution also means that the calculated q-values are very conservative, but this method is still more inclusive than using, for example, the Bonferroni correction. The most significant tests, where the qvalue is less than 0.05 for the comparison with insertions of all other genotypes and/or with only wildtype insertions, and that therefore suggest a bias towards, or away from, a particular tumour genotype, are presented in Tables 3.5A and 3.5B, respectively. Results are grouped by gene, with genes ordered according to the most significant association obtained from the comparison with insertions of all other genotypes. The two methods gave similar results, although the comparison of a given genotype versus wildtype gave fewer significant results. This may be because some genes always require the co-

A				-	vs. insertio	ons of all		
					other genotypes		vs. wildtype	insertions
6	Frankl Grant ID	Total number of	Construct	Insertions of given			<i>P</i> -value	
Gene Evi5	Ensembl Gene ID ENSMUSG00000011831	insertions	Genotype p27 all	genotype 134	P-value 1.87E-29	q-value 8.19E-27	1.77E-15	q-value 4.50E-13
EVIS	ENSMUSG0000011831			100	1.87E-29 1.57E-10	3.45E-08	2.30E-06	4.50E-13 5.06E-04
			p21 all	71	7.83E-10	3.45E-08 1.72E-07	2.30E-06 7.17E-07	1.57E-04
			p21, p27	66	2.93E-10	1.29E-07	7.91E-07	1.57E-04 3.47E-04
			p27					
			p27 ko	40	3.89E-08	1.71E-05	9.34E-07	4.10E-04
Gfi1	ENSMUSG0000029275		p21 ko, p27 ko	35 127	1.38E-06 2.44E-26	3.03E-04 5.35E-24	8.50E-06 2.05E-15	1.86E-03 4.50E-13
GIII	ENSMUSG0000029275		p27 all	77				
			p21, p27		1.01E-12	4.44E-10	1.40E-09	6.16E-07
			p21 all	102	1.19E-11	5.23E-09	7.63E-08	3.35E-05
			p21 ko, p27 ko	37	6.98E-08	3.07E-05	2.07E-07	9.10E-05
			p27 ko	35	4.21E-06	9.24E-04	8.24E-06	1.81E-03
			p27	55	1.27E-05	2.79E-03	3.97E-05	8.71E-03
	ENOLUS 000000000000		p21 ko, p27 het	35	8.59E-05	3.77E-02	8.13E-05	3.57E-02
Map3k8	ENSMUSG0000024235		p16 ko, p19 ko	27	2.77E-21	1.21E-18	5.59E-10	2.45E-07
			p16 all	28	1.67E-20	7.33E-18	5.33E-09	2.34E-06
			p16, p19	28	1.67E-20	7.33E-18	5.33E-09	2.34E-06
	ENOLUS 00000000000		p19 all	31	2.13E-09	7.57E-07	6.05E-03	5.67E-01
Мус	ENSMUSG0000022346		p27 all	74	6.14E-09	8.98E-07	3.56E-06	5.20E-04
Myb	ENSMUSG0000019982		p27 all	51	1.30E-06	1.35E-04	1.37E-03	1.00E-01
Art2b	ENSMUSG0000030651		p27 all	8	1.53E-06	1.35E-04	1.99E-03	1.22E-01
Pvt1	ENSMUSG0000072566		p27 all	55	1.72E-05	1.26E-03	2.23E-03	1.22E-01
A530013C23Rik	ENSMUSG0000006462		p16 all	15	1.75E-05	3.57E-03	7.46E-04	8.08E-02
			р16, р19	15	1.75E-05	3.57E-03	7.46E-04	8.08E-02
			p16 het, p19 het	6	1.22E-04	5.36E-02	8.57E-02	1.95E-02
Ccnd3	ENSMUSG0000034165		p53 ko	59	2.23E-05	9.79E-03	1.75E-06	7.68E-04
			p53	59	5.64E-05	2.47E-02	3.18E-06	7.68E-04
Zfp438	ENSMUSG0000050945		p19 ko	27	2.51E-05	3.67E-03	7.06E-03	8.13E-01
	ENSMUSG0000059894		p27 all	36	3.30E-05	2.07E-03	1.80E-02	6.64E-01
Pim1	ENSMUSG0000024014		p21 all	29	4.81E-05	7.04E-03	4.72E-03	4.14E-01
Rras2	ENSMUSG00000055723		p27 all	43	5.42E-05	2.97E-03	2.19E-04	2.40E-02
	2358 ENSMUSG0000052248		p16 ko, p19 ko	11	4.48E-04	3.93E-02	8.20E-03	5.14E-01
Zeb2	ENSMUSG0000026872		p16 ko, p19 ko	11	4.48E-04	3.93E-02	8.20E-03	5.14E-01
Mycn	ENSMUSG0000037169	81	p27 all	19	6.07E-04	2.96E-02	2.80E-04	2.46E-02
Ahi1	ENSMUSG0000019986	124	p27 all	25	9.15E-04	4.02E-02	5.17E-02	8.61E-01

В			-	<i>vs</i> . insertio other gen		<i>vs</i> . wildtype	insertions
Gene	Ensembl Gene ID	Total number of insertions Gen	Insertions of given otype genotype	<i>P</i> -value	<i>q</i> -value	<i>P</i> -value	<i>q</i> -value
Evi5	ENSMUSG0000011831	466 p19 ko	71	2.43E-07	1.07E-04	7.20E-03	8.13E-01
Gfi1	ENSMUSG0000029275	458 p19 all p19 ko	114 76	9.17E-07 1.51E-05	1.34E-04 3.31E-03	1.56E-01 9.18E-02	1.00E+00 1.00E+00
Rasgrp1	ENSMUSG00000027347	237 p16 all p16, p1 p16 ko,		2.44E-05 2.44E-05 1.73E-04	3.57E-03 3.57E-03 2.53E-02	2.76E-05 2.76E-05 1.65E-04	6.06E-03 6.06E-03 2.41E-02
Ccnd3	ENSMUSG00000034165	206 p16 ko, p16 all p16, p1	. 7	9.19E-05 2.27E-04 2.27E-04	2.02E-02 2.49E-02 2.49E-02	1.25E-01 3.15E-01 3.15E-01	1.00E+00 1.00E+00 1.00E+00
Ikzf1	ENSMUSG0000018654	93 p53 p53 ko	4	2.75E-04 4.14E-04	5.30E-02 7.49E-02	5.99E-05 6.54E-05	8.76E-03 9.58E-03
Zmiz1	ENSMUSG0000007817	62 p16 ko, p53 p53 ko	p19 ko 2 0 2	5.79E-03 6.36E-03 9.62E-03	2.06E-01 1.99E-01 3.02E-01	9.60E-05 4.95E-05 5.37E-05	2.11E-02 8.76E-03 9.58E-03

Table 3.5. Genes containing an over-representation (A) or underrepresentation (B) of insertions on a given tumour background compared with all other backgrounds and compared with wild-type insertions only. All genes identified in tests with a q-value of less than 0.05 for one or both methods are shown. p27 all = all genotypes that include a mutation in p27 (homozygous or heterozygous, single or double mutant); p27 = single homozygous or heterozygous mutation of p27; p27 ko = single homozygous mutation of p27. All other genotypes follow the same rules. Genes are listed in order of decreasing significance (increasing P-value) with respect to the comparison with insertions from all other genotypes. operation of other cancer genes and are therefore rarely or never mutated on a wildtype background.

The most significant result was the bias of insertions in Evi5/Gfi1 towards p27-deficient genetic backgrounds. Interestingly, while insertions within this locus were identified in an MuLV screen performed on p27-deficient mice by Hwang and coworkers (2002), no significant difference was observed between the frequency of insertions in $p27^{-/-}$ and wildtype mice. The screen, which involved 50 tumours, was smaller than the screen described in this thesis, and therefore the difference may reflect the increased power of this larger dataset and supports the use of larger insertional mutagenesis screens for identifying co-operating oncogenes. In accordance with the observations of Hwang et al. (2002), insertions in Myc showed a significant bias towards p27-deficient genotypes. This is also supported by the finding that p27-deficient lymphomas show an increased frequency of Mvc activation, and that Mvc-induced tumourigenesis may be enhanced upon loss of p27 (Martins and Berns, 2002). Mycn, which is structurally and functionally related to Myc, was also associated with p27-deficient tumours. MYCN amplification in human neuroblastomas is associated with poor prognosis (Seeger et al., 1985). Low expression of p27 is also correlated with poor prognosis in patients with neuroblastoma, yet p27 expression and MYCN amplification are prognostic independent and are not significantly associated in neuroblastomas (Bergmann *et al.*, 2001). While this seems to suggest that disrupted p27 and MYCN are not collaborating in neuroblastoma, it does indicate that the genes may act in different genetic pathways, as is generally expected for genes that collaborate in tumourigenesis.

Insertions in *Map3k8* showed the most significant bias to the $p16^{-r}p19^{-r}$ (or *Cdkn2a*^{-/-}) tumour genotype. *Map3k8* has previously been identified as *Cdkn2a*^{-/-}-specific in an MuLV screen performed on 115 mice (Lund *et al.*, 2002). Activation of Mek by Map3k8 in the mitogen-activated protein kinase (MAPK) signalling pathway (Salmeron *et al.*, 1996) induces p16 and p53, resulting in the permanent arrest of mouse fibroblasts (Lin *et al.*, 1998). However, in the absence of p16 or p53, the activation of the MAPK cascade causes cells to undergo uncontrolled mitogenesis and transformation (Lin *et al.*, 1998).

Insertions affecting the gene encoding zinc finger E-box-binding homeobox 2 (Zeb2 or Sip1) were also significantly associated with the $Cdkn2a^{-/-}$ genotype. *SIP1* plays a role in replicative senescence, which controls the number of cell divisions in human somatic

tissues and so prevents the indefinite proliferation associated with tumour cells (Ozturk et al., 2006). Inactivation of SIP1 causes reactivation of the human telomerase reverse transcriptase (hTERT), resulting in the rescue of hepatocellular carcinoma cells from senescence arrest (Ozturk et al., 2006). Replicative immortality also requires the inactivation of Trp53 and p16 (Ozturk et al., 2006), therefore suggesting co-operation between p16 and SIP1 in tumourigenesis. Interestingly, all of the insertions within Zeb2 are flanking an internal gene, and most are in the upstream sense orientation, suggesting that they are promoter insertions that upregulate the internal gene. This internal gene is a natural antisense transcript that, when overexpressed in epithelial cells, prevents splicing of the Zeb2 5' UTR (Beltran et al., 2008). However, this is proposed to increase the levels of Zeb2 (Beltran et al., 2008), which conflicts with the observations described above. Zeb2 also directly represses cyclin D1, resulting in initiation of the epithelialmesenchymal transition (EMT), in which cells switch from a proliferative to an invasive state (Mejlvang et al., 2007). Cyclin D1 (Ccnd1) was also biased towards the Cdkn2a^{-/-} tumour background, albeit with lower significance ($P=2.16 \times 10^{-3}$, q=0.135). p16 binds to CDK4 and prevents it from forming a complex with cyclin D1, resulting in cell cycle arrest at the G1/S transition (Serrano et al., 1993). An enhanced gene ratio of CCND1:CDKN2A, i.e. a high copy number of CCND1 combined with deletion of CDKN2A, correlates with poor survival in patients with squamous cell carcinoma of the head and neck (Akervall et al., 2003), while the combined loss of p16 and overexpression of cyclin D1 has been observed in 49% of gastric carcinomas (Kishimoto et al., 2008). It therefore appears that *Ccnd1*, *Zeb2* and *Cdkn2a* may collaborate in tumourigenesis, where *Ccnd1* causes uncontrolled cell growth in the absence of *Cdnk2a*, and *Zeb2* represses *Ccnd1*, causing hyperproliferating cells to undergo EMT.

Insertions in the oncogene *Pim1* were associated with *p21*-deficient tumours. Phosphorylation of *p21* by Pim1 results in the cytoplasmic localisation (Wang *et al.*, 2002b) or stabilisation of p21 (Zhang *et al.*, 2007), and this is proposed to be a contributing factor in the tumourigenesis of cells overexpressing *Pim1* (Zhang *et al.*, 2007). However, the fact that *Pim1* mutagenesis is favoured in a *p21*-deficient background suggests that overexpression of *Pim1* alone cannot fully inactivate *p21* and the genes may have a more complex relationship that has not been elucidated. Insertions in *Runx1* were biased towards the *p53^{-/-}* genetic background. If a gene contains fewer insertions than expected in tumours bearing a particular inactivated tumour suppressor gene, this suggests that the CIS gene and the inactivated tumour suppressor gene may act in the same cancer pathway. Insertions in Zmiz1 and *Ikaros* (*Ikzf1*) were under-represented in $p53^{-/-}$ tumours. Zmiz1 is a transcriptional coactivator of p53 (Lee et al., 2007) and therefore, in the absence of p53, mutation of Zmiz1 does not provide any additional growth advantage. The results for Ikaros are more surprising since, in chemically induced murine lymphomas, allelic loss of Ikaros was more frequently found in $p53^{-1}$ lymphomas than in wildtype p53 lymphomas, suggesting cooperation in lymphomagenesis (Okano et al., 1999). Further functional evidence is required to validate this proposal. None of the genes identified as containing p53 binding sites in Section 3.2 were positively or negatively associated with the $p53^{-/-}$ tumour background. Some of the genes contained few insertions, and there may not be enough power to identify a significant association. For example, Chd1 did not contain any insertions in $p53^{-/-}$ tumours but only contained 7 insertions overall. Alternatively, the relationship with p53 may not be relevant in the setting of MuLV-induced lymphomagenesis. For example, p53-mediated upregulation of Notch1 contributes to cell fate determination (Alimirah et al., 2007), but in MuLV insertional mutagenesis, Notch1 is activated by truncating mutations and is therefore not dependent on p53 ($P=3.73\times10^{-4}$, $q=5.30 \times 10^{-2}$).

Further genes for which there was a strong bias towards or against a particular tumour genotype are listed in Table 3.5. Supporting evidence in the literature for the genes described above indicates that this may be a powerful method for identifying cooperating cancer genes.

3.5.2 Co-occurrence and mutual exclusivity of disrupted genes

The "genotype-specific" approach for identifying collaborating cancer genes only allows for the identification of collaborations with selected oncogenes or tumour suppressor genes. Identifying CIS genes that co-occur in tumours more often than expected by chance enables the identification of collaborations without any predetermined conditions. In Section 1.4.2.1.3, oligoclonality is cited as a potential disadvantage of this approach. However, since only significant CISs are utilised, insertions within these CISs are likely to be present, and to co-occur, in the dominant clone, rather than being rare insertions in less successful sublines of the tumour.

For each pair of CIS genes, the number of tumours that contained an insertion in both genes, or in one or other gene, was counted. The 2-tailed Fisher Exact Test was used to determine whether the number of tumours containing a co-occurrence of each gene pair was significantly different to the number expected by chance.

а	b
С	d

- *a* = Number of tumours containing an insertion in both genes
- b = Number of tumours containing an insertion in first gene
- c = Number of tumours containing an insertion in second gene
- d = Number of tumours containing an insertion in neither gene

To account for multiple testing, the R package QVALUE (Storey and Tibshirani, 2003), and specifically the *Bootstrap* method, was applied to all tests in which one or more co-occurrences were observed. Over- and under-represented co-occurrences with a *q*-value of less than 0.05 are shown in Tables 3.6A and 3.6B, respectively.

The most significant association was between genes A530013C23Rik and leukocytespecific protein tyrosine kinase *Lck*. Lck initiates a tyrosine phosphorylation cascade in lymphocytes that results in T-cell antigen receptor signal transduction, and it is overexpressed in lymphomas, breast cancer and colon cancer (for review, see Palacios and Weiss, 2004). Interestingly, insertions in both *Lck* and *A530013C23Rik* were biased towards a *Cdkn2a*^{-/-} genotype (*Lck*: *P*=9.12x10⁻⁴ and *q*=5.01x10⁻²; *A530013C23Rik*: see Table 3.5A), suggesting that all 3 genes collaborate in tumourigenesis.

Co-occuring insertions were also identified in *Lck* and signal transducer and activation of transcription 5b (*Stat5b*). LCK has been shown to interact with STAT5b in cells, and induces tyrosine phosphorylation and DNA-binding of STAT5b (Shi *et al.*, 2006). Exogenous expression of wildtype *STAT5b* increases LCK-mediated cellular transformation (Shi *et al.*, 2006). This is consistent with the pattern of insertions in and around *Stat5b*, which suggests that the gene is upregulated by promoter and enhancer mutations that increase the levels of the wildtype protein. Finally, activation of *Lck* was also significantly associated with activation of the c-src tyrosine kinase gene *Csk*. Csk negatively regulates Lck by phosphorylation of a C-terminal tyrosine (Tyr-505) (Bergman *et al.*, 1992). The distribution of insertions in and around *Csk* suggests that the gene is

1	۸
r	1

		Total tumours			Total tumours in	Number of tumours		
		in which Gene			which Gene 2	in which Gene 1 and		
Gene name 1	Ensembl ID 1	1 disrupted	Gene name 2	Ensembl ID 2	disrupted	Gene 2 disrupted	P-value	q-value
A530013C23Rik	ENSMUSG0000006462	43	Lck	ENSMUSG0000000409	26	10	2.47263E-08	8.59E-06
Ikzf1	ENSMUSG0000018654	93	Notch1	ENSMUSG0000026923	127	30	1.6337E-07	4.80E-05
Zfp438	ENSMUSG0000050945	51	Ntn1	ENSMUSG0000020902	59	14	3.41404E-07	9.66E-05
Pik3r5	ENSMUSG0000020901	64	Zfp438	ENSMUSG0000050945	51	14	1.02005E-06	2.52E-04
Epha6	ENSMUSG0000055540	20	Pim1	ENSMUSG0000024014	118	11	2.78357E-06	6.26E-04
Runx1	ENSMUSG0000022952	143	Rasgrp1	ENSMUSG0000027347	237	54	4.93807E-05	9.95E-03
Stat5b	ENSMUSG0000020919		Lck	ENSMUSG0000000409	26	5	5.55279E-05	1.06E-02
Nid1	ENSMUSG0000005397	12	Cd3e	ENSMUSG0000032093	8	3	7.27389E-05	1.32E-02
Lfng	ENSMUSG0000029570	33	Notch1	ENSMUSG0000026923	127	13	7.96187E-05	1.38E-02
Ppp2r5a	ENSMUSG0000026626	10	Vps13d	ENSMUSG0000020220	10	3	8.4762E-05	1.44E-02
Cd48	ENSMUSG0000015355		Arhqap26	ENSMUSG0000036452	11	3	1.16E-04	1.81E-02
Psma1	ENSMUSG0000030751		mmu-mir-17	ENSMUSG0000065508	33	4	1.13E-04	1.81E-02
Fgr	ENSMUSG0000028874	7	Dad1	ENSMUSG0000022174	16	3	1.15E-04	1.81E-02
Ubxd5	ENSMUSG0000012126	14	Thra	ENSMUSG0000058756	9	3	1.78E-04	2.56E-02
Rras2	ENSMUSG0000055723		Rasgrp1	ENSMUSG0000027347	237	75	1.75E-04	2.56E-02
Sdk1	ENSMUSG0000039683		Mns1	ENSMUSG0000032221	10	3	1.99E-04	2.77E-02
Zbtb7b	ENSMUSG0000028042		Notch1	ENSMUSG0000026923	127	7	2.15E-04	2.94E-02
Evi1	ENSMUSG0000027684		AB041803	ENSMUSG0000044471	14	5	2.23E-04	3.00E-02
Cd48	ENSMUSG00000015355		Fqfr2	ENSMUSG0000030849	14	3	2.52E-04	3.12E-02
Hycn1	ENSMUSG0000064267		Pyqm	ENSMUSG00000032648	4	2	2.53E-04	3.12E-02
D12Frtd553e	ENSMUSG00000020589		Ptpre	ENSMUSG00000041836	10	3	2.52E-04	3.12E-02
Eng	ENSMUSG00000026814		Gse1	ENSMUSG0000031822	25	3	2.67E-04	3.24E-02
Mvlc2pl	ENSMUSG0000005474		Bcl11a	ENSMUSG0000000861	13	3	2.71E-04	3.24E-02
mmu-mir-802	ENSMUSG00000076457		Rasgrp1	ENSMUSG0000027347	237	52	2.79E-04	3.28E-02
Csk	ENSMUSG00000032312		Lck	ENSMUSG00000000409	26	4	3.08E-04	3.57E-02
Smq6	ENSMUSG0000038290		Pik3r5	ENSMUSG00000020901	64	10	3.18E-04	3.63E-02
Fafr2	ENSMUSG00000030849		Plac8	ENSMUSG0000029322	11	3	3.43E-04	3.86E-02
A530013C23Rik	ENSMUSG0000006462		Hhex	ENSMUSG00000024986	16	5	3.66E-04	4.00E-02
A530013C23Rik	ENSMUSG0000006462		Exoc6	ENSMUSG00000024300	16	5	3.66E-04	4.00E-02
Spsb4	ENSMUSG00000046997			kENSMUSG00000045348	5	2	4.20E-04	4.23E-02
Arid3a	ENSMUSG00000019564		Rreb1	ENSMUSG00000039087	29	3	4.20E-04 4.21E-04	4.23E-02
Tcfap4	ENSMUSG0000005718		Jph4	ENSMUSG00000022208	5	2	4.20E-04	4.23E-02
Zfp608	ENSMUSG00000052713		Olfr56	ENSMUSG00000022208	7	3	4.06E-04	4.23E-02
Parvg	ENSMUSG00000022439		Prr6	ENSMUSG00000040328	6	2	4.50E-04	4.23E-02 4.29E-02
Frmd8	ENSMUSG0000022439 ENSMUSG00000043488		Ubxd5	ENSMUSG0000012126	14	2	4.50E-04 4.54E-04	4.29E-02
Frmd8	ENSMUSG0000043488		AB041803	ENSMUSG00000012128 ENSMUSG00000044471	14	3	4.54E-04 4.54E-04	4.29E-02 4.29E-02
Ubxd5					14	3	4.54E-04 4.54E-04	4.29E-02
AB041803	ENSMUSG00000012126		Scyl1	ENSMUSG00000024941	12	3	4.54E-04 4.54E-04	4.29E-02 4.29E-02
Gimap6	ENSMUSG00000044471 ENSMUSG00000047867		Scyl1 Bcl11a	ENSMUSG00000024941 ENSMUSG0000000861	12	3	4.54E-04 4.70E-04	4.29E-02 4.38E-02
Zmiz1 Cecr5	ENSMUSG0000007817		Notch1	ENSMUSG0000026923	127 11	18 3	4.96E-04	4.58E-02 4.59E-02
	ENSMUSG00000058979		Nfkb1	ENSMUSG0000028163			5.22E-04	
B3gntl1	ENSMUSG0000046605		Hhex	ENSMUSG0000024986	16	3	5.22E-04	4.59E-02
B3gntl1	ENSMUSG0000046605		Exoc6	ENSMUSG0000053799	16	3	5.22E-04	4.59E-02
Irf2bp2	ENSMUSG0000051495		Nsmce1	ENSMUSG00000030750	7	3	5.18E-04	4.59E-02
Jundm2	ENSMUSG0000034271		Runx1	ENSMUSG00000022952	143	28	5.67E-04	4.87E-02
Myb	ENSMUSG0000019982	247	Rras2	ENSMUSG0000055723	224	76	5.79E-04	4.92E-02

В

		Total tumours			Total tumours in	Number of tumours		
		in which Gene			which Gene 2	in which Gene 1 and		
Gene name 1	Ensembl ID 1	1 disrupted	Gene name 2	Ensembl ID 2	disrupted	Gene 2 disrupted	P-value	q-value
Ikzf1	ENSMUSG0000018654	93	Evi5	ENSMUSG0000011831	466	11	6.30E-14	3.01E-11
Evi5	ENSMUSG0000011831	466	Notch1	ENSMUSG0000026923	127	25	1.79E-11	8.06E-09
Ikzf1	ENSMUSG0000018654	93	Gfi1	ENSMUSG0000029275	458	16	1.41E-09	5.68E-07
Evi5	ENSMUSG0000011831		Rasgrp1	ENSMUSG0000027347	237	71	1.92E-09	7.35E-07
Gfi1	ENSMUSG0000029275	458	Rasgrp1	ENSMUSG0000027347	237	72	2.73E-08	9.08E-06
Ikzf1	ENSMUSG0000018654	93	Myc	ENSMUSG0000022346	359	11	5.30E-08	1.69E-05
Gfi1	ENSMUSG0000029275	458	Jundm2	ENSMUSG0000034271	105	23	8.30E-08	2.54E-05
Jundm2	ENSMUSG0000034271	105	Evi5	ENSMUSG0000011831	466	25	4.60E-07	1.26E-04
Gfi1	ENSMUSG0000029275	458	Notch1	ENSMUSG0000026923	127	33	9.12E-07	2.41E-04
Myc	ENSMUSG0000022346	359	Notch1	ENSMUSG0000026923	127	22	9.88E-07	2.52E-04
Map3k8	ENSMUSG0000024235	37	Myc	ENSMUSG0000022346	359	1	1.68E-06	4.02E-04
Gfi1	ENSMUSG0000029275	458	Lck	ENSMUSG0000000409	26	1	2.60E-06	6.03E-04
Mycn	ENSMUSG0000037169	81	Myc	ENSMUSG0000022346	359	12	1.74E-05	3.70E-03
Evi5	ENSMUSG0000011831	466	Lck	ENSMUSG0000000409	26	2	2.57E-05	5.31E-03
Gfi1	ENSMUSG0000029275	458	Zfp438	ENSMUSG0000050945	51	10	7.48E-05	1.33E-02
A530013C23Rik	ENSMUSG0000006462	43	Evi5	ENSMUSG0000011831	466	8	1.25E-04	1.92E-02
Ccnd3	ENSMUSG0000034165	206	Ikzf1	ENSMUSG0000018654	93	6	1.34E-04	2.00E-02
Mycn	ENSMUSG0000037169	81	Notch1	ENSMUSG0000026923	127	1	1.99E-04	2.77E-02
Gfi1	ENSMUSG0000029275	458	Rras2	ENSMUSG0000055723	224	79	2.51E-04	3.12E-02
Ccr7	ENSMUSG0000037944	50	Evi5	ENSMUSG0000011831	466	11	2.42E-04	3.12E-02
Pvt1	ENSMUSG0000072566	296	Mycn	ENSMUSG0000037169	81	11	5.62E-04	4.87E-02

Table 3.6. Gene pairs in which insertions co-occur more often (A) or less often (B) than expected by chance. All tests with a *q*-value of less than 0.05 are shown.

upregulated by promoter and enhancer insertions, rather than being inactivated, as would be expected for co-operation in tumourigenesis. The distribution of *Lck*-associated insertions suggests that the full-length protein is produced (see Section 3.4.1), and can therefore be phosphorylated by Csk. Further experimental analysis of these genes is therefore required to understand their cooperative role.

A highly significant association was also identified between *Ikaros* (*Ikzf1*) and *Notch1*. An MuLV screen performed on transgenic mice expressing the oncogenic *Notch1* intracellular domain has previously identified the disruption of *Ikaros* as a co-operating event in lymphomagenesis (Beverly and Capobianco, 2003). Loss of heterozygosity of *Ikaros* and activation of *Notch1* have also been shown to co-occur in mouse thymic lymphomas induced by gamma-irradiation (Lopez-Nieva *et al.*, 2004; Ohi *et al.*, 2007). Activating insertions also co-occurred in *Notch1* and lunatic fringe (*Lfng*). *Lfng* encodes a glycosyltransferase that initiates elongation of *O*-linked fucose residues attached to the extracellular epidermal growth factor-like domain of Notch1 (Moloney *et al.*, 2000). This increases the sensitivity of Notch1 to Delta-like, rather than Jagged, Notch ligands and so promotes T cell, rather than B cell, development from haematopoietic progenitors (Besseyrias *et al.*, 2007; Haines and Irvine, 2003; Visan *et al.*, 2006). Upregulation of *Lfng* by insertional mutagenesis may therefore contribute to tumourigenesis by mediating an increase in the binding of oncogenic Notch1 to Delta-like ligands.

A significant co-occurrence was also identified between *Runx1* and *Rasgrp1*. The *Runx1* gene encodes the DNA binding alpha subunit of the Runt domain transcription factor PEBP2/CBF. *Runx1* translocations and point mutations are frequently implicated in human leukaemias and are often associated with activation of the Ras pathway (Goemans *et al.*, 2005). *Rasgrp1* is a Ras GTPase-specific guanine nucleotide exchange factor that activates Ras in lymphocytes (Roose *et al.*, 2007) and, in support of the observed co-occurrence, it was shown to be preferentially targeted by the endogenous retrovirus in BXH2-*Runx1*^{+/-} mice (Yamashita *et al.*, 2005).

There were also numerous genes for which the number of co-occurrences was lower than expected. The lack of co-operation between *Myc* and *Mycn* reflects the fact that they are structurally and functionally related. Co-occurring insertions disrupting *Myc* and either *Ikaros* or *Notch1* were also under-represented. *Myc* is a transcriptional target of the Notch signalling pathway in T cell acute lymphoblastic leukaemia, and Notch1 is

required to sustain the high levels of Myc that are required for continued growth and survival of the cancer (Sharma *et al.*, 2007). The mutual exclusivity of activated *Notch1* and *Myc* in mouse tumours suggests that *Notch1* activation may not provide a significant growth advantage when high levels of Myc are sustained by constitutive overexpression. *Mycn* and *Notch1* were also mutually exclusive, suggesting that Notch1 may play a similar role in the maintenance of *Mycn* expression during tumourigenesis. Co-occurring insertions that disrupt *Gfi1* and either *Rras2* or *Rasgrp1* were also under-represented. *Rasgrp1* and Ras-related *Rras2* were significantly associated, suggesting that Rasgrp1 activates *Rras2* and that overexpression of both genes contributes to tumourigenesis. The mutual exclusivity of disrupted *Gfi1* with both of these activated genes suggests that they may act in a common cancer pathway.

Many of the significant associations identified in this analysis are supported by observations in the literature, yet there are many more for which there is no evidence, in many cases because little is known about the genes involved. The list of co-occurring and mutually exclusive genes therefore provides a basis for future functional analyses, and demonstrates the potential of large scale insertional mutagenesis screens in the identification of cancer gene collaborations in mouse, and human, tumourigenesis.

3.6 Discussion

The purpose of the work described in this chapter was to characterise the candidate cancer genes identified by insertional mutagenesis, and to demonstrate their relevance to human tumourigenesis. The candidates showed a significant overlap with human mutation datasets associated with, or biased towards, cancers of haematopoietic and lymphoid tissue, but not with breast and colon candidate cancer genes. This suggests that the screen may only be effective in identifying novel candidates involved in the development of lymphomas and/or leukaemias. A number of the over-represented GO terms were also associated with the development, differentiation or carcinogenesis of T- and B-cells, but others were associated with general features of cancers, such as cell proliferation and apoptosis. An exciting observation to be followed up was the positive association between candidate genes and genes containing Nanog and Oct4 binding sites. This suggests that a significant proportion of the candidate genes may be involved in tumour cell self-renewal, which has not been previously reported in insertional mutagenesis screens. This chapter also presents evidence for an overlap between genes identified by

insertional mutagenesis and regions of copy number change in human acute lymphoblastic leukaemias, therefore providing a justification for the cross-species comparative analyses performed in Chapters 4 and 5.

The comparison of candidate genes identified in the MuLV and Sleeping Beauty screens demonstrated differences in the mutational profiles of the two mutagens. This suggests that the use of different mutagens can increase the spectrum of candidate cancer genes, but the difference in profiles can only be fully appreciated by comparing fully saturated screens, since some CISs may be missing from one screen simply because of an insufficient number of PCRs or low sequencing depth. However, comparison of the screens does provide strong evidence that overlapping genes are involved in tumourigenesis, rather than resulting from insertional bias, which differs in MuLV and Sleeping Beauty (see Section 1.4.2.1.1). Osk was flagged as a promising candidate following its identification in both screens. In light of this finding, Fanni Gergely at the Cambridge Research Institute performed RNAi-mediated knockdown of OSK in HeLa cells, which are an immortal cell line derived from human cervical cancer cells, and scored chromosome lagging in 40 late anaphase/early telophase cells in 2 separate experiments. Chromosome lagging was observed in 12.3 ± 2.2 control cells and $28.1 \pm$ 4.0 cells with OSK knocked down by 95-100% (Figure 3.8). No other mitotic defects Chromosome lagging at anaphase can result in the failure of a were observed. chromosome or chromatid to become incorporated into one of the daughter nuclei following cell division. This causes an uploidy and can therefore contribute to genomic instability and cancer formation. This study is ongoing, but suggests that QSK does play important role in tumourigenesis. Likewise, *p116Rip*, Zmiz1 and an ENSMUSG0000075015 contained both MuLV and T2/Onc insertions and are therefore promising candidates for which functional validation is required.

Co-occurring MuLV and T2/Onc insertions were also used in the prediction of the mechanisms of mutation of candidate cancer genes. While these may not always recapitulate the mutations observed in human cancer, as demonstrated for *Flt3*, in other cases, e.g. *Notch1*, similar mutations are observed. Identifying the structure and function of the mutant products of oncogenes is valuable in the development of therapeutic drugs that target those proteins. Experimental approaches are required to validate the predictions, although the efficacy of gene expression analysis appears to be variable, since in the limited analysis performed in Section 3.4.5, many CIS genes did not show

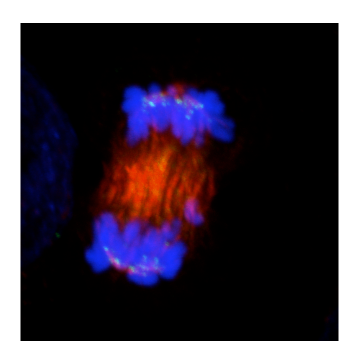


Figure 3.8. Knockdown of *QSK* **in human HeLa cells is associated with increased chromosome lagging at anaphase.** Figure shows a single cell at anaphase, with chromosomes stained blue and spindle fibres stained red. Image provided by Fanni Gergely at the Cambridge Research Institute.

significant differential expression in tumours containing insertions versus those without. Promoter and enhancer mutations appeared to be the most common mechanisms of mutation, with upregulation of the wildtype gene being the most common type of mutation overall. Initial comparison against a predicted set of regulatory features suggests that disruption of regulatory elements is not a common mechanism of mutation in insertional mutagenesis, although a more accurate analysis could be performed using a set of regulatory features specific for the mouse, rather than human. Analysis of the distribution of insertions within genes can also facilitate the identification of tumour suppressor genes, which are expected to contain only intragenic, truncating mutations, and may show a more random distribution of insertions that includes multiple insertions from the same tumour. For a number of the genes studied in this chapter (i.e. Etv6, Myb, Fli1, Erg, Foxp1), some of the insertions appeared to be associated with Ensembl EST genes for which there was no associated Ensembl gene transcript. Therefore, analysis of the distribution of insertions from insertional mutagenesis screens may also facilitate the identification of novel gene transcripts.

The identification of collaborating cancer genes is important for the development of targeted cancer therapies. As discussed in Section 1.2.7, cancers can develop resistance to targeted therapies but this may be alleviated by developing therapies that target multiple genes simultaneously. Collaborating cancer genes can also help in deciphering the complex landscape of cancer genomes and the events involved in the multi-step process of tumour evolution. The analyses described in Section 3.5 have identified a number of collaborations for which there is supporting evidence in the literature, as well as many novel collaborations.

In summary, this chapter demonstrates that insertional mutagenesis is a powerful tool for identifying both novel candidate cancer genes and collaborations between candidate cancer genes that are relevant to mouse and human tumourigenesis. In order to maximise the candidates and collaborations identified by this approach, the combined use of a variety of insertional mutagens and genetic backgrounds is recommended. In the future, the development of mutagens that can induce the formation of solid tumours should facilitate the identification of a larger repertoire of cancer gene candidates.

Chapter 4 Using mouse candidate cancer genes to narrow down the candidates in regions of copy number change in human cancers

4.1 Introduction

As discussed in Section 1.3.3, copy number changes are a common feature of cancer genomes, and can be identified using comparative genomic hybridisation (CGH)-based techniques. However, regions of copy number change are often large and encompass many genes, making it difficult to identify the "critical" genes that contribute to the tumourigenic process. Candidate cancer genes identified by insertional mutagenesis in the mouse can be used in a cross-species oncogenomics approach to narrow down the candidates within regions of copy number change in human tumours. The use of cross-species comparative analysis for cancer gene discovery is discussed in Section 1.5. In this chapter, mouse candidate cancer genes are used to identify orthologous candidates within regions of copy number change in 713 human cancer cell lines generated using SNP array CGH. The analyses were performed as part of a collaboration with the Netherlands Cancer Institute (NKI), published in Cell (Uren *et al.*, 2008), and therefore, rather than using the mouse candidates were provided by the NKI.

The datasets are introduced in Section 4.2. This is followed, in Section 4.3, by a description of the methods used to process the copy number data into regions of copy number change, and gains and losses within the human cancer cell lines are characterised in Section 4.4. In Section 4.5.1, the mouse and human datasets are compared to determine whether retroviral insertional mutagenesis is relevant to the discovery of amplified and deleted cancer genes in humans. Promising cancer gene candidates that are both disrupted by insertional mutagenesis in the mouse and amplified or deleted in human cancers are presented in Section 4.5.2. A range of algorithms have been developed for identifying regions of copy number change within CGH data, and these are described and compared in Section 4.6. Finally, in Section 4.7, the mouse candidate cancer genes are combined with copy number variation (CNV) data from apparently healthy individuals to determine whether there is any overlap between candidates and regions of CNV.

Since the ploidy of the cell lines, and therefore the exact copy number of alterations, is difficult to establish, the terms "gain and "amplicon" are used interchangeably throughout this thesis to mean any gain of copy number, irrespective of the size or nature of the alteration.

4.2 Description of the datasets

As well as the datasets described below, the set of known cancer genes from the Cancer Gene Census (Futreal *et al.*, 2004) was also used. This is described in Section 2.2.3.

4.2.1 Mouse candidate cancer genes identified by retroviral insertional mutagenesis

As mentioned in the introduction, some of the work described in this chapter was undertaken as part of a collaboration with the NKI (Uren et al., 2008, reprinted on p.365). The gene lists used in this chapter were therefore provided by the NKI but were generated from the analysis of insertion sites identified in the retroviral insertional mutagenesis screen described in Chapter 2. There were 6 lists of putative tumour suppressor genes. These included 3 lists comprising all genes in which there were insertions in the entire transcribed region, including UTRs and introns, only in the translated region (no UTRs) but including introns, and only in the coding region (no UTRs or introns). These lists are described throughout this thesis as genes in the transcribed region, translated region, and coding region, respectively. A further 3 lists contained genes with insertions in the same regions, but only where insertions comprised 2 or more sequence reads. Insertions represented by only 1 read are considered less likely to contribute to tumourigenesis (see Section 2.8) and are therefore predicted to have a reduced overlap with human deletions. 2 additional lists contained genes that were closest to CISs with *P*-values of less than 0.05 and 0.001, as determined using the kernel convolution (KC)-based statistical method (de Ridder et al., 2006, see Sections 1.4.2.1.2 and 2.10.2). From these, lists were also generated for genes that were adjacent to CISs of P<0.05 and P<0.001 but were further away than the closest gene. For each gene list, the human orthologues and their genomic coordinates were extracted from Ensembl version 37 using Ensembl BioMart (see Section 3.2.1). Table 4.1 shows the number of mouse genes and human orthologues in each gene list. The P<0.001 and P<0.05 CISs and their associated nearest and further mouse genes

Gene List	Number of mouse genes		Number of human orthologues in CIS gene list	% of human orthologues in CIS gene list
ORF only	266	240	41	17.1
ORF only (no singletons)	86	75	22	29.3
Translated region only	3024	2647	216	8.2
Translated region only (no singletons)	1331	1163	173	14.9
Transcribed region only	3773	3316	275	8.3
Transcribed region only (no singletons)	1706	1498	227	15.2
CIS nearest P<0.05	559	424	196	46.2
CIS nearest P<0.001	355	265	155	58.5
CIS further P<0.05	505	362	85	23.5
CIS further P<0.001	313	219	66	30.1

Table 4.1. Description of the lists of mouse candidate cancer genes used for comparison with human cancer copy number data. "[ORF, Translated region, Transcribed region] only" are lists of genes containing insertions only in the open reading frame, translated region (but including introns) or transcribed region, respectively. "no singletons" means that the list does not include genes that only contain insertions represented by a single read. "CIS nearest P<0.05" and "CIS nearest P<0.001" contain genes nearest to CISs identified by the kernel convolution (KC)-based method. "CIS further P<0.05" and "CIS further P<0.001" contain genes that flank CISs identified by the KC-based method but are not the nearest genes. The columns labelled "Number/% of human orthologues in CIS gene lists" show the overlap of each list with the list of candidate cancer genes generated and described in Chapters 2 and 3.

are listed in Appendix D. Due to their length, the lists of candidate tumour suppressor genes are not included, but are available on request.

Table 4.1 also shows the overlap of each gene list with the list of candidate cancer genes generated and described in Chapters 2 and 3 (shown in Appendix B2 and referred to here as the CIS gene list). The CIS gene list contains only genes that are associated with a significant CIS and this, together with the fact that the screen identifies mainly oncogenes, accounts for the small overlap with the tumour suppressor gene lists, in which genes may contain any number of insertions. The differences between the CIS gene list and the remaining lists may reflect differences in gene selection, i.e. a more sophisticated method was used to assign insertions to genes in the CIS gene list, and in read and insertion site processing, which were more conservative for the CIS gene list. Candidates from the CIS gene list are used in Chapter 5, where it is compared to higher resolution human CGH data (Section 5.3), as well as to the CGH data described in this chapter (Section 5.4).

4.2.2 Copy number data for human cancer cell lines

Comparative genomic hybridisation (CGH) data were generated by the Wellcome Trust Sanger Institute (WTSI) Cancer Genome Project for 713 human cancer cell lines from 29 tissues. A list of all cell lines and their tissue of origin is provided in Appendix E and is summarised in Table 4.2. None of the chosen cell lines had a common ancestor, according to cell line identity typing also performed by the WTSI Cancer Genome Project (http://www.sanger.ac.uk/genetics/CGP/Genotyping/synlinestable.shtml). This is important, since an amplicon or deletion might otherwise appear to be recurrent simply because it is within synonymous cell lines. CGH was performed using two Affymetrix GeneChip® Human Mapping Arrays. The 10K array, which comprises 11,555 SNPs, was used for 313 cell lines, while the 10K 2.0 array, comprising 10,204 SNPs, was used for the remaining 400 lines. 10,136 SNPs were shared between the two arrays, and both used the Affymetrix GeneChip® Mapping 10K assay, described in Section 1.3.3.2. The SNPs were mapped to the NCBI 35 human genome assembly. The mean distance between SNPs was 258.50 (±634.21) kb in the 10K array, and 292.82 (±683.49) kb in the 10K 2.0 array. The minimum distance was 2 bp and 11 bp for the 10K and 10K 2.0 arrays, respectively, and the maximum distance was 24.81 Mb for both arrays. 9.4% of human protein-coding genes in Ensembl v37 (extracted using Ensembl BioMart, see

Tissue of origin	Number of cell lines
Lung	131
Haematopoietic and lymphoid	117
Breast	43
Skin	42
Central nervous system	40
Unknown	39
Large intestine	38
Autonomic ganglia	29
Bone	23
Kidney	21
Soft tissue	20
Oesophagus	20
Stomach	19
Upper aerodigestive tract	19
Ovary	18
Pancreas	14
Urinary tract	13
Liver	11
Thyroid	11
Cervix	11
Endometrium	10
Biliary Tract	6
Pleura	5
Testis	3 2 2
Vulva	2
Prostate	2
Eye	2
Placenta	2
Adrenal gland	1
Small intestine	1
<u>Total</u>	713

Table 4.2. Tissues of origin of human cancer cell lines used in the 10K SNP arrayCGH analysis.

Section 3.2.1) contained at least one SNP in the 10K array, while 9.0% contained at least one SNP in the 10K 2.0 array. Genes were defined as the longest Ensembl gene transcript. The 10K and 10K 2.0 arrays contained an average of 0.176 (\pm 0.735) and 0.157 (\pm 0.648) SNPs per protein-coding gene, respectively. The interSNP distances and number of SNPs per gene are shown in Figures 4.1 and 4.2, respectively. The largest gaps between adjacent SNPs occur at the centromeres, while some gaps correspond to other regions of the genome that have not been assembled, e.g. due to highly repetitive sequences.

For each cell line, the raw intensity values were normalised internally. This involved calculating the value for each SNP as a total of all the SNPs on the array, and obtaining a copy number ratio for each SNP by dividing the SNP value by the value for the same SNP from a pool of reference normal samples. This is the point at which I received the data. The copy number data for all cell lines are available for download from ftp://ftp.sanger.ac.uk/pub/CGP/10kData. Data generated on the 10K and 10K 2.0 arrays is pooled in subsequent analyses and is collectively referred to as 10K data.

4.2.3 Copy number variants (CNVs)

CNVs are regions within the genome that vary in copy number. Germline CNV regions identified within 270 HapMap samples from Redon *et al.* (2006) were downloaded from http://www.sanger.ac.uk/humgen/cnv/data/cnv_data/. Merged CNVs identified using the Whole Genome Tilepath (WGTP) array and Affymetrix GeneChip Human Mapping 500K early access array (500K EA) were used. The WGTP array comprises 26,574 BAC clones, while the 500K EA array covers 474,642 SNPs. The WGTP and 500K EA platforms are complementary, since they are able to detect smaller and larger CNVs, respectively (Kehrer-Sawatzki, 2007). There are 1,447 merged CNVs that cover ~12% of the genome. 1,390 CNVs that mapped to autosomes in the NCBI 35 human build were used in this analysis.

4.3 Processing the copy number data

The copy number ratios at individual SNPs must be processed into regions of copy number change. As discussed in Section 1.3.3.2, a variety of methods have been

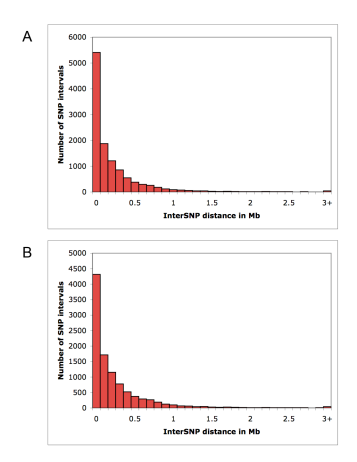


Figure 4.1. The distance between the genomic coordinates of adjacent SNPs on the 10K (A) and 10K 2.0 (B) SNP arrays.

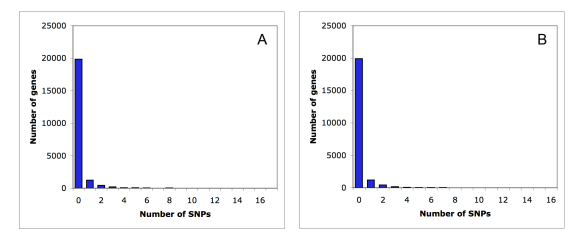


Figure 4.2. The number of SNPs per human protein-coding gene on the 10K (A) and 10K 2.0 (B) SNP arrays.

developed for this purpose. At the time of the analysis, most of the available algorithms had been developed primarily for conventional array CGH, i.e. using large genomic clones (see Section 1.3.3.2). In a comparison of 11 methods, DNAcopy (Olshen *et al.*, 2004) performed consistently well (Lai *et al.*, 2005), and a comparison of 3 segmentation methods by Willenbrock and Fridlyand (2005) demonstrated that DNAcopy performed better than GLAD (Hupe *et al.*, 2004) and HMM (Fridlyand *et al.*, 2004). A further benefit of DNAcopy is that it is freely available as an R package in BioConductor (http://www.bioconductor.org/). BioConductor is an open source software project that provides tools, mostly written in R, for analysing genomic data. DNAcopy (version 1.4.0) was therefore chosen as the method for detecting regions of copy number change in the 10K CGH data.

DNAcopy uses a method called circular binary segmentation (CBS) to identify changepoints in CGH data, which is input as log₂ intensity ratios at consecutive positions in the genome. The change-points correspond to positions in the genome where the DNA copy number has significantly changed. For each cell line, the copy number ratios for all SNPs were converted to log₂-ratios and were smoothed, using a method within DNAcopy, to remove single point outliers before segmentation. Copy number ratios of 0 were given a log₂-ratio of -6. Change-points may result from local trends in the data, and therefore all change-points that were less than 3 standard deviations apart were removed. Default parameters were used for the segmentation. Different values were tested for the parameter alpha but, upon visual inspection of the graphical outputs, the default value of alpha=0.01 appeared to be most suitable. Increasing alpha increases the sensitivity, resulting in more change-points but, potentially, more false positive change-points. Decreasing alpha results in fewer change-points, and regions of copy number change may therefore be missed. Increasing the number of standard deviations below which changepoints were removed resulted in the loss of potentially important change-points. Figure 4.3 shows an example of how changing the parameters can affect the output of DNAcopy for chromosomes 1 and 6 of ovarian cancer cell line 41M-CISR. The removal of changepoints less than 3 standard deviations apart results in the loss of a change-point in chromosome 1 (Figure 4.3B). However, the slight difference in copy number between the 2 arms of the chromosome may be due to trends in the data, and the difference in copy number is small. Increasing the number of standard deviations to 4 results in the loss of a change-point in chromosome 6, for which there is a clear step in copy number that does look real (Figure 4.3C). Increasing alpha from 0.01 to 0.05 results in the inclusion of

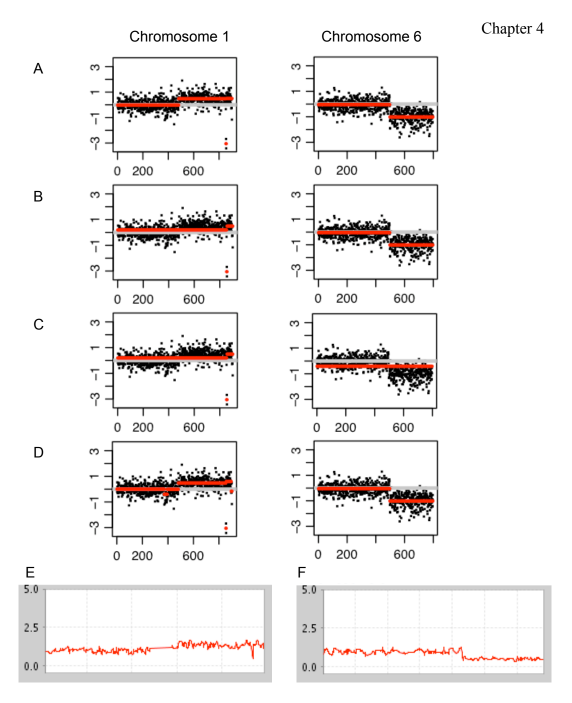


Figure 4.3. Altering the values for parameters in DNAcopy leads to differences in the regions of copy number change detected by the algorithm, as demonstrated for chromosomes 1 and 6 of ovarian cancer cell line 41M-CISR. (A) Default parameters. (B) Default parameters and removal of change-points less than 3 standard deviations apart. (C) Default parameters, smoothing and removal of change-points less than 4 standard deviations apart. (D) Alpha = 0.05 plus smoothing. (E) Copy number for chromosome 1, with values averaged across 3 consecutive SNPs. (F) Copy number for chromosome 6, with values averaged across 3 consecutive SNPs. Figures E and F taken from the WTSI Cancer Genome Project website are (http://www.sanger.ac.uk/genetics/CGP/) and give a clearer picture of the copy number across the chromosome. Figures A-D are extracted from the output of DNAcopy. Removing change-points that are close together results in fewer regions being detected, and the larger the number of standard deviations below which change-points are removed, the more regions are missed. Increasing alpha leads to the inclusion of additional changepoints and, therefore, regions of copy number change.

additional change-points in chromosome 1 (Figure 4.3D). Since the data is relatively low resolution, it is highly possible that a region of copy number change may be represented by just 1 or 2 SNPs. However, it is also possible that such SNPs are anomalies and, to avoid the identification of false positives, this is the preferred assumption.

DNAcopy identifies changes in DNA copy number but does not indicate which regions are unchanged and which are gains or losses. It is therefore the responsibility of the user to set thresholds for calling gains and losses based on the mean log₂-ratios of predicted segments. A disadvantage of DNAcopy is that it operates on individual chromosomes rather than the entire genome and the mean log₂-ratios of segments representing no copy number change, or representing a gain or loss of a certain number of copies, will differ slightly across the genome. This makes it difficult to determine what is "normal" and therefore to call gains and losses, and the exact number of copies within a gain or loss cannot be clearly determined. Willenbrock and Fridlyand (2005) have developed an algorithm called MergeLevels that merges segments across the genome that are not significantly different from one another and so produces a more interpretable set of copy number levels. Combining DNAcopy and MergeLevels was found to be more effective than using DNAcopy alone (Willenbrock and Fridlyand, 2005). MergeLevels is freely available within an R/BioConductor package called aCGH. Therefore, for each cell line, the DNAcopy segmentation results were merged across all autosomes using MergeLevels with default parameters, which were considered appropriate upon inspection of the graphical outputs. Example outputs for the kidney cancer cell line 786-0 and endometrial cancer cell line AN3-CA are shown in Figure 4.4. The merged segments with a log₂-ratio closest to 0 were defined as the level of no copy number change and, to enable comparison across cell lines, this log₂-ratio was set to 0 and all other log₂-ratios were normalised accordingly. Figure 4.5A shows the distribution of log₂-ratios of the segments predicted by DNAcopy across all cell lines, while Figure 4.5B shows the distribution of log₂-ratios of the merged segments. The log₂-ratios of the merged segments show a series of peaks and troughs that may reflect distinct copy number levels. The large peak at -6 represents segments for which the copy number ratio of individual SNPs was 0. The log₂ratios of the merged segments were converted to copy number ratios (Figure 4.6), and troughs in the distribution were used to set thresholds for subsequent analyses (see Sections 4.4 and 4.5.1.1).

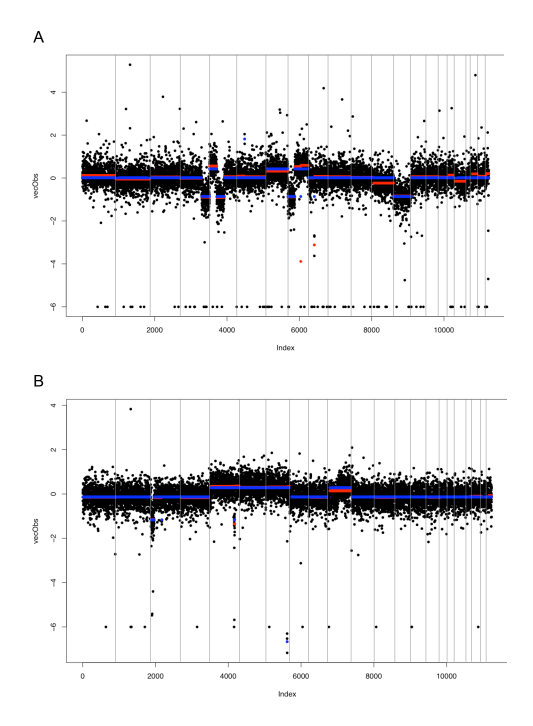


Figure 4.4. Graphical output from MergeLevels for human cancer cell lines 786-0 (A) and AN3-CA (B). Black dots represent the log_2 -ratios for individual SNPs ordered across the genome. The mean log_2 -ratios for segments identified by DNAcopy are shown in red. Merged segments generated by MergeLevels are shown in blue. Segments are merged across chromosomes into a set of copy number levels. The x-axis shows the position within the genome, while the y-axis measures the log_2 -ratio. Vertical lines represent the division of chromosomes.

Chapter 4

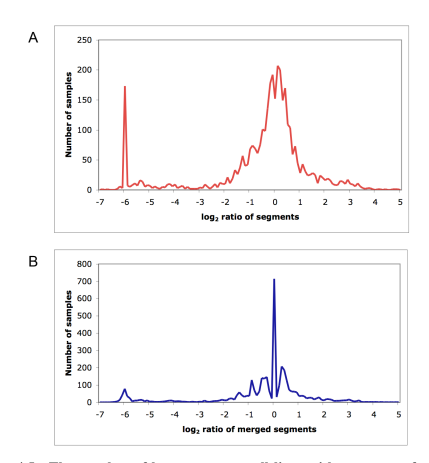


Figure 4.5. The number of human cancer cell lines with segments of varying log₂ratio following processing with DNAcopy (A) and DNAcopy plus MergeLevels (B).

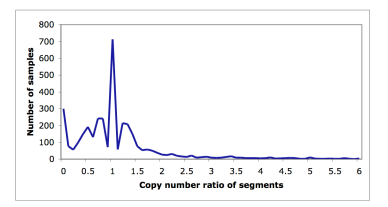


Figure 4.6. The number of human cancer cell lines with segments of varying copy number ratio following processing with DNAcopy plus MergeLevels. Troughs in the data were used to set thresholds for the analysis of gains and losses.

4.4 Characterising gains and losses in cancer genomes

All segments with a copy number of 1.6 or more were designated gains, and all segments with a copy number of 0.6 or less were designated losses. Segments with a copy number of 0.2 or less were designated homozygous deletions. Each of these thresholds was within a trough in the distribution of copy numbers for merged segments in Figure 4.6. The average number of gains per cancer cell line was $1.47 (\pm 2.02)$, and the average size across all cell lines was 21.37 (±34.15) Mb. Amplicons contained an average of 213.68 (± 341.45) genes. The average number of losses per cell line was 4.43 (± 3.69) and the average size was 19.56 (±33.83) Mb, encompassing 195.62 (±338.32) genes. The average number of homozygous deletions was 1.53 (±1.99) per cell line. The average size was 0.98 (±2.68) Mb, encompassing 19.64 (±20.69) genes. Therefore, homozygous deletions were significantly smaller than amplicons and heterozygous deletions and contained fewer genes. Homozygous deletions have been previously shown to contain fewer genes than other regions of the genome (Cox et al., 2005, see Section 1.3.3.3), and this analysis shows that, in general, the deletion of both copies of a gene is more likely to be deleterious to a cell than the loss of one copy or the gain of copies. The distributions of amplicon and deletion lengths are shown in Figure 4.7.

For each cancer type, the number of cell lines containing gains was counted, and the 2tailed Fisher Exact Test was used to determine whether there was any significant difference between the observed and expected number of each cancer type. Cell lines derived from the oesophagus were over-represented ($P=7.11\times10^{-4}$), suggesting that oesophageal tumours are particularly prone to genomic instability. Haematopoietic and lymphoid cancer cell lines were under-represented ($P=5.32\times10^{-5}$), reflecting the fact that they often contain balanced translocations that do not show a change in DNA copy number (see Section 1.3.3.4) and that, in some cases, few genetic events are thought to be required for tumour development. For example, acute lymphoblastic leukaemias contain an average of 3.83 deletions and focal amplifications are rare (Mullighan *et al.*, 2007). Since most cell lines contained deletions, there was no significant difference between the observed and expected numbers of cancer types containing deletions.

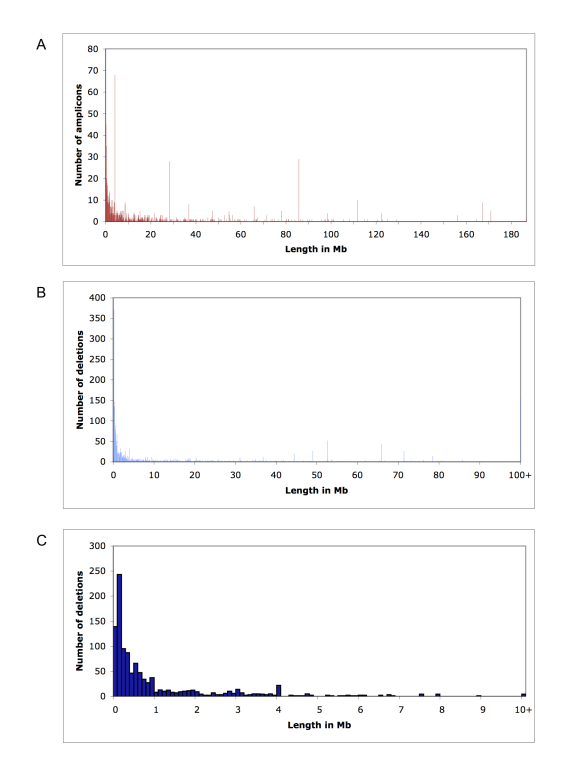


Figure 4.7. Distribution of the lengths of amplicons (A), deletions (B) and homozygous deletions (C) in 713 human cancer cell lines. Amplicons are defined as regions with a copy number greater than or equal to 1.6. Deletions and homozygous deletions are defined as regions with a copy number less than or equal to 0.6 and 0.2, respectively.

4.5 Comparative analysis of mouse candidate cancer genes and CGH data from human cancers

4.5.1 Global comparison

The purpose of the global comparison is to determine whether the human orthologues of candidate cancer genes identified by retroviral insertional mutagenesis by the Netherlands Cancer Institute are over-represented within regions of copy number change in the human cancer cell lines. Specifically, an over-representation of candidate oncogenes in human amplicons, and candidate tumour suppressor genes in human deletions, suggests that the retroviral insertional mutagenesis screen is relevant to human cancer, and may help to identify human cancer gene candidates within regions of copy number change.

4.5.1.1 Method

Rather than setting single copy number thresholds for gains and losses, a range of copy number thresholds were investigated. Thresholds were set as the centre-point of troughs in the graph shown in Figure 4.6, since these may represent transitions in the number of gene copies. The chosen thresholds were copy number ratios of less than or equal to 0.9, 0.6 and 0.2, and greater than or equal to 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9. The genomic coordinates of the human orthologues of all mouse genes were extracted from Ensembl version 37 using Ensembl BioMart. For each gene list described in Section 4.2.1, the number of mouse genes with human orthologues was counted. The same number of genes was selected randomly from among all mouse genes with human orthologues and this was repeated 1,000 times. For each of the 1,000 iterations, the number of human orthologues that resided within human cancer cell line segments with a mean copy number above or below a given threshold was counted. This produced a normal distribution of counts. The number of human orthologues of mouse candidate cancer genes in segments above or below each threshold was also counted. The Z-test was used to calculate the probability of obtaining a number greater than or equal to the observed count for the mouse candidates, based on the distribution of counts for the randomised genes. The procedure is summarised in Figure 4.8.

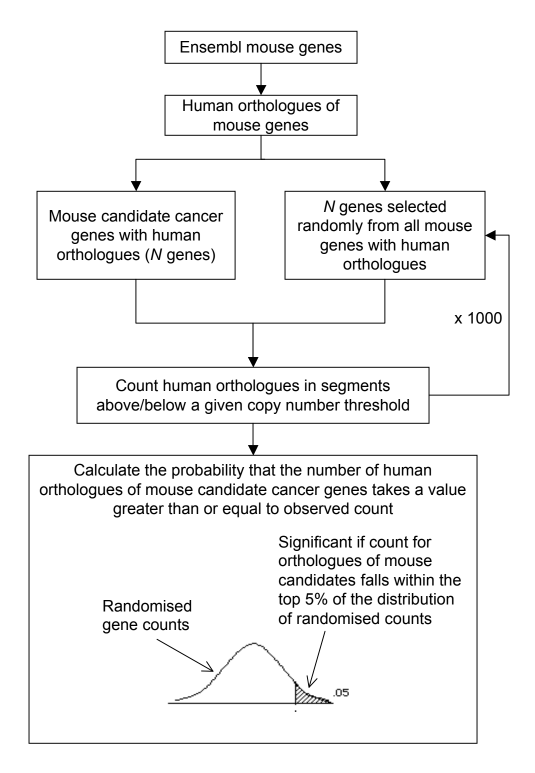


Figure 4.8. Overview of the method for identifying over-representation of the human orthologues of mouse candidate cancer genes in regions of human copy number change.

4.5.1.2 Setting the boundaries of amplicons and deletions

The start and end coordinates of the copy number segments generated by DNAcopy and MergeLevels correspond to the first and last SNPs for which the log₂-ratios are not significantly different to other SNPs in the segment. Therefore, the copy number can be determined for all coordinates between these positions. It is, however, impossible to determine the copy number for coordinates within the interval between the first SNP and the preceding SNP, which corresponds to the end coordinate of the preceding segment, and between the last SNP and the proceeding SNP, which corresponds to the start coordinate of the proceeding segment. As shown in Section 4.2.2, the distance between SNPs can be very large, especially across unassembled regions of the genome such as centromeres. Setting the boundaries of an amplicon or deletion as the end of the previous segment and start of the next segment, or even using half-way points, could therefore result in a very high number of false positives among genes predicted to be amplified or deleted.

In order to choose an appropriate distance for the boundaries of amplicons and deletions, the global comparison was performed using a range of distances. Assuming that CIS genes are more likely to be amplified or deleted in human cancers than are other genes, the most appropriate distance should be that which gives the highest over-representation of CIS genes. The list of genes nearest to CISs with P<0.001 was used in this analysis. Amplicon boundaries were extended beyond the first and last amplified SNP by a distance of 0 kb, 200 kb, 500 kb, 1 Mb, 3 Mb and 5 Mb, or as far as the adjacent SNP, whichever was closer. The results are shown in Figure 4.9. The association between CIS genes and amplicons was strongest when the boundaries were not extended at all. However, at lower copy numbers and in greater numbers of cell lines at higher copy number, the association was less significant than when the boundaries were extended to 500 kb. Extending the boundaries to 1 Mb and beyond resulted in a considerable decrease in the association between CIS genes and amplicons. Therefore, 500 kb was chosen as the most suitable distance.

Known oncogenes from among the CIS genes that were identified within full-length amplicons (i.e. where the amplicon was extended as far as the adjacent, non-amplified SNPs) were compared to those identified within amplicons with a 0 kb or 500 kb extension of the amplicon boundaries (Table 4.3). While the non-extended amplicons

Chapter 4

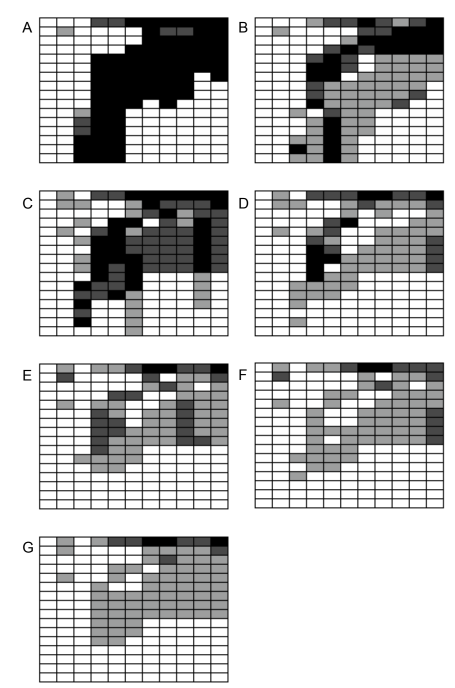


Figure 4.9. Over-representation of human orthologues of genes nearest to CISs in amplicons with boundaries extended beyond the first and last amplified SNP by a maximum distance of 0 kb (A), 200 kb (B), 500 kb (C), 1 Mb (D), 3 Mb (E), 5 Mb (F) and up to the adjacent, non-amplified SNPs (G). Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Columns represent amplicons, with (from left to right) copy number thresholds of greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4, 4.8, 5.2 and 5.9. Rows represent the number of cell lines, which increases in increments of 1, up to a cut-off of 16 cell lines. For example, the box in the bottom right-hand corner of each figure represents the P-value for the over-representation of CIS genes that occur in amplicons of copy number greater than or equal to 5.9 in at least 16 cancer cell lines.

Copy number	Gene	Full length	0 kb	500 kb
4.8	МҮС	14	10	14
	MYCN	9	4	9
	CCND1	1	1	1
4.3	МҮС	14	10	14
	MYCN	9	4	9
	CCND1	1	1	1
	ZFPN1A1	1	1	1
	LMO2	1	1	1
	CCND3	2	0	2
2.7	МҮС	29	22	29
	MYCN	12	7	12
	CCND1	4	4	4
	ZFPN1A1	1	1	1
	LMO2	2	2	1
	CCND3	3	0	2
	CCND2	1	1	1
	PIM1	2	1	2
	EVI1	1	1	1
	IRF4	1	1	1

Table 4.3. The number of amplicons in which known cancer genes among genes nearest to CISs are identified when the amplicon boundaries are altered. "Full length" applies to amplicons extended to the next, non-amplified SNP. "0 kb" applies to amplicons where the start and end correspond to the first and last amplified SNP. "500 kb" applies to amplicons extended to a maximum of 500 kb. Copy number values are given as the minimum copy number of amplicons.

missed some of the occurrences of amplified *MYC* and *MYCN* that were identified in the full-length amplicons, all occurrences were identified in the amplicons extended by up to 500 kb. Likewise, occurrences of amplified *CCND3* and *PIM1* were identified in the 500 kb amplicons but not the non-extended amplicons.

To demonstrate that the observed association between candidate cancer genes and human amplicons was real, full-length amplicons were shuffled across the genome. The length and mean copy number of each amplicon were conserved, but the location was shuffled. The method from Section 4.5.1.1 was then performed on the shuffled amplicons. As shown in Figure 4.10, the association of candidates with regions of copy number gain was completely abolished.

4.5.1.3 Comparison with lists of candidate cancer genes

Having chosen 500 kb as the maximum distance for extending amplicon and deletion boundaries, the method of Section 4.5.1.1 was applied to all of the gene lists outlined in Section 4.2.1. The results are shown in Figure 4.11. The lists of genes nearest to CISs with P < 0.001 or P < 0.05 are lists of candidate oncogenes, with those nearest to CISs with P < 0.001 being stronger candidates for a role in tumourigenesis. This is reflected in the results, since both lists showed an over-representation of candidates within regions of amplification, but the association was stronger for genes near to a CIS with P < 0.001. For both gene lists, the association became significant at copy number 1.6 and above, but for low-level copy numbers, the association was generally strongest for genes that were amplified in higher numbers of cell lines. Figure 4.12A shows the over-representation of known oncogenes within regions of copy number gain. The pattern of association was very similar to that obtained using genes nearest to CISs with P < 0.001, suggesting that this list contains oncogenes that are relevant to human cancer. Almost all of the mouse tumours generated in the retroviral screen were lymphomas, and therefore it could be assumed that the candidate cancer genes identified in the screen are only relevant to similar cancers within humans. Therefore, the human cancer cell lines were divided into haematopoietic and lymphoid cell lines and all other cell lines (from solid tumours) and the global comparison was performed on each subset using genes nearest to CISs with P < 0.001. As shown in Figure 4.13, the association was much weaker when only haematopoietic and lymphoid cell lines were considered. This may be partly because amplification is not a common mechanism of mutation in these cell types (see Section

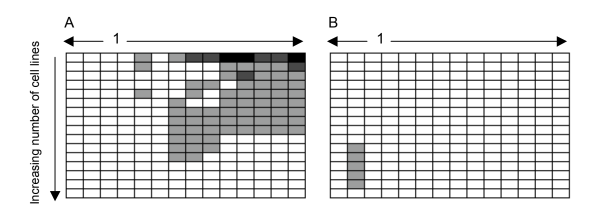


Figure 4.10. Over-representation of human orthologues of genes nearest to CISs in full-length human amplicons (A) and shuffled full-length amplicons (B). Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Copy number thresholds below 1 represent deletions, with (from left to right) copy number thresholds of less than 0.2, 0.6 and 0.9. Copy number thresholds above 1 represent amplicons, with (from left to right) copy number thresholds of greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9. The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

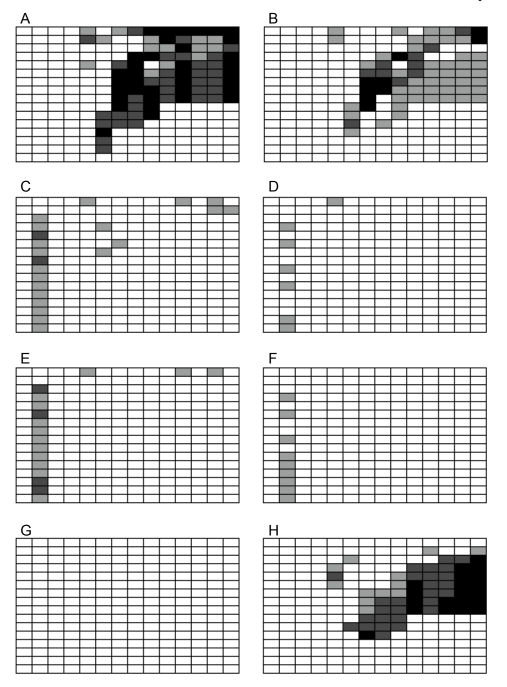


Figure 4.11. Over-representation of human orthologues of candidate cancer genes in regions of copy number change. (A) Genes nearest to CISs with P<0.001. (B) Genes nearest to CISs with P<0.05. (C) Genes with insertions within the transcribed region. (D) Genes with insertions but no singletons in the transcribed region. (E) Genes with insertions within the translated region. (F) Genes with insertions but no singletons in the translated region. (H) Genes with insertions but no singletons in the coding region. (H) Genes with insertions but no singletons in the coding region. (H) Genes with insertions but no singletons in the coding region. (H) Genes with insertions but no singletons in the coding region. Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Columns from left to right represent copy number thresholds of less than 0.2, 0.6 and 0.9 (deletions) and greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9 (amplicons). The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

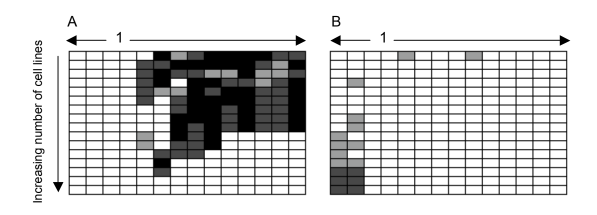


Figure 4.12. Over-representation of known oncogenes (A) and known tumour suppressor genes (B) in regions of copy number change in human cancer cell lines. Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Copy number thresholds below 1 represent deletions, with (from left to right) copy number thresholds of less than 0.2, 0.6 and 0.9. Copy number thresholds above 1 represent amplicons, with (from left to right) copy number thresholds of greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9. The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

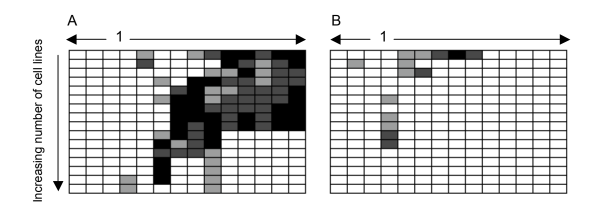


Figure 4.13. Over-representation of human orthologues of genes nearest to CISs with a *P*-value of <0.001 in regions of copy number change in human cancer cell lines derived from solid tumours (A) and haematopoietic and lymphoid cancers (B). See Figure 4.13 above for a description of how to interpret the figures.

4.4), but may also reflect the fact that the set of cell lines is smaller and therefore the comparison lacks power. Importantly, the pattern of association in solid tumours was similar to that for all cell lines and was highly significant. This demonstrates the relevance of retroviral insertional mutagenesis to the discovery of cancer genes in diverse human cancers, and shows that analysis of the full set of human cancer cell lines is warranted. Each cancer type provided in Table 4.2 was then separately tested for an association with the candidate cancer genes. Splitting the cancer cell lines into different types reduces the power of the analysis, and for most tumour types there was no clear association. However, cell lines derived from the autonomic ganglia, breast, upper aerodigestive tract, large intestine, oesophagus and stomach did show a significant overlap between mouse candidates and regions of copy number gain, although in the large intestine cell lines, there was also a significant overlap with regions of copy number loss (Figure 4.14).

The remaining lists are expected to contain candidate tumour suppressor genes. The results were similar for genes with insertions in transcribed and translated regions (Figure 4.11C-F). In both cases, including all genes containing insertions, rather than just those containing insertions represented by more than one read, generated a more significant association. This suggests that insertions represented by a single read ("singletons") in this retroviral screen are often important in tumourigenesis. As discussed in Section 2.7, the screen is not fully saturated due to the use of an insufficient number of enzymes in PCR and insufficient sequencing depth. Therefore, singleton insertions may result from these limitations, rather than because they are rare in the tumour mixture. However, for genes with insertions in the coding region, the reverse was observed, with a significant association only occurring when singleton insertions were omitted (Figure 4.11G-H, see below). As expected for tumour suppressor genes, the lists of genes with insertions in the transcribed and/or translated region were associated with deletions of copy number less than or equal to 0.6. However, the significance of the association was weak. When singleton insertions were included, there was also evidence of a weak association with regions of copy number gain. This is not surprising since the lists are likely to be contaminated with candidate oncogenes, as well as genes that do not play a role in tumourigenesis. The gene lists are long and yet tumour suppressor genes are less likely to be identified by insertional mutagenesis than are oncogenes and, as shown in Chapter 3, oncogenes are often disrupted by intragenic insertions. The association between known tumour suppressor genes and regions of copy number change is shown in Figure 4.12B.

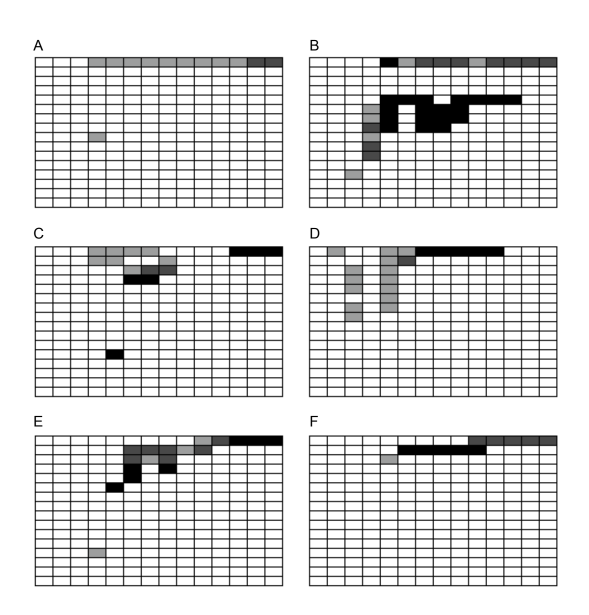


Figure 4.14. Over-representation of human orthologues of candidate cancer genes in regions of copy number change in cancer cell lines derived from the upper aerodigestive tract (A), autonomic ganglia (B), breast (C), large intestine (D), oesophagus (E) and stomach (F). Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Columns from left to right represent copy number thresholds of less than 0.2, 0.6 and 0.9 (deletions) and greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9 (amplicons). The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

There was a more significant over-representation of genes within deletions of copy number less than or equal to 0.6, but also in deletions of copy number less than or equal to 0.2. However, the fact that the pattern was broadly similar for genes with insertions in transcribed and translated regions is encouraging, and suggests that the lists do contain tumour suppressor genes that are relevant to human cancer.

The results for genes containing insertions within the coding region did not show the expected pattern for tumour suppressor genes (Figure 4.11G-H). As mentioned above, when singleton insertions were included, a significant association was not observed with either amplicons or deletions. Omission of singleton insertions resulted in a pattern of association representative of oncogenes, i.e. showing an over-representation of genes within amplicons. The identities of genes that reside within human amplicons and deletions are provided in Section 4.5.2.

4.5.1.4 Determining whether the nearest gene to a CIS is the most likely candidate cancer gene

As discussed in Chapter 2, it can be difficult to determine which gene is being mutated by insertions within a CIS, especially when the insertions are intergenic and disrupt genes by enhancer mutation. CISs are often assigned to the nearest gene. Therefore, to test whether this is a sensible assumption, the overlap of the human CGH data with candidate genes nearest to CISs was compared to that observed for the next nearest genes to CISs. The method was performed as described in Section 4.5.1.1, whereby the number of genes closest to CISs that occurred within amplicons or deletions was compared to the number of randomly occurring genes in amplicons or deletions. This was then repeated for genes adjacent to, but further from, CISs. Thresholds in this analysis were the same as for previous comparisons. The method was performed on CISs with a P-value of <0.05 and < 0.001, and the results are shown in Figure 4.15. As previously shown, there was a more significant over-representation within human amplicons of genes nearest to CISs with P < 0.001 than P < 0.05. However, for both significance levels, the clear overlap between human amplicons and genes nearest to CISs was almost absent for genes further from CISs. This suggests that the nearest gene to a CIS is generally the disrupted gene. *Plekhf1* and *Ltap* (also known as *Vangl2*) were the only two genes in the set of genes that are further from the CIS for which the human orthologues were amplified to a copy number greater than or equal to 5.2. However, in both cases, the nearest gene to the CIS

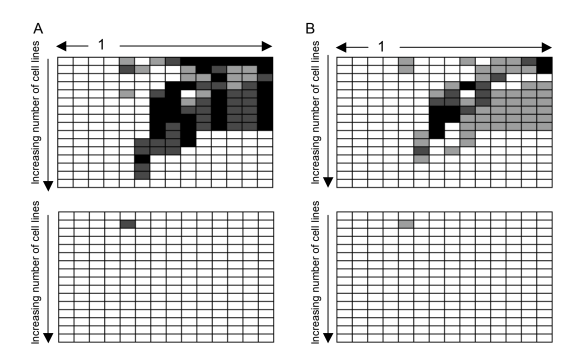


Figure 4.15. Over-representation of human orthologues of genes nearest to CISs (above) and genes further from CISs (below) in amplicons and deletions, where CISs have a *P*-value of <0.001 (A) and <0.05 (B). Each box represents the significance of the association between the selected genes and amplicons/deletions at a given copy number threshold and cell line number. *P*<0.0001, black; *P*<0.001, dark grey, *P*<0.05, light grey. Copy number thresholds below 1 represent deletions, with (from left to right) copy number thresholds of less than 0.2, 0.6 and 0.9. Copy number thresholds above 1 represent amplicons, with (from left to right) copy number thresholds of greater than 1.1, 1.6, 2.1, 2.5, 2.7, 3.1, 3.8, 4.3, 4.8, 5.2 and 5.9. The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

(1600014C10Rik and Slamf6, respectively) was also amplified, and analysis of the insertions around these genes suggests that the nearest gene is more likely to be disrupted by MuLV (Figure 4.16). *PLEKHF1* and *LTAP* may therefore be non-tumourigenic passengers within the amplified regions. *Tpcn2* and *Ccnd1* are neighbouring genes that both have nearby CISs, and both human orthologues were amplified, suggesting that both may be involved in tumourigenesis. For the remaining 9 genes nearest to CISs that were amplified to a copy number greater than or equal to 5.2, a human orthologue could not be found for the further gene. This explains why the lists of human orthologues of nearest genes are longer than the lists of human orthologues of further genes (see Table 4.1). Therefore, in some cases, the further gene may have an unidentified human orthologue that is also amplified in cancer. However, the fact that the nearest gene list contains a higher proportion of genes with human orthologues is itself significant, since cancer and cancer-related functions, such as cell growth, are well-studied and implicated genes may therefore be more likely to be conserved between species.

To investigate whether the difference between comparisons of nearest and further genes is most likely to be due to the further gene not being amplified or not having a human orthologue, the 66 human orthologues nearest to CISs that were amplified to a copy number greater than or equal to 2.7 were analysed in greater detail. 10 of the amplified genes, including *TPCN2* and *CCND1*, were neighbouring genes for which both mouse orthologues had nearby CISs. For 27 genes, the human orthologue of the further gene could not be identified. For a further 27 genes, the further gene was also amplified and, in all cases, both the nearest and the further genes were amplified in the same number of cell lines. There were only 2 amplified nearest genes, *Slc9a8* and *Mafk*, for which the further gene, *B4galt5* and *1110015K06Rik* respectively, had a human orthologue that was not amplified, suggesting that the nearest genes are the likely candidate cancer genes. In both cases, the human and mouse regions containing these genes are syntenic, and thus the lack of amplification is not due to a break in synteny in the human genome.

The reciprocal analysis was also performed, whereby the 36 human orthologues further from CISs that were amplified to copy number 2.7 or above were also investigated. 29 had neighbouring, nearer genes that were also amplified. This number is higher than the reciprocal count of 27 genes because 2 of the genes were adjacent to nearer genes that had more than one adjacent gene because they contained multiple CISs. For the 7 remaining

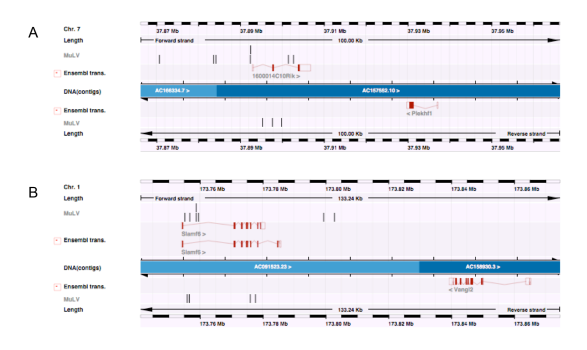


Figure 4.16. Insertions appear to be associated with the gene nearest to the CIS, i.e. *1600014C10Rik* (A) and *Slamf6* (B), even though adjacent genes are also amplified. Insertions are shown as black vertical lines. Those above the blue bar labelled DNA(contigs) are in the sense orientation, those below are in the antisense orientation. Ensembl genes are shown in red.

genes, there was no human orthologue for the nearest gene to the CIS. Therefore, it appears that the stronger overlap between the human amplicons and the human orthologues of genes nearest to CISs mainly reflects the inability to identify human orthologues for a higher percentage of the genes further from CISs. Since amplicons are generally large and encompass many genes (see Section 4.4), this is the more sensible explanation. However, several genes nearest to CISs were amplified in the absence of the further gene, while there were no examples of the reciprocal association. Choosing the nearest gene is the simplest method for assigning insertions to genes and will correctly identify genes that contain intragenic CISs. There is no evidence to suggest that genes nearest to CISs are preferentially amplified, but the fact that they are more likely to have a human orthologue does indicate that they may be the more likely candidates for a role in cancer. However, a more sophisticated method for assigning insertions to genes, such as the approach described in Sections 2.9 and 2.10, is likely to yield the most reliable list of cancer gene candidates. Comparative analyses involving these genes are discussed in Chapter 5.

4.5.2 Identification of individual candidates for a role in human cancer

For each gene list, the amplicons and deletions containing the human orthologues of candidate cancer genes were analysed in more detail to find the most promising candidates for a role in human cancer. The minimal amplified/deleted region was calculated by taking all of the amplicons/deletions in which the gene resided and finding the most 3' start coordinate and the most 5' end coordinate. The identities of other genes within the minimal region were established using the coordinates of human genes in Ensembl v37. For each gene within a minimum amplified or deleted region, the total number of cell lines in which that gene was amplified or deleted was calculated. In order to filter out the less likely candidates, genes in amplicons were discarded if they were co-amplified with known oncogenes or other mouse candidates from the gene list. Likewise, genes in deletions were discarded if they were co-deleted with known tumour suppressor genes or other mouse candidates from the gene list.

4.5.2.1 Candidate oncogenes among genes nearest to CISs

4.5.2.1.1 Protein-coding genes

The strongest candidates from the list of genes nearest to CISs with P<0.001 are shown in Table 4.4. Among 242 genes amplified to copy number 1.6 or above, 60 co-occurred with 1 or more known oncogenes and 128 co-occurred with other candidates in the list, of which 105 co-occurred with genes that were amplified in a greater number of cell lines. The filtered list of 54 candidates contained 14 known oncogenes, including *EVI1* and *FGFR2*, for which the murine insertions and human amplicons of less than 70 Mb are shown in Figure 4.17. 70 Mb is an arbitrary cut-off, but omits amplicons that are very large and for which there is therefore a low degree of certainty that the CIS genes are the targets of amplification. The kinase insert domain protein receptor gene (*KDR*) was amplified in 5 cell lines. Analysis of the insertions around *Kdr* in the mouse suggests that the adjacent gene, known oncogene *Kit*, may in fact be disrupted by the insertions assigned to *Kdr* (see Figure 2.11B, page 94). Likewise, the minimal amplified region containing *KDR* also contained *KIT*, which was amplified in an additional cell line (Figure 4.17) and is therefore the more likely target of amplification.

Further implicated oncogenes were also identified (Table 4.4). For example, the homeobox gene *MEIS1* is implicated in neuroblastoma. It was found to be amplified in the neuroblastoma cell line IMR-32 and was overexpressed in further neuroblastoma cell lines (Jones *et al.*, 2000). The single cell line in which it was amplified (to copy number 9.8) in this analysis was the neuroblastoma cell line GI-LI-N, which, according to the cell line typing analysis of the Cancer Genome Project (see Section 4.2.2), shares 96.0% identity with IMR-32, suggesting that they are derived from the same cancer. Even genes that are rarely amplified may therefore contribute to tumourigenesis. Likewise, the NF- κ B transcription factor family member *NFKB1* was amplified to copy number 4.8 in one cell line (HH) derived from an adult T-cell lymphoma-leukaemia. Polymorphisms of *NFKB1* are associated with susceptibility to a number of cancers, including oral squamous cell carcinoma, myeloma, and cancers of the colon, liver and breast (for review, see Sun and Zhang, 2007). Interestingly, *NFKB1* maps to a region that is involved in translocations in certain types of acute lymphoblastic leukaemia (Liptay *et al.*, 1992).

Other implicated oncogenes that were amplified in human cancer and disrupted by retroviral insertions include matrix metalloproteinase-13 (*MMP13*) and mothers against decapentaplegic homolog 7 (*SMAD7*). *MMP13* shows recurrent amplification and overexpression in cervical cancer (Narayan *et al.*, 2007) and 2 of the 12 cell lines in this

CIS P-value	Gene name	Mouse Ensembl ID	Human Ensembl ID	Number of cell lines	Genes in minimal amplified region	Maximum copy number	Known oncogene?
0.001	Мус	ENSMUSG0000022346	ENSG00000136997	71	3	5.9+	Y
0.001	Ccnd1	ENSMUSG0000031071	ENSG00000110092	24	10	5.9+	Y
	Nmyc1	ENSMUSG0000037169	ENSG00000134323	14	9	5.9+	Y
	Slamf6	ENSMUSG0000015314	ENSG00000162739	14	21	5.9+	
	Smad7	ENSMUSG0000025880	ENSG00000101665	6	27	5.9+	
	Fgfr2	ENSMUSG0000030849	ENSG0000066468	5	7	5.9+	Y
0.001		ENSMUSG0000062960	ENSG00000128052	5	15	5.9+	
	Tnfrsf7	ENSMUSG0000030336	ENSG00000139193	5	26	5.9+	
	Meis1	ENSMUSG0000020160	ENSG00000143995	1	2	5.9+	
	Mmp13	ENSMUSG00000050578	ENSG00000137745	12	21	4.8	
	Nfkb1	ENSMUSG0000028163	ENSG00000109320	1	7	4.8	
	Zfp217	ENSMUSG0000052056	ENSG00000171940	43	15	4.3	
	Zfpn1a1	ENSMUSG0000018654	ENSG00000185811	18	103	4.3	Y
	Ccnd3	ENSMUSG0000034165	ENSG00000112576	8	37	4.3	Y
	Lmo2	ENSMUSG0000032698	ENSG00000135363	4	43	4.3	Y
0.001		ENSMUSG0000024014	ENSG00000137193	7	12	3.8	Y
	Ccnd2	ENSMUSG0000000184	ENSG00000118971	6	15	3.8	Y
0.001		ENSMUSG0000027684	ENSG0000085276	17	27	3.1	Y
0.001		ENSMUSG0000020423	ENSG00000159388	10	35	3.1	
0.001		ENSMUSG0000028459	ENSG00000137101	8	26	3.1	
	Rreb1	ENSMUSG0000039087	ENSG00000124782	7	10	3.1	
0.001		ENSMUSG0000023938	ENSG00000124608	6	19	3.1	
0.001	Taok3	ENSMUSG0000061288	ENSG00000135090	3	9	3.1	
0.001		ENSMUSG0000020902	ENSG0000065320	2	31	3.1	
	Pik3r5	ENSMUSG0000020901	ENSG00000141506	2	31	3.1	
0.001	Eif4e3	ENSMUSG0000030068	ENSG00000163412	2	33	3.1	
0.001		ENSMUSG0000021356	ENSG00000137265	7	27	2.7	Y
0.001	Ubb	ENSMUSG0000019505	ENSG00000170315	5	72	2.5	
0.001	Cd69	ENSMUSG0000030156	ENSG00000110848	4	52	2.5	
0.001	Lrrc5	ENSMUSG0000046079	ENSG00000171492	2	25	2.5	
0.001	Ptpn1	ENSMUSG0000027540	ENSG00000196396	42	27	2.1	
0.001		ENSMUSG0000027636	ENSG00000101082	41	84	2.1	
	E030003N15Rik	ENSMUSG0000036661	ENSG00000105339	40	68	2.1	
	2310007D09Rik	ENSMUSG0000027654	ENSG00000101447	38	61	2.1	
	Capsl	ENSMUSG0000039676	ENSG00000152611	32	40	2.1	
	Cldn10	ENSMUSG0000022132	ENSG00000134873	20	37	2.1	
0.001		ENSMUSG0000051212	ENSG00000169508	18	31	2.1	
0.001		ENSMUSG0000042817	ENSG00000122025	11	122	2.1	Y
	Chc1l	ENSMUSG0000022106	ENSG00000136161	10	53	2.1	
0.001		ENSMUSG0000021998	ENSG00000136167	10	128	2.1	Y
	4933403F05Rik	ENSMUSG0000038121	ENSG00000177150	10	159	2.1	
0.001		ENSMUSG0000037474	ENSG00000143476	8	70	2.1	
	2410129E14Rik	ENSMUSG0000045136	ENSG00000137285	7	18	2.1	
	1110036003Rik	ENSMUSG0000006931	ENSG00000141696	6	43	2.1	
	Fmnl1	ENSMUSG0000055805	ENSG00000184922	6	85	2.1	
0.001		ENSMUSG0000018334	ENSG00000141068	5	34	2.1	
	Jundm2	ENSMUSG0000034271	ENSG00000140044	13	42	1.6	
	Tomm20	ENSMUSG0000058779	ENSG00000173726	8	252	1.6	
0.001		ENSMUSG0000024646	ENSG00000166347	6	20	1.6	
0.001		ENSMUSG0000031958	ENSG00000166816	6	74	1.6	
	Cbfa2t3h	ENSMUSG0000006362	ENSG00000129993	5	133	1.6	Y
	Zfp608	ENSMUSG0000052713	ENSG00000168916	3	30	1.6	
	2610307008Rik	ENSMUSG0000024349	ENSG00000184584	3	95	1.6	
	Hhex-rs2	ENSMUSG0000024986	ENSG00000152804	2	40	1.6	
	D930036F22Rik	ENSMUSG0000035181	ENSG00000129493	17	19	5.9+	
	Laptm5	ENSMUSG0000028581	ENSG00000162511	1	11	2.7	
	Emp3	ENSMUSG0000040212	ENSG00000142227	7	33	2.7	
0.05	Rai1	ENSMUSG0000062115	ENSG00000108557	5	72	2.1	

Table 4.4. Genes that are nearest to CISs in mouse lymphomas and are also promising candidates for targets of amplification in human cancer cell lines. "CIS *P*-value" is the minimum threshold for the significance of the CIS nearest to the given gene. "Number of cell lines" is the number of samples in which the gene is amplified to a copy number of greater than or equal to 1.6. "Genes in minimal amplified region" is the number of genes that co-occur with the CIS gene in the smallest region of amplification. "Maximum copy number" is the maximum copy number threshold above which the gene is identified as being amplified. "Known oncogene?" indicates whether the gene is a dominant cancer gene listed in the Cancer Gene Census.

Chapter 4

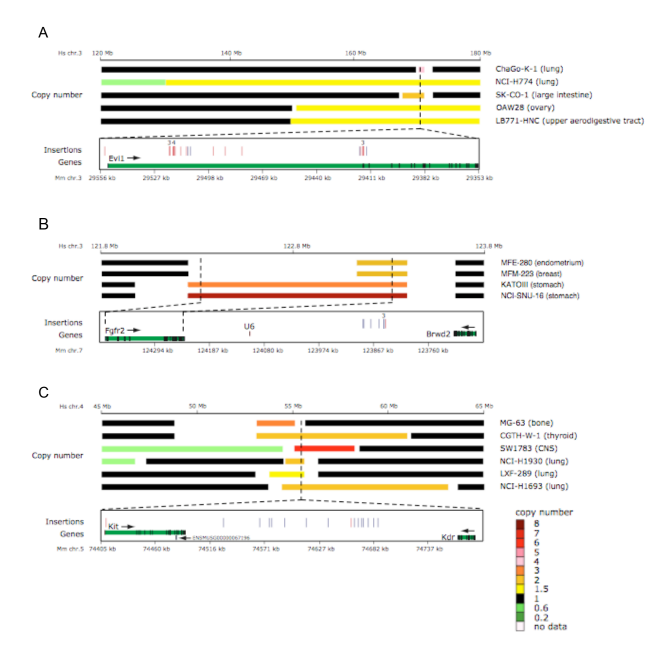


Figure 4.17. Known human oncogenes *EVI1* (A), *FGFR2* (B) and *KIT* (C) are amplified in human cancer cell lines and are disrupted by retroviral insertional mutagenesis in mouse lymphomas. The copy number of chromosomal regions in the human cell lines is depicted in colour. Names of human cell lines and tissue of origin are provided Only cell lines in which the amplicon containing the oncogene is less than 70 Mb are shown. The lower part of each figure shows insertions within mouse tumours, and was kindly provided by Jaap Kool and Jeroen de Ridder. Blue vertical lines represent insertions in the sense orientation, while red vertical lines represent antisense insertions. Genes are shown in green, with exons marked in black. Positions on the murine and human chromosomes are indicated on the black horizontal bars in kb and Mb, respectively. These figures can also be seen in Uren *et al.* (2008).

study in which *MMP13* was amplified were indeed derived from cervical cancers. *MMP13* has not been shown to be amplified in any other cancer types, but overexpression has been observed, e.g. in squamous cell carcinomas of the head and neck (Johansson *et al.*, 1997) and vulva (Johansson *et al.*, 1999). The results of this analysis suggest that *MMP13* is amplified in, and implicated in, a range of cancer types. The cell lines containing amplicons of less than 70 Mb that encompass *MMP13* are shown in Figure 4.18. Among these types are oesophageal, skin and breast cancers, in which *MMP13* overexpression has been observed (Freije *et al.*, 1994; Hu *et al.*, 2001; Kuivanen *et al.*, 2006). The minimal amplified region on chromosome 11 contains 21 genes, including a cluster of genes encoding matrix metalloproteinases, of which a number have been previously implicated in cancer. However, *MMP13* was the only gene disrupted by insertional mutagenesis.

SMAD7 duplication has been demonstrated in colorectal cancer (Boulay et al., 2001) and the gene is overexpressed in a number of cancer types, including basal cell carcinoma (Gambichler et al., 2007), endometrial cancer (Dowdy et al., 2005) and thyroid follicular carcinoma cell lines (Cerutti et al., 2003). The highest amplification of SMAD7 was in the retinoblastoma cell line Y79. Interestingly, SMAD7 has been shown to suppress TGF-β1-mediated growth inhibition in pancreatic cancer cells through the inactivation of the retinoblastoma protein (Boyer Arnold and Korc, 2005) and it inhibits growth arrest and apoptosis in mouse B cells through the inactivation of retinoblastoma (Ishisaki et al., 1998; Nakahara et al., 2003). In addition, SMAD7 is expressed in the eye, and suppresses TGF-β2-mediated inhibition of corneal endothelial cell proliferation, resulting in accelerated wound healing (Funaki et al., 2003). SMAD7 is therefore a promising target for amplification in the retinoblastoma cell line. Likewise, one of the amplicons encompassing SMAD7 was identified in a Ewing's sarcoma cell line (EW-24) and, in osteogenesis, SMAD7 suppresses osteoblast differentiation and bone formation (Koinuma and Imamura, 2005) and inhibits Saos2 osteosarcoma cell differentiation (Eliseev et al., 2006). SMAD7 was also amplified in 2 haematopoietic and 2 lung cancer cell lines. SMAD7 promotes self-renewal of haematopoietic stem cells (Blank et al., 2006) and is highly expressed in metastatic lung cancer cell lines (Shen et al., 2003).

Other interesting candidates include SLAM family member 6 precursor (*SLAMF6*), serine/threonine-protein kinase TAO3 (*TAOK3*), RAS-responsive element-binding protein 1 (*RREB1*) and leucine-rich repeat-containing protein 8D (*LRRC5*). The minimal

Chapter 4

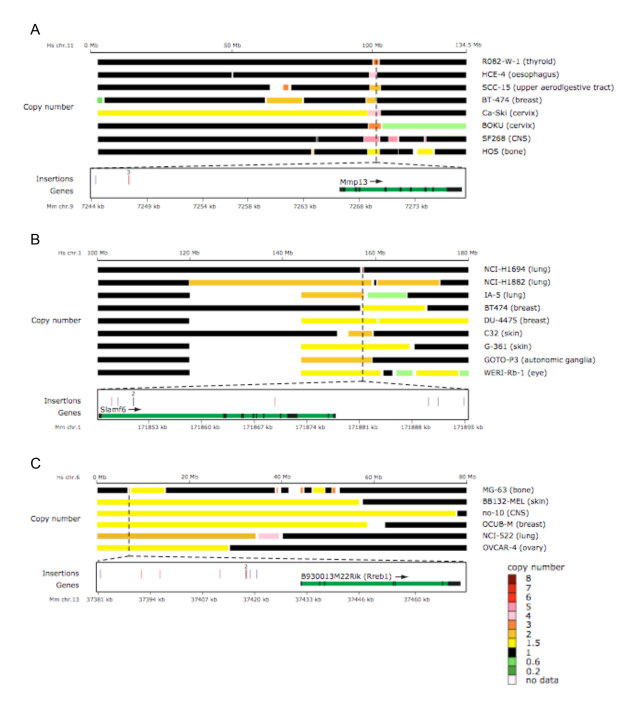


Figure 4.18. Candidate oncogenes *MMP13* (A), *SLAMF6* (B) and *RREB1* (C) are amplified in human cancer cell lines and are disrupted by retroviral insertional mutagenesis in mouse lymphomas. The copy number of chromosomal regions in the human cell lines is depicted in colour. Names of human cell lines and tissue of origin are provided Only cell lines in which the amplicon containing the oncogene is less than 70 Mb are shown. The lower part of each figure shows insertions within mouse tumours, and was kindly provided by Jaap Kool and Jeroen de Ridder. Blue vertical lines represent insertions in the sense orientation, while red vertical lines represent antisense insertions. Genes are shown in green, with exons marked in black. Positions on the murine and human chromosomes are indicated on the black horizontal bars in kb and Mb, respectively. These figures can also be seen in Uren *et al.* (2008).

amplified region encompassing SLAMF6 comprised 21 genes and was recurrent across 14 cell lines. Cell lines containing an amplicon of less than 70 Mb are shown in Figure 4.18. The highest amplification of SLAMF6 was within the lung cancer cell line NCI-H1694. However, it has been proposed that SLAMF6, also known as Ly108, is only expressed in lymphoid tissues (Peck and Ruley, 2000), where it regulates T cell development (Jordan et al., 2007) and B cell tolerance (Kumar et al., 2006). Polymorphisms within the gene are associated with systemic lupus erythematosus (Wandstrat et al., 2004), which has been widely associated with an increased risk of developing a range of cancers, but most strongly with cancers arising from B lymphocytes (Bernatsky et al., 2007). TAOK3 was only amplified in 3 cell lines, but the minimum region contained just 9 genes. TAOK3 is poorly characterised, but contains a somatic missense mutation in 2 lung cancers (small cell carcinoma cell line NCI-H28 and a primary adenocarcinoma) in the COSMIC database (Forbes et al., 2006). Although a role in tumourigenesis has not been demonstrated for TAOK3, protein kinases are widely implicated in cancer (see Sections 1.2.5.2 and 1.3.1). *RREB1* was amplified in 7 cell lines within a minimal region of 10 genes. Cell lines containing an amplicon of less than 70 Mb are shown in Figure 4.18. Each cell line was derived from a different tissue, but *RREB1* has been shown to be ubiquitously expressed in human tissues apart from the adult brain (Thiagalingam et al., 1997). Rreb1 binds to, and represses expression of, the *p16^{lnk4a}* promoter, and the development of pristine-induced plasma cell tumours in Balb/C mice is attributable to a polymorphism in this Rreb1 binding site (Zhang et al., 2003). In addition, RREB1 is important in reducing cell-cell adhesion and collective migration of epithelial cells (Melani et al., 2008), and it may therefore play a role in metastasis. RREB1 has also been identified as a transcriptional effector of RAL (Oxford et al., 2007), and RALA is itself implicated in cancer cell migration, as well as other cancer-related functions (Oxford et al., 2005). The most amplified occurrence of RREB1 was in the osteosarcoma cell line MG-63, but no role for *RREB1* has previously been elucidated in bone tissue. *LRRC5* was amplified in just 2 cell lines, with a minimal amplified region of 25 genes. Although little is known about this gene, it is thought that it might be implicated in the proliferation and activation of lymphocytes and monocytes, suggesting a possible role in the oncogenesis of B cells which would account for the insertions disrupting Lrrc5 in mouse lymphomas. However, LRRC5 was amplified in human cancer cell lines derived from the ovary and upper aerodigestive tract.

Only 4 additional candidates (D930036F22Rik, LAPTM5, EMP3 and RAI1) were identified using genes nearest to CISs with P < 0.05 (see Table 4.4). D930036F22Rik is also known as HEAT repeat containing 5A (HEATR5A). The minimal amplified region also included Rho-GTPase-activating protein 5 (p190-B), which is known to be overexpressed in breast cancer (Chakravarty et al., 2000), although only 1 breast cancer cell line contained an amplification of this region. Based on the distribution of insertions in the CIS, it is entirely possible that nearby genes Hectd1 and/or EG544864 were in fact the targets of MuLV mutagenesis (Figure 4.19). However, none of these genes have been previously implicated in tumourigenesis. Lysosomal-associated protein transmembrane 5 (LAPTM5) was amplified in a single cell line derived from an endometrial carcinoma (MFE-280). LAPTM5 is inactivated by chromosomal rearrangement and DNA methylation in human multiple myeloma (Hayami et al., 2003) but is overexpressed in malignant B lymphomas (Seimiya et al., 2003) and is a predictor for early intrahepatic recurrence of hepatocellular carcinoma (Somura et al., 2008). However, the amplified region in MFE-280 also contained the syndecan-3 gene, which is expressed in the human endometrium (Germeyer et al., 2007) and is thought to play a role in uterine growth (Russo et al., 2001). Epithelial membrane protein gene EMP3 was proposed as a candidate tumour suppressor in glioma and neuroblastoma (Alaminos et al., 2005), but it has since been shown to be overexpressed in oligodendroglial tumours (Li et al., 2007a) and primary glioblastomas (Kunitz et al., 2007). It is also overexpressed in invasive human mammary carcinoma cell lines (Evtimova et al., 2003) and contains a polymorphism in prostate cancers (Burmester et al., 2004). Retinoic acid induced 1 gene (RAI1) has not been previously implicated in cancer.

4.5.2.1.2 miRNA genes

The list of genes nearest to CISs with P < 0.001 contained 6 miRNA genes, while the list nearest to CISs with P < 0.05 contained 9. As mentioned in Section 3.2.2, deregulated miRNAs are implicated in promoting and suppressing tumourigenesis in a range of tissues. Currently, only protein-coding genes have human orthologues in Ensembl, and miRNA genes were therefore omitted from the global comparison and principal analysis of CIS genes within amplicons. However, it is possible to manually identify the human equivalents based on the miRNA name and the conserved synteny between the mouse and human genomes. Table 4.5 shows the name of the murine miRNA and the corresponding human miRNA for genes nearest to CISs, as well as lists of the miRNA genes within

Chr. 12 Length	52.70 Mb	52.75 Mb 52.80 Mb	52.85 Mb 52.90 Mb
MuLV	1		
 Ensembl trans. 	Ap4s1 >	EG544864 >	
DNA(contigs)	AC159644.3 >	< AC157213.2	< AC161114.4
 Ensembl trans. 	<pre>Hetd1 <hetd1 <hetd1="" <hetd<="" th=""><th>< Hoatr5a</th><th>₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩ < Q8I</th></hetd1></pre>	< Hoatr5a	₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩₩ < Q8I
	 Hectd1 		
MuLV			
Length	- <u> </u>	264.77 Kb	Reverse strand
	52.70 Mb	52.75 Mb 52.80 Mb	52.85 Mb 52.90 Mb

Figure 4.19. Insertions assigned to *Heatr5a* may be associated with *Hectd1* or EG544864. Insertions are shown as black vertical lines. Those above the blue bar labelled DNA(contigs) are in the sense orientation, those below are in the antisense orientation. Ensembl genes are shown in red.

A		
P-value	Mouse miRNA	Human miRNA
0.001	rno-mir-128b	hsa-mir-128b
0.001	hsa-mir-142	hsa-mir-142
0.001	mmu-mir-21	hsa-mir-21
0.001	mmu-mir-23a	hsa-mir-23a
0.001	mmu-mir-17	hsa-mir-17
0.001	hsa-mir-106a	hsa-mir-106a
0.05	mmu-mir-26b	hsa-mir-26b
0.05	mmu-mir-22	hsa-mir-22
0.05	rno-mir-200b	hsa-mir-200b

В

Mouse		Human		Number of	Maximum	Known oncogenes
miRNA gene	Mouse Ensembl ID	miRNA gene	Human Ensembl ID	cell lines	copy number	in minimal region?
mmu-mir-17	ENSMUSG0000065508	hsa-mir-17	ENSG00000198999	23	3.1	
hsa-mir-142	ENSMUSG0000065420	hsa-mir-142	ENSG00000199166	9	2.1	Y
mmu-mir-21	ENSMUSG0000065455	hsa-mir-21	ENSG00000199004	9	2.1	Y
mmu-mir-23a	ENSMUSG0000065611	hsa-mir-23a	ENSG00000199028	8	2.1	Y
rno-mir-128b	ENSMUSG0000065441	hsa-mir-128b	ENSG00000199105	1	2.1	

С

0					
Mouse		Human		Number of	Minimum
miRNA gene	Mouse Ensembl ID	miRNA gene	Human Ensembl ID	cell lines	copy number
rno-mir-128b	ENSMUSG0000065441	hsa-mir-128b	ENSG00000199105	40(2)	0.2
mmu-mir-17	ENSMUSG0000065508	hsa-mir-17	ENSG00000198999	61	0.6
mmu-mir-22	ENSMUSG0000065529	hsa-mir-22	ENSG00000199060	17	0.6
mmu-mir-26b	ENSMUSG0000065468	hsa-mir-26b	ENSG00000199121	8	0.6
mmu-mir-21	ENSMUSG0000065455	hsa-mir-21	ENSG00000199004	6	0.6
hsa-mir-142	ENSMUSG0000065420	hsa-mir-142	ENSG00000199166	5	0.6
mmu-mir-23a	ENSMUSG0000065611	hsa-mir-23a	ENSG00000199028	3	0.6

Table 4.5. miRNA genes that are nearest to CISs in mouse lymphomas and are amplified and/or deleted in human cancer cell lines. (A) Names of the murine miRNAs and their human orthologues. (B) Amplified miRNA genes. (C) Deleted miRNA genes. "P-value" is the minimum threshold for the significance of the CIS nearest to the given gene. "Number of cell lines" is the number of samples in which the gene is amplified to a copy number of greater than or equal to 1.6 (B) or deleted to a copy number of less than or equal to 0.6, with the number for deletions of copy number 0.2 or below shown in brackets (C). "Maximum copy number" is the maximum copy number threshold above which the gene is identified as being amplified. "Minimum copy number" is the minimum copy number.

human amplicons and deletions. Genes encoding 5 of the miRNAs were amplified in human cancer cell lines. The minimal amplified regions for hsa-miR-142 and hsa-miR-23a were very large and encompassed 4 and 3 known oncogenes, respectively, while hsamiR-21 was co-amplified with hsa-miR-23a, and hsa-miR-128b was amplified in just 1 cell line. The minimal amplified region of hsa-miR-17 contained 14 genes, of which none were oncogenes and only 2 had a description in Ensembl. hsa-miR-17 is part of the miR-17-92 cluster of 6 miRNAs. All 3 of the cell lines in which hsa-miR-17 was amplified to copy number 2.5 or above were derived from haematopoietic and lymphoid cancers, which is consistent with a role for miR-17-92 in both B-lymphocyte development and Blymphoproliferative disorders (Garzon and Croce, 2008). The cluster is also overexpressed in other human cancers, including colorectal cancers (Monzo *et al.*, 2008), anaplastic thyroid cancer cells (Takakura et al., 2008), neuroblastomas with MYCN amplification (Schulte et al., 2008), bladder cancers (Gottardo et al., 2007) and lung cancers (Hayashita et al., 2005). Of the 23 cell lines in which hsa-mir-17 was amplified to copy number 1.6 or above, 7 were from colon cancers, 7 were from haematopoietic and lymphoid cancers, 4 were from lung cancers, and 1 each were from cancers of the stomach, soft tissue, central nervous system, breast and eye. miR-17-92 is the likely target for amplification within this region, and demonstrates that the function of miRNAs may be conserved between species.

7 miRNA genes were identified within deletions of copy number 0.6 or below, but only *hsa-miR-17* and *hsa-miR-128b* were deleted in a large number of cell lines, and only *hsa-miR-128b* was within homozygous deletions. *hsa-miR-128b* was shown to be downregulated in classic Hodgkin lymphomas infected with Epstein-Barr virus (Navarro *et al.*, 2008). However, it is also upregulated in acute lymphoblastic leukaemia (Zanette *et al.*, 2007). The two homozygous deletions of *hsa-miR-128b* were within glioma and neuroblastoma cell lines, respectively. Interestingly, *hsa-miR-128* is highly expressed in the adult brain and preferentially in neurons, where it is thought to play a role in neural differentiation (Smirnova *et al.*, 2005). Downregulation of *hsa-miR-128* has been previously demonstrated in glioblastoma (Ciafre *et al.*, 2005), but deletion of the gene has not been previously demonstrated.

4.5.2.2 Candidate cancer genes among genes containing insertions within the coding region

Analyses involving the remaining lists were primarily designed to identify tumour suppressor genes, but as shown in Section 4.5.1.3, it is likely that the lists are contaminated with candidate oncogenes. The list for which the pattern of distribution was most similar to that of oncogenes was the list of genes containing insertions, not including those represented by a single read, within the coding region. The strongest candidates in regions of copy number gain, identified using the same filtering procedure as used for genes nearest to CISs, included 4 oncogenes (Table 4.6). As discussed in Section 3.4.1, insertions within the 3' UTR of *Mycn* and *Pim1* result in the formation of a more stable protein product, rather than gene inactivation. Some of the insertions were within the last exon of these genes, which explains their inclusion within the current list.

Candidate tumour suppressor genes were identified within regions of copy number loss. Among these was the gene encoding transmembrane protease, serine 2 precursor (TMPRSS2), which is a known oncogene that forms fusions with the ETS transcription factor genes ERG and ETV1 in prostate cancer (Tomlins et al., 2005) and is overexpressed in most prostate cancers (Vaarala et al., 2001). A hemizygous microdeletion within the fusion has been observed on chromosome 21 between ERG and TMPRSS2 (Yoshimoto et al., 2006), but this does not explain the deletion of the entire gene, sometimes in both copies. In addition, the homozygous deletions, which were also the most focal deletions, were identified in cancer cell lines derived from the pancreas and upper aerodigestive tract, rather than the prostate. Most of the heterozygous deletions were very large, encompassing many genes, and the minimal deleted region contained 18 genes. Based on the known role of *TMPRSS2*, it seems unlikely that this is the target of deletion within this region. Likewise, although deleted in human cancers, MAP3K8 and IL6RA are also more likely to act as oncogenes. MAP3K8 is overexpressed in, for example, invasive endometrioid cancer (Aparecida Alves et al., 2006), T-cell neoplasias (Christoforidou et al., 2004) and breast cancer (Sourvinos et al., 1999), while expression of the interleukin 6 receptor gene *IL6RA* is promoted by Epstein-Barr virus in immortalised B cells and Burkitt's lymphoma cells (Klein et al., 1995). The minimal amplified region containing the gene encoding MAGUK p55 subfamily member 4 (MMP4) comprised just 4 genes. Interestingly, MMP4 is a homologue of the Drosophila Stardust gene, which is involved in establishing and maintaining epithelial tissue polarity,

Cono nomo	Mourse Encombl TD	Human Encombl TD	Number of cell lines	Genes in minimal region	Copy	Singletons only?	Known
Gene name Capsl	Mouse Ensembl ID ENSMUSG00000039676	Human Ensembl ID ENSG00000152611	32	40	number 1.6+	Onlys	oncogene?
Bcl9	ENSMUSG0000039070	ENSG000001122011	17	40	1.6+		Y
Mycn	ENSMUSG00000037169	ENSG00000134323	14	9	1.6+		Y
Ccnd3	ENSMUSG00000034165	ENSG00000134325	8	37	1.6+		Y
Pim1	ENSMUSG00000034105	ENSG00000112578	7	12	1.6+		Y
NM 009283.2	ENSMUSG0000024014 ENSMUSG00000026104	ENSG00000115415	2	12	1.6+		1
Mrps18b	ENSMUSG0000024436	ENSG00000113413	12	12	0.6		
Mpp4	ENSMUSG0000024436	ENSG00000137330	9	4	0.6		
Il6ra	ENSMUSG0000028024 ENSMUSG00000027947	ENSG000000160712	3	23	0.6		
	ENSMUSG00000027347	ENSG00000134882			0.0		
Phgdhl1 Map3k8	ENSMUSG00000041765 ENSMUSG00000024235	ENSG00000134882 ENSG00000107968	63(1)	9(21)	0.2		
			35(1)	13(20)			v
Tmprss2	ENSMUSG0000000385	ENSG00000184012	14(2)	18(18)	0.2	V	Y
Tmem16f	ENSMUSG0000064210	ENSG00000177119	3	1	1.6+	Y	
Nfkb1	ENSMUSG0000028163	ENSG00000109320	1	7	1.6+	Y	
9030611019Rik	ENSMUSG0000036136	ENSG00000184731	3	16	1.6+	Y	
Olfr1509	ENSMUSG0000035626	ENSG00000182735	12	26	1.6+	Y	
	ENSMUSG0000046186	ENSG00000156535	3	35	1.6+	Y	
Rasgrp4	ENSMUSG0000030589	ENSG00000171777	12	56	1.6+	Y	
Dsg1b	ENSMUSG0000061928	ENSG00000134760	76	89	0.6	Y	
XP_484397.2	ENSMUSG0000034731	ENSG00000102780	68	8	0.6	Y	
Riok3	ENSMUSG0000024404	ENSG00000101782	68	47	0.6	Y	
6330406I15Rik	ENSMUSG0000029659	ENSG00000102802	64	9	0.6	Y	
Il17rb	ENSMUSG0000015966	ENSG0000056736	46	14	0.6	Y	
Zmynd11	ENSMUSG0000021156	ENSG00000015171	46	32	0.6	Y	
Hmgb2	ENSMUSG0000054717	ENSG00000164104	43	13	0.6	Y	
Gtse1	ENSMUSG0000022385	ENSG00000075218	41	90	0.6	Y	
1700020C11Rik	ENSMUSG0000004748	ENSG00000100010	36	61	0.6	Y	
Slc37a2	ENSMUSG0000032122	ENSG00000134955	34	70	0.6	Y	
Man1a	ENSMUSG0000003746	ENSG00000111885	33	8	0.6	Y	
Ate1	ENSMUSG0000030850	ENSG00000107669	32	6	0.6	Y	
Nrap	ENSMUSG00000049134	ENSG00000197893	32	9	0.6	Y	
Q91VN2_MOUSE	ENSMUSG0000042293	ENSG00000180425	32	16	0.6	Y	
Snf1lk2	ENSMUSG0000037112	ENSG00000170145	32	40	0.6	Y	
Dnajc9	ENSMUSG0000021811	ENSG00000182180	31	26	0.6	Y	
3110003A17Rik	ENSMUSG0000019855	ENSG00000146386	30	11	0.6	Y	
Shb	ENSMUSG0000044813	ENSG00000107338	29	16	0.6	Y	
Hp1bp3	ENSMUSG0000028759	ENSG00000127483	24	84	0.6	Y	
1200009I06Rik	ENSMUSG0000021280	ENSG00000185215	23	94	0.6	Y	
8430406I07Rik	ENSMUSG0000027424	ENSG00000125871	21	20	0.6	Y	
Wdr5b	ENSMUSG0000034379	ENSG00000196981	17	14	0.6	Y	
Zdhhc23	ENSMUSG0000036304	ENSG00000184307	17	44	0.6	Y	
9030611019Rik	ENSMUSG0000036136	ENSG00000184731	15	22	0.6	Y	
Gcnt2	ENSMUSG0000021360	ENSG00000111846	14	47	0.6	Y	
Itk	ENSMUSG0000020395	ENSG00000113263	13	31	0.6	Y	
	ENSMUSG0000039153	ENSG00000124813	13	97	0.6	Y	
Dok3	ENSMUSG0000035711	ENSG00000146094	13	183	0.6	Y	
Hivep3	ENSMUSG0000028634	ENSG00000127124	12	2	0.6	Y	
CSDE1_MOUSE	ENSMUSG0000068823	ENSG0000009307	11	25	0.6	Ŷ	
8430438D04Rik	ENSMUSG0000036019	ENSG00000179104	10	4	0.6	Ŷ	
Bcl10	ENSMUSG0000028191	ENSG00000142867	10	33	0.6	Ŷ	
Rgs2	ENSMUSG0000026360	ENSG00000116741		16	0.6	Ŷ	
Jmjd4	ENSMUSG00000036819	ENSG00000081692	, 7	93	0.6	Ý	
Tssk6	ENSMUSG00000047654	ENSG00000178093	, 6	60	0.6	Ý	
Tdrd5	ENSMUSG00000047034	ENSG000001/8093	5	46	0.6	Y	
Sell	ENSMUSG00000026581	ENSG00000182782	4	40 51	0.6	Y	
Leprel1	ENSMUSG0000026581 ENSMUSG00000038168	ENSG00000188404 ENSG00000090530	4 8(1)	29(33)	0.8	ř Y	
Olfr1509	ENSMUSG0000035626	ENSG00000090530			0.2	ř Y	
			19(3)	19(27)		ř Y	
Dut	ENSMUSG00000027203	ENSG00000128951	14(1)	7(7)	0.2		
3200002M19Rik	ENSMUSG0000030649	ENSG00000110200	11(1)	24(52)	0.2	Y	

Table 4.6. Mouse genes that contain retroviral insertions within the coding region and are also promising candidates for targets of amplification or deletion in human cancer cell lines. "Number of cell lines" is the number of samples in which the gene is amplified or deleted. "Copy number" is the maximum copy number threshold above which the gene is identified as being amplified, or the minimal threshold below which the gene is deleted. Where the copy number is 0.2, the number of cell lines and number of genes in the minimal deleted region are given for deletions of copy number ≤ 0.6 , with numbers for copy number ≤ 0.2 being shown in brackets. "Genes in minimal region" is the number of genes that co-occur with the CIS gene in the smallest region of amplification/deletion. "Singletons only?" indicates whether the gene contains insertions other than those represented by a single read. "Known oncogene?" indicates whether the gene is a dominant cancer gene listed in the Cancer Gene Census. which is disrupted in epithelial tumours. Although *MMP4* has not been implicated in cancer, expression of another family member, known as *MMP7*, has been demonstrated in tumours of the uterus and bladder, and in lymphomas (Katoh and Katoh, 2004).

Among the genes that contained insertions represented by a single read, 3 genes (*SHB*, *HIVEP3* and *BCL10*) stood out as potential tumour suppressor genes. Overexpression of the gene encoding the SHB adaptor protein causes increased activity of the pro-apoptotic kinase c-ABL, resulting in reduced tumour growth in PC3 prostate cancer cells (Davoodpour *et al.*, 2007). Therefore, it is possible that deletion of the gene may lead to tumour cell growth and proliferation. The human immunodeficiency virus type 1 enhancer binding protein 3 gene (*Hivep3*, also known as *Krc*), positively regulates transcription of the mouse metastasis-associated gene, *S100A4/mts1* (Hjelmsoe *et al.*, 2000). In addition, *KRC* was proposed as a potential tumour suppressor gene following the development of a teratoma from *KRC*-deficient embryonic stem cells introduced into an animal model (Allen *et al.*, 2002). Finally, the B-cell lymphoma/leukaemia 10 gene (*BCL10*) is a "hotspot" within the commonly deleted region 1p22.3 in mantle cell lymphomas. Interestingly, 5 of the 10 cell lines containing a deletion within this region were derived from tumours of the autonomic ganglia, but no role for *BCL10* has previously been demonstrated in these cancers (Balakrishnan *et al.*, 2006).

4.5.2.3 Candidate tumour suppressor genes among genes containing insertions within the translated or transcribed region

Candidates among the lists of genes containing insertions within the translated or transcribed region are combined in Table 4.7. The gene that was most frequently deleted below the copy number thresholds of both 0.6 and 0.2 was the known tumour suppressor gene *CDKN2A* (also known as the *INK4A/ARF* locus, and described in Section 1.2.6). This demonstrates the efficacy of the analysis, since homozygous and heterozygous deletions of *CDKN2A* are commonly observed in a wide range of cancers. The only other known tumour suppressor gene in the list, according to the Cancer Gene Census, was the gene encoding FAS, which is a member of the TNF receptor superfamily. Binding of the FAS ligand to the FAS receptor results in the formation of the death-inducing complex (DISC), which triggers apoptosis (for review, see Wajant, 2002). The implicated tumour suppressor gene *WWOX* was also frequently deleted. *WWOX* resides in a fragile site and therefore while it is frequently deleted in cancers, it is unclear whether it contributes to

Chapter 4

			Number of	Genes in minimal	Сору	Insertions in translated	Singletons	Known
Gene name Cdkn2a	Mouse Ensembl ID ENSMUSG00000044303	Human Ensembl ID ENSG00000147889	cell lines 207(145)	region	number 0.2	region?	only?	TSG?
	ENSMUSG00000044303 ENSMUSG00000033016	ENSG00000131196	112(2)	1(3) 15(15)	0.2	r Y		Ť
	ENSMUSG0000042439	ENSG0000074657	100(1)	14(14)	0.2			
	ENSMUSG0000052085 ENSMUSG00000033107	ENSG00000107099 ENSG00000101695	88(5) 78(1)	15(15) 14(14)	0.2			
	ENSMUSG00000048279	ENSG00000151835	64(1)	4(5)	0.2	Y		
	ENSMUSG0000021895	ENSG00000163947	47(2)	10(10)	0.2	Y		
	ENSMUSG0000039607	ENSG00000144642	43(4)	1(3)	0.2	Y		
	ENSMUSG0000032503 ENSMUSG00000019828	ENSG00000172995 ENSG00000152822	40(2) 37(6)	2(2) 2(1)	0.2 0.2	Y		
	ENSMUSG0000028029	ENSG00000164022	36(1)	12(17)	0.2	Ŷ		
	ENSMUSG0000024778	ENSG0000026103	35(2)	11(13)	0.2	Y		Y
	ENSMUSG00000040675 ENSMUSG00000024235	ENSG00000120254 ENSG00000107968	35(1) 35(1)	5(5) 13(20)	0.2	Y Y		
	ENSMUSG0000004637	ENSG00000186153	34(3)	1(2)	0.2	Ŷ		
	ENSMUSG0000019768	ENSG0000091831	34(1)	5(5)	0.2	Y		
	ENSMUSG0000052920 ENSMUSG00000019849	ENSG00000185532 ENSG00000085377	33(1) 33(1)	2(2) 10(19)	0.2 0.2	Y Y		
	ENSMUSG0000019820	ENSG00000152818	32(1)	2(1)	0.2	Ý		
	ENSMUSG0000033102	ENSG0000081377	31(1)	19(19)	0.2			
	ENSMUSG0000069601 ENSMUSG00000038578	ENSG00000151150 ENSG00000106868	29(1) 25(2)	1(1) 5(5)	0.2	Y Y		
	ENSMUSG00000054150	ENSG00000176438	25(2)	2(7)	0.2			
	ENSMUSG00000050103	ENSG00000187546	24(7)	1(1)	0.2	Y		
	ENSMUSG0000033713	ENSG0000053254	24(1)	17(17)	0.2	v		
	ENSMUSG00000056924 ENSMUSG00000027347	ENSG00000158321 ENSG00000172575	22(1) 21(6)	1(14) 6(7)	0.2	Y Y		
	ENSMUSG0000029763	ENSG00000131558	20(1)	1(3)	0.2	Ý		
lars2	ENSMUSG0000027430	ENSG00000125821	20(1)	10(20)	0.2	Y		
	ENSMUSG00000059060 ENSMUSG00000040003	ENSG00000182185 ENSG00000187391	19(1) 18(3)	3(11) 7(6)	0.2	Y Y		
	ENSMUSG00000030244	ENSG00000111713	17(1)	14(18)	0.2	Y		
Atg10 I	ENSMUSG0000021619	ENSG00000152348	16(4)	5(5)	0.2	Y		
	ENSMUSG0000030839 ENSMUSG00000037416	ENSG00000129158 ENSG00000172869	16(2) 15(1)	2(4) 23(23)	0.2	Y Y		
	ENSMUSG00000037416 ENSMUSG00000048285	ENSG00000172889 ENSG00000139926	15(1) 14(1)	12(12)	0.2	Y		
810060J02Rik	ENSMUSG0000030301	ENSG00000123106	14(1)	13(13)	0.2	Y		
	ENSMUSG0000001995	ENSG00000116991	12(1)	7(9)	0.2	Y		
	ENSMUSG0000020472 ENSMUSG00000048058	ENSG00000162714 ENSG00000179241	11(2) 11(2)	16(16) 3(3)	0.2	Y		
	ENSMUSG00000039167	ENSG00000162618	11(1)	2(2)	0.2	Ý		
	ENSMUSG0000024074	ENSG00000150938	9(1)	1(1)	0.2	Y		
	ENSMUSG0000027562 ENSMUSG00000034101	ENSG00000104267 ENSG00000198561	8(1) 8(1)	12(12) 17(17)	0.2	Y		
	ENSMUSG00000027684	ENSG00000085276	8(1)	19(26)	0.2			
Slc15a4	ENSMUSG0000029416	ENSG00000139370	7(1)	3(3)	0.2	Y		
	ENSMUSG0000033306	ENSG00000145012	7(1)	33(33)	0.2	Y Y		
	ENSMUSG0000028497 ENSMUSG00000024513	ENSG00000188921 ENSG00000134046	105 95	10 1	0.6 0.6	Y		
	ENSMUSG0000052942	ENSG00000107249	84	2	0.6	Ý		
	ENSMUSG0000022021	ENSG00000139734	74	5	0.6			
	ENSMUSG0000035078 ENSMUSG00000039945	ENSG00000104643 ENSG00000120162	70 66	26 1	0.6 0.6	Y		
	ENSMUSG00000022120	ENSG00000152193	66	4	0.6			
	ENSMUSG0000024052	ENSG00000101577	62	17	0.6			
	ENSMUSG0000024544 ENSMUSG00000022031	ENSG00000168675 ENSG00000134014	62 59	63 9	0.6 0.6	Y Y		
	ENSMUSG00000049717	ENSG00000174405	59	18	0.6			
	ENSMUSG0000018796	ENSG00000151726	51	44	0.6	Y		
	ENSMUSG0000030064 ENSMUSG00000035828	ENSG00000114541 ENSG00000198355	43 43	8 39	0.6 0.6	Y		
	ENSMUSG00000330067	ENSG00000114861	43	39	0.6	Y		
	ENSMUSG0000000708	ENSG00000114166	42	4	0.6	Y		
	ENSMUSG0000032035	ENSG00000134954	41	2	0.6	Y		
	ENSMUSG0000016087 ENSMUSG00000029084	ENSG00000151702 ENSG00000004468	40 40	8 20	0.6 0.6	Y		
	ENSMUSG0000042496	ENSG00000170325	39	10	0.6			
	ENSMUSG0000026723	ENSG00000107614	39	31	0.6	Y		
	ENSMUSG0000054328 ENSMUSG00000023868	ENSG00000177199 ENSG00000112541	38 37	94 15	0.6 0.6	Y		
	ENSMUSG0000023868 ENSMUSG00000022443	ENSG00000112541 ENSG00000100345	37	21	0.6	ř		
.ef1	ENSMUSG0000027985	ENSG00000138795	36	6	0.6			
	ENSMUSG00000039048	ENSG00000110074	36 35	8 4	0.6 0.6	Y		
	ENSMUSG0000039031 ENSMUSG00000041836	ENSG00000146376 ENSG00000132334	35	4	0.6	ř Y		
	ENSMUSG00000019845	ENSG00000074935	35	8	0.6	Ý		
	ENSMUSG0000037999	ENSG0000047365	34	6	0.6			
	ENSMUSG0000062110 ENSMUSG00000028931	ENSG00000184178 ENSG00000069424	33 33	1 11	0.6 0.6	Y		
	ENSMUSG00000027993	ENSG000000109654	33	32	0.6	Y		
CA_MOUSE	ENSMUSG00000041018	ENSG00000166056	32	1	0.6	Y		
	ENSMUSG00000049134 ENSMUSG00000058013	ENSG00000197893 ENSG00000138758	32 32	9 29	0.6 0.6	Y		
	ENSMUSG00000045973	ENSG00000138758	29	29 16	0.6			
Pip5k1a I	ENSMUSG0000024867	ENSG00000107242	29	17	0.6			
	ENSMUSG00000031785	ENSG00000159618	27	58	0.6			
	ENSMUSG00000048251 ENSMUSG00000027309	ENSG00000127152 ENSG00000088854	24 23	17 16	0.6 0.6	Y Y		
	ENSMUSG00000027459	ENSG00000125898	23	40	0.6			
	ENSMUSG0000021758	ENSG00000152670	20	15	0.6	Y		
	ENSMUSG0000052013 ENSMUSG00000030921	ENSG00000186265 ENSG00000132256	19 19	6 20	0.6 0.6	Y		
	ENSMUSG00000030921 ENSMUSG00000040321	ENSG00000132256 ENSG00000198146	19	17	0.6			
Man2a1 I	ENSMUSG0000024085	ENSG00000112893	17	9	0.6			
	ENSMUSG0000056219	ENSG00000198133	17	14	0.6			
	ENSMUSG0000026764 ENSMUSG00000021366	ENSG00000168280 ENSG00000095951	17 16	32 6	0.6 0.6	Y		
	ENSMUSG0000021388 ENSMUSG00000018995	ENSG00000095951 ENSG00000137513	16	8	0.6	r Y		
A <i>I875199</i> I	ENSMUSG00000021629	ENSG00000145740	16	10	0.6	Y		
SIc30a5			10	11	0.6	Y		
Slc30a5 I Pde3b I	ENSMUSG0000030671	ENSG00000152270	16					
Slc30a5 I Pde3b I Pnn I	ENSMUSG00000020994	ENSG00000100941	16	12	0.6	Ŷ		
5lc30a5 Pde3b Pnn Rff								

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Chapter 4

				Genes in		Insertions in		
C	Maura Engement ID	Human Encombl TD	Number of	minimal	Сору	translated	Singletons	Known
Gene name Slco3a1	Mouse Ensembl ID ENSMUSG0000025790	Human Ensembl ID ENSG00000176463	cell lines 15	region 1	number 0.6	region? Y	only?	TSG?
	LINSMUSG00000033147	ENSG00000163393	15	4	0.6	Ý		
St3gal6	ENSMUSG00000022747	ENSG00000064225	15	. 7	0.6	Ŷ		
Slc36a3	ENSMUSG0000049491	ENSG00000186334	15	8	0.6	Y		
[tpr5	ENSMUSG0000030287	ENSG00000123104	15	10	0.6	Y		
	ENSMUSG0000042590	ENSG0000086200	15	36	0.6			
ND 070550 1	ENSMUSG0000062252	ENSG00000197753	15	40	0.6	V		
NP_079558.1 Phf14	ENSMUSG0000005583 ENSMUSG00000029629	ENSG00000081189 ENSG00000106443	14 14	2 9	0.6 0.6	Y Y		
2810013C04Rik	ENSMUSG00000066411	ENSG00000173575	14	18	0.6	1		
Lyn	ENSMUSG00000042228	ENSG00000147507	14	20	0.6			
Cd53	ENSMUSG0000040747	ENSG00000143119	13	2	0.6			
St6galnac3	ENSMUSG0000052544	ENSG00000184005	13	2	0.6	Y		
Grik1	ENSMUSG0000022935	ENSG00000171189	13	6	0.6	Y		
Rab27a	ENSMUSG0000032202	ENSG0000069974	13	8	0.6	Y		
Zfhx1b	ENSMUSG0000026872	ENSG00000169554	13	10	0.6			
	ENSMUSG0000021703 ENSMUSG00000022913	ENSG00000164300 ENSG00000183527	13 13	13 14	0.6	Y		
Dscr2 1700001D09Rik		ENSG00000121933	13	26	0.6 0.6	Y		
Sh3gl3	ENSMUSG00000030638	ENSG00000140600	13	20	0.6	Y		
5dk1	ENSMUSG00000039683	ENSG00000146555	12	1	0.6	Ŷ		
livep3	ENSMUSG0000028634	ENSG00000127124	12	2	0.6	Y		
	ENSMUSG0000021676	ENSG00000145703	12	3	0.6			
Mgat5	ENSMUSG0000036155	ENSG00000152127	12	17	0.6			
Cdc42se2	ENSMUSG0000052298	ENSG00000158985	12	54	0.6			
Vdfy1	ENSMUSG0000004377	ENSG0000085449	11	1	0.6	Y		
Bard1	ENSMUSG0000026196	ENSG00000138376	11	3	0.6			
012Ertd553e	ENSMUSG0000020589	ENSG00000197872	10	1	0.6			
Vfia	ENSMUSG0000028565 ENSMUSG00000036019	ENSG00000162599 ENSG00000179104	10 10	1	0.6 0.6	Y Y		
8430438D04Rik Acvr1	ENSMUSG00000026836	ENSG00000179104 ENSG00000115170	10	4	0.6	T		
1pp4	ENSMUSG0000026024	ENSG0000003393	9	4	0.6	Y		
Slc39a11	ENSMUSG00000041654	ENSG00000133195	9	12	0.6	Ŷ		
	ENSMUSG0000053396	ENSG00000185676	8	32	0.6			
Dnmt3a	ENSMUSG0000020661	ENSG00000119772	7	22	0.6	Y		
1110014D18Rik	ENSMUSG0000059586	ENSG00000156831	7	29	0.6	Y		
Ccnl1	ENSMUSG0000027829	ENSG00000163660	7	29	0.6	Y		
Мус	ENSMUSG0000022346	ENSG00000136997	6	3	0.6			
	ENSMUSG0000054676	ENSG00000131943	6	12	0.6	Y		
Phf21a	ENSMUSG0000058318	ENSG00000135365	6	19	0.6	Y		
VM 011210 1	ENSMUSG00000057788 ENSMUSG00000026395	ENSG00000105671 ENSG00000081237	6 5	60 3	0.6	Y Y		
VM_011210.1 .rp12	ENSMUSG00000022305	ENSG00000147650	5	6	0.6 0.6	Y		
Stxbp4	ENSMUSG0000022546	ENSG00000166263	5	8	0.6	Ý		
Galnt14	ENSMUSG0000024064	ENSG00000158089	5	16	0.6	Ý		
Meis1	ENSMUSG0000020160	ENSG00000143995	4	3	0.6	Y		
Ccdc19	ENSMUSG0000026546	ENSG00000158710	4	40	0.6	Y		
Ndr7	ENSMUSG0000040560	ENSG0000091157	98(2)	2(3)	0.2	Y	Y	
Rfx3	ENSMUSG0000040929	ENSG0000080298	85(3)	3(3)	0.2		Y	
Vfib	ENSMUSG0000008575	ENSG00000147862	84(4)	2(2)	0.2	Y	Y	
ltr2a	ENSMUSG0000064286	ENSG00000189076	75(3)	354(6)	0.2	Y Y	Y Y	
5430573F11Rik	ENSMUSG0000034997 ENSMUSG00000039620	ENSG00000102468 ENSG00000170941	72(1) 67(1)	5(5) 3(11)	0.2	1	Y	
Gpc5	ENSMUSG00000022112	ENSG00000170391	65(2)	4(2)	0.2	Y	Y	
Tt1	ENSMUSG0000022112	ENSG00000102755	63(1)	6(6)	0.2	Ý	Ý	
Gas7	ENSMUSG0000033066	ENSG0000007237	57(2)	1(16)	0.2		Y	
Rac1	ENSMUSG0000001847	ENSG00000136238	45(1)	126(22)	0.2	Y	Y	
Park2	ENSMUSG0000023826	ENSG00000185345	42(5)	2(1)	0.2	Y	Y	
Robo1	ENSMUSG0000022883	ENSG00000169855	42(1)	4(6)	0.2	Y	Y	
Htr1f	ENSMUSG00000050783	ENSG00000179097	37(2)	9(9)	0.2		Y	
115 Pank1	ENSMUSG0000031712	ENSG00000164136	34(1)	4(4)	0.2	Y	Y Y	
Pank1 Slc44a1	ENSMUSG0000033610 ENSMUSG0000028412	ENSG00000152782 ENSG00000070214	33(2) 25(1)	1(1) 20(20)	0.2	ř Y	Y Y	
016Ertd472e	ENSMUSG00000022864	ENSG00000154642	24(4)	13(13)	0.2	Y	Y	
Ibxd3	ENSMUSG00000043621	ENSG00000162543	24(1)	5(5)	0.2		Y	
Arfrp2	ENSMUSG0000042348	ENSG00000185305	20(2)	1(7)	0.2		Ŷ	
Col19a1	ENSMUSG0000026141	ENSG0000082293	20(1)	2(15)	0.2	Y	Y	
Accn1	ENSMUSG0000020704	ENSG00000108684	18(1)	1(1)	0.2	Y	Y	
Rhoj	ENSMUSG0000046768	ENSG00000126785	17(1)	14(14)	0.2	Y	Y	
Jsp47	ENSMUSG0000059263	ENSG00000170242	16(2)	4(4)	0.2		Y	
	ENSMUSG0000058152	ENSG00000198108	16(1)	3(2)	0.2	Y	Y	
	ENSMUSG0000027203	ENSG00000128951	14(1)	7(7)	0.2	Y	Y	
	ENSMUSG0000025959	ENSG00000118263	13(1)	1(1)	0.2	Y	Y	
KIF7					0.2	Y	Y	
Klf7 Ifngr2	ENSMUSG0000022965	ENSG00000159128	12(1)	12(12)				
Dut Klf7 Ifngr2 Tmem16f	ENSMUSG00000022965 ENSMUSG00000064210	ENSG00000177119	10(1)	21(21)	0.2	Y	Y	
Klf7 Ifngr2	ENSMUSG0000022965							

Table 4.7. Mouse genes that contain retroviral insertions within the transcribed or translated region and are also promising candidates for targets of deletion in human cancer cell lines. "Number of cell lines" is the number of samples in which the gene is deleted. "Genes in minimal region" is the number of genes that co-occur with the CIS gene in the smallest region of deletion. "Copy number" is the minimal threshold below which the gene is deleted. Where the copy number is 0.2, the number of cell lines and number of genes in the minimal deleted region are given for deletions of copy number <= 0.6, with numbers for copy number <= 0.2 being shown in brackets. "Insertions in translated region?" indicates whether any of the insertions are within the translated region of the gene. "Singletons only?" indicates whether the gene contains insertions other than those represented by a single read. "Known TSG?" indicates whether the gene is a recessive cancer gene listed in the Cancer Gene Census.

tumourigenesis (see Section 1.3.3.3). The identification of insertions within the gene provides strong evidence that it does contribute to cancer (see also Section 3.4.3). Deletions of less than 70 Mb encompassing *WWOX* and insertions in *Wwox* are shown in Figure 4.20. In Section 3.4.3, *Foxp1* was proposed as a putative tumour suppressor gene. Deletion of *FOXP1* was observed in 42 cell lines, with a minimal amplified region of 3 genes, therefore providing additional evidence that this gene contributes to cancer and that it does so in both species. *Mobkl2a* was also presented as a putative tumour suppressor gene in Section 3.4.3, and while the human orthologue of this gene was not deleted in cancer, the human orthologue of paralogue *Mobkl2b* was deleted. Another implicated tumour suppressor gene identified in this analysis was *DOCK8*, which is deleted and under-expressed in human lung cancers (Takahashi *et al.*, 2006).

Known oncogenes EVII, MYC and FLII were also identified in the analysis, demonstrating that the results must be viewed with caution and that functional validation, as well as analysis of the distribution of insertions within the mouse candidate, is required to determine whether deletion of the identified genes is likely to contribute to tumourigenesis. Other candidates that have been implicated as oncogenes include *GRM1*, which plays an important role in the transformation of melanocytes in melanoma (Shin et al., 2008), RASGPR1, which contributes to tumour progression in murine skin cancer (Luke et al., 2007; Oki-Idouchi and Lorenzo, 2007) and, as mentioned in the previous sections, MAP3K8, MPP4 and MEIS1. Likewise, amplification and overexpression of genes encoding cyclin L1 (CCNL1), low-density lipoprotein receptor-related protein 12 precursor (LRP12) and glypican-5 (GPC5) have been demonstrated in human head and neck squamous cell carcinomas (Muller et al., 2006; Redon et al., 2002), oral squamous cell carcinomas (Garnis et al., 2004) and rhabdomyosarcomas (Williamson et al., 2007), respectively. It is therefore likely that other genes are the targets of deletion in the regions containing these known and implicated oncogenes. *GRM1* was the only gene for which the minimal deleted region did not contain additional genes. However, this does not prove that GRM1 must be the critical gene, since deletions affecting upstream and downstream genes may simply overlap at GRM1.

The list contains many genes for which there is limited evidence in the literature to suggest that they may act as tumour suppressor genes. The results of this analysis therefore lend further support to these findings. Some of these candidates (*RBMS3*, *PCAF*, *UTRN*, *ANK3*, *ACCN1*, *CDC14B*, *CHES1* and *PARK2*) are briefly discussed

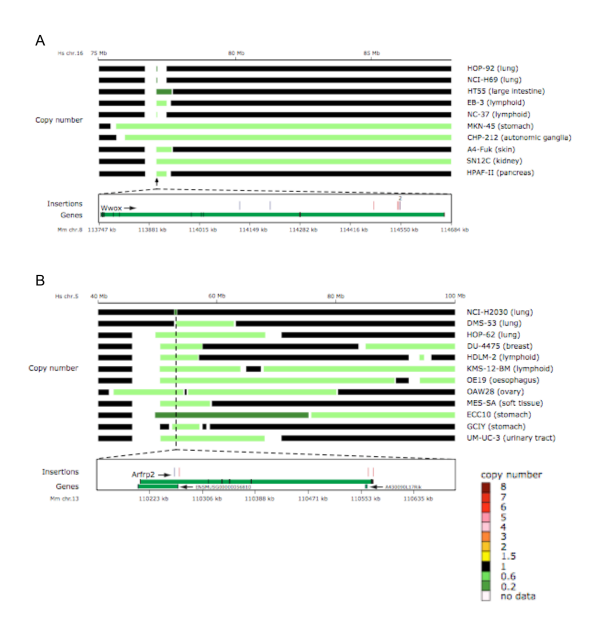


Figure 4.20. Candidate tumour suppressor genes *WWOX* (A) and *ARFRP2* (B) are deleted in human cancer cell lines and are disrupted by retroviral insertional mutagenesis in mouse lymphomas. The copy number of chromosomal regions in the human cell lines is depicted in colour. Names of human cell lines and tissue of origin are provided Only cell lines in which the deletion containing the gene is less than 70 Mb are shown. The lower part of each figure shows insertions within mouse tumours, and was kindly provided by Jaap Kool and Jeroen de Ridder. Blue vertical lines represent insertions. Genes are shown in green, with exons marked in black. Positions on the murine and human chromosomes are indicated on the black horizontal bars in kb and Mb, respectively. These figures can also be seen in Uren *et al.* (2008).

below. *RBMS3* and *PCAF* reside in commonly deleted regions, and are down-regulated, in oesophageal squamous cell carcinomas (Qin et al., 2008). One of the homozygous deletions that contained *RBMS3* was from an oesophageal cancer cell line (COLO-608N), but the remaining three were from the large intestine (NCI-H747), ovary (TYK-nu) and cervix (SKG-IIIa). The single homozygous deletion of PCAF was in a biliary tract cell line (EGI-1), and neither gene was deleted below copy number 0.6 in any additional oesophageal cancer cell lines. This suggests that the genes may also contribute to other cancers. The utrophin gene (UTRN) resides within a deletion of the long arm of chromosome 6 that is frequently observed in a range of tumours, and UTRN has been recently proposed as a putative tumour suppressor gene within this region (Li et al., 2007b). Ankyrin-3 (ANK3) is a target of the transcription factor hepatocyte nuclear factor 4 alpha that down-regulates cell proliferation in kidney cells (Grigo *et al.*, 2008). None of the 29 deletions containing ANK3 were within cell lines derived from kidney cancer, but the fact that this was the only gene in the minimal deleted region provides support for a role in tumour suppression. ACCN1 was proposed as a putative glioma tumour suppressor gene following the observation that surface expression of one of the two isoforms reduces cell growth and migration (Vila-Carriles et al., 2006), while the gene was also shown to be disrupted by a translocation within a neuroblastoma (Vandepoele et al., 2008). Notably, the single homozygous deletion containing this gene was within a glioma cell line (8-MG-BA), while 3 of the remaining 17 deletions of copy number less than or equal to 0.6 were within neuroblastomas. The rest of the deletions were in a range of tumours, including 3 breast, 2 bone, 2 lung and 2 ovarian. CDC14B and CHES1 are both involved in regulating cell cycle checkpoints related to DNA damage response (Bassermann et al., 2008; Busygina et al., 2006), and the deletion of these genes could therefore contribute to tumourigenesis by allowing damaged cells to enter mitosis. Like WWOX, the Parkin gene (PARK2) resides within a common fragile site (FRA6E) and, therefore, while the gene is frequently deleted in cancer, it is unclear whether it contributes to cancer development. However, deletions involving PARK2 are associated with ovarian cancer (Denison et al., 2003) and glioblastoma multiforme (Mulholland et al., 2006), and promoter hypermethylation of *PARK2*, resulting in down-regulation of gene expression, has been observed in leukaemias (Agirre et al., 2006). PARK2 is a long gene, measuring 994.53 kb, and contains just 2 insertions that could have occurred by Therefore, the presence of insertions within the gene does not provide chance. convincing support for a role in tumourigenesis. Interestingly, a break in FRA6E was associated with poor outcome in breast carcinomas, but expression of PARK2 was not

associated, while the loss of *AF-6* gene, which is telomeric of *PARK2*, was associated, suggesting that this may be a tumour suppressor gene affected by the break (Letessier *et al.*, 2007). Other candidates for which there is evidence in the literature of a tumour suppressive role in cancer include *BARD1*, *DMXL1*, *GPR56*, *HIVEP1*, *KCNAB2*, *LEF1*, *LIG4*, *PHF14*, *RAD51L1* and *RIPK3*. Further candidates *SDK1*, *BCL11B* and *MBD2* are discussed in Section 5.3.2.2.

ARFRP2 is a novel candidate tumour suppressor gene for which there is currently no evidence in the literature for a role in cancer. ARFRP2, also known as ARL15, is a member of the ADP-ribosylation factor-like family. Another member of this family, *ARL11*, is a tumour suppressor gene for which truncating germline mutations and promoter methylation contribute to leukaemia, breast cancer, ovarian cancer and melanoma (Frank *et al.*, 2005; Petrocca *et al.*, 2006). Deletions of less than 70 Mb that encompass *ARFRP2* are shown in Figure 4.20. There is also no evidence in the literature to suggest that the sec1 family domain containing gene *SCFD2* is a tumour suppressor gene. However, *SCFD2* is a transcriptional target of p53 (Krieg *et al.*, 2006), and it is the only gene within the minimal deleted region of 33 cancer cell lines.

4.6 Comparison of methods for calling gains and losses

As discussed in Section 4.3, DNAcopy and MergeLevels were the algorithms chosen for detecting regions of copy number change because they had been shown to perform better than other methods, and were freely available. However, it is not known whether DNAcopy and MergeLevels out-perform other methods in processing copy number data generated on the 10K SNP array CGH platform, and a variety of methods were therefore compared. The methods tested were DNAcopy alone (Olshen *et al.*, 2004), DNAcopy and MergeLevels (Olshen *et al.*, 2004; Willenbrock and Fridlyand, 2005), FASeg (Yu *et al.*, 2007), BioHMM (Marioni *et al.*, 2006) and a selection of the methods included within ADaCGH (Diaz-Uriarte and Rueda, 2007), i.e. CGHseg (Picard *et al.*, 2005), HMM (Fridlyand *et al.*, 2004), Wavelets (Hsu *et al.*, 2005) and GLAD (Hupe *et al.*, 2004).

27 different runs of DNAcopy version 1.4.0 were performed, each time varying the parameters. Alpha values of 0.1, 0.05, 0.01, 0.005 and 0.001 were tested, change-points that differed by less than 1, 2, 3 or 4 standard deviations were removed or all change-points were retained, and the smoothing step was either performed or was omitted from

the process (see Section 4.3 for details of these parameters). A further 17 runs of DNAcopy plus MergeLevels were performed, with various combinations of values for the DNAcopy parameters and the Wilcoxon and Ansari-Bradley thresholds within MergeLevels. The Wilcoxon rank sum test is used to determine whether there is a significant difference (according to the Wilcoxon threshold) between the observed values for two copy number levels, or whether they should be merged. The Ansari-Bradley 2-sample test determines whether there is any significant difference between the distribution of merged values minus observed log₂-ratios (i.e. the original ratios at individual SNPs) compared with the distribution of original segmented values minus observed log₂-ratios. The optimal Ansari-Bradley threshold is the largest threshold where the distributions do not differ significantly (Willenbrock and Fridlyand, 2005).

BioHMM is available as part of the BioConductor/R package, snapCGH. It is the only method that takes into account the distance between clones (or in this case SNPs), rather than simply ordering the clones or SNPs along the chromosome. BioHMM uses a Hidden Markov Model to segment data into a finite number of hidden states, where all of the data-points within a state have an equivalent copy number (Marioni *et al.*, 2006). A single run of BioHMM version 1.2.0 was performed using default parameters.

ADaCGH (analysis of data from aCGH) is a web-based tool that provides a selection of the best-performing methods via a simple user interface. DNAcopy and MergeLevels are available within this tool, but it is only possible to use default parameters and the MergeLevels output has been post-processed into three states: -1 (loss), 0 (no change) and 1 (gain). Methods within ADaCGH were chosen because they have been shown to perform well in the comparisons by Lai et al. (2005) and Willenbrock and Fridlyand (2005) and/or because they help to present a cross-section of the types of algorithm available for detecting copy number changes. CGHseg models the CGH data as a random Gaussian process and segments the data at points where the mean log₂-ratio changes abruptly. A threshold must be set for the adaptive penalisation, which is a threshold used to estimate the number of segments in the data. Picard et al. (2005) proposed a threshold of -0.05 as the default value, but Diaz-Uriarte and Rueda (2007) found that values around -0.005 were more appropriate but recommended experimenting with different values, which must be less than 0. For this analysis, 5 runs of CGHseg were performed, using thresholds of -0.005, -0.01, -0.05, -0.1 and -0.2. The smoothing approach of Hsu et al. (2005) uses wavelets to "denoise" the DNA copy number data and so to capture copy

number changes while smoothing out the noise. HMM is another method in which Hidden Markov Models are fitted to the data to identify different states, or copy number levels (Fridlyand *et al.*, 2004). However, unlike BioHMM, it does not take account of distances between data-points. Finally, the detection of breakpoints in GLAD is based on the Adaptive Weights Smoothing (AWS) procedure. This method finds the maximal neighbourhood around each data-point in which the local constant assumption holds true. In other words, it finds regions within which the copy number does not differ significantly and the boundaries of these regions represent breakpoints where the copy number changes. Default parameters were used for GLAD, HMM and the wavelets approach. All runs were performed in December 2007 on the website http://adacgh.bioinfo.cnio.es/.

FASeg, or Forward-Backward Fragment-Annealing Segmentation, is available as an R package from http://www.sph.emory.edu/bios/FASeg/. It is proposed to be especially suitable for SNP array CGH, which has a higher probe density but lower signal-to-noise ratio than traditional array CGH. According to the developers, the performance of FASeg was superior to 6 R packages, including DNAcopy, GLAD, BioHMM and CGHseg, in the detection of small segments with a low signal-to-noise ratio, although GLAD and BioHMM also performed well when the signal-to-noise ratio was low and the segments flanking copy number changes were long. When the signal-to-noise ratio was high, most methods performed well, although the HMM-based methods were less effective when there were multiple copy number levels within a single chromosome. This is a significant drawback, since multiple states are common in unstable cancer genomes. FASeg breaks each chromosome into small segments in an over-sensitive edge (or breakpoint) detection step that involves LOESS smoothing. It then iteratively merges consecutive segments until all remaining edges pass a significance threshold, based on testing for equal means between the groups of copy number values for SNPs before and after the edge using the unpaired Student's t-test. 15 different runs of FASeg version 1.2 were performed, in which parameters were altered for the smoothing span, which is the number of SNPs used to calculate the weights around each probe in the LOESS smoother, and the P-value cutoff for defining the significance of each edge. (See Yu et al., 2007)

In total, 69 different method and/or parameter combinations were compared. Each method was performed on the same 50 randomly selected cancer cell lines. The results were compared using Matthew's Correlation Coefficient (MCC), which is described in

Section 2.10.2. 280 Ensembl genes corresponding to known oncogenes involved in translocations or amplifications were extracted from the Cancer Gene Census. The number of known oncogenes and the number of other Ensembl genes within, and outside of, amplicons of copy number greater than or equal to 2.7 were counted. Oncogenes and other genes within amplicons were defined as true positives and false positives, respectively. Oncogenes and other genes that were not within amplicons were defined as false negatives and true negatives, respectively. The numbers of true and false positives and negatives in each cell line were then added together to give the number across all cell lines, and the MCC score was calculated. This analysis was performed individually on each method. It is possible that some of the known oncogenes that are involved in translocations are not amplified in human cancer, and of course there will be a proportion of non-oncogenes that are amplified in, and contribute to the development of, cancer. However, this analysis gives an indication of the performance of the method in comparison to other methods. The coverage was defined as the proportion of known oncogenes that were represented in amplicons, and the accuracy was defined as the proportion of genes in amplicons that were known oncogenes. The coverage, accuracy and MCC score for each method are shown in Table 4.8.

The wavelet, HMM and BioHMM algorithms all performed poorly. In the case of HMM and BioHMM, this may reflect the fact that there are often multiple copy number levels within a chromosome (see above). The low signal-to-noise ratio may account for the poor performance of the wavelet approach, since this method involves "denoising" the data but was developed for conventional array CGH data, which has a higher signal-tonoise ratio. Denoising the SNP CGH data may result in the removal of biologically relevant copy number changes. In addition, only the default parameters were used for this method. Changing the penalty constant in CGHseg made a considerable difference to the number of amplicons that were detected. This demonstrates the importance of choosing suitable parameter values. The closer the value was to 0, the greater the number of amplicons and the higher the coverage. However, the accuracy fell considerably. The default parameter value of -0.05 gave the best overall results, but this was lower than many of the results obtained using FASeg or DNAcopy. The value suggested in ADaCGH, i.e. -0.005, produced the highest coverage of all methods, but at the expense of a low accuracy. Although only the default parameters were used, GLAD performed reasonably well, obtaining similar results to the best-performing DNAcopy runs.

Method	Parameters	ТР	FP	TN	FN	Coverage	Accuracy	мсс
FASeg	p=0.01, smooth=7	31	1027	883573	13969	0.00221	0.02930	0.00380
FASeg	p=0.001, smooth=7	25	830	883770	13975	0.00179	0.02924	0.00340
FASeg	p=0.001, smooth=5	22	757	883843	13978	0.00157	0.02824	0.00301
DNAcopy & MergeLevels	alpha=0.05, smooth, SD=1	25	907	883693	13975	0.00179	0.02682	0.00293
DNAcopy & MergeLevels	alpha=0.1, smooth, w=0.00001	28	1055	883545	13972	0.00200	0.02585	0.00288
DNAcopy	alpha=0.05, smooth, SD=3	17	560	884040	13983	0.00121	0.02946	0.00284
FASeg	p=0.0001, smooth=10	18	608	883992	13982	0.00129	0.02875	0.00281
GLAD		25	931	883669	13975	0.00179	0.02615	0.00279
DNAcopy	alpha=0.01, smooth, SD=2	17	571	884029	13983	0.00121	0.02891	0.00275
DNAcopy	alpha=0.01, smooth, SD=1	18 28	620 1087	883980 883513	13982 13972	0.00129 0.00200	0.02821 0.02511	0.00272
DNAcopy & MergeLevels DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1 w=0.00001 alpha=0.1, smooth, SD=1 w=0.00001, ans=0.01	28		883513	13972	0.00200	0.02511	0.00271 0.00271
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1 w=0.00001, ans=0.01 alpha=0.1, smooth, SD=1 w=0.00001, ans=0.1	28	1087	883513	13972	0.00200	0.02511	0.00271
DNAcopy	alpha=0.05, smooth, $SD=1$ w=0.00001, ans=0.1	23	854	883746	13977	0.00164	0.02623	0.00269
DNAcopy	alpha=0.1, smooth, SD=2	23	855	883745	13977	0.00164	0.02620	0.00268
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1	28	1103	883497	13972	0.00200	0.02476	0.00263
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1, ans=0.01	28	1103	883497	13972	0.00200	0.02476	0.00263
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1, ans=0.1	28	1103		13972	0.00200	0.02476	0.00263
FASeg	p=0.01, smooth=10	24	911	883689	13976	0.00171	0.02567	0.00263
DNAcopy	alpha=0.1, smooth, SD=1	26	1007	883593	13974	0.00186	0.02517	0.00263
DNAcopy	alpha=0.05, smooth	23	865	883735	13977	0.00164	0.02590	0.00262
DNAcopy	alpha=0.05, smooth, SD=4	13	415	884185	13987	0.00093	0.03037	0.00261
DNAcopy	alpha=0.01, smooth	18	636	883964	13982	0.00129	0.02752	0.00260
DNAcopy & MergeLevels	alpha=0.1, smooth	27	1061	883539	13973	0.00193	0.02482	0.00260
DNAcopy	alpha=0.005, smooth, SD=2	16	558	884042	13984	0.00114	0.02787	0.00251
DNAcopy	alpha=0.1, smooth	26	1029	883571	13974	0.00186	0.02464	0.00251
DNAcopy	alpha=0.01, smooth, SD=4	12	383	884217	13988	0.00086	0.03038	0.00251
DNAcopy	alpha=0.05, smooth, SD=2	20	744	883856	13980	0.00143	0.02618	0.00250
DNAcopy CGHseg	alpha=0.01, smooth, SD=3 penalty=-0.05	14 16	471 561	884129 884039	13986 13984	0.00100 0.00114	0.02887 0.02773	0.00249 0.00249
DNAcopy	alpha= 0.001 , smooth, SD= 2	15	526	884074	13985	0.00114	0.02773	0.00249
FASeg	p=0.001, smooth=10	20	758	883842	13980	0.00107	0.02571	0.00241
DNAcopy	alpha=0.001, smooth	16	575	884025	13984	0.00113	0.02707	0.00238
DNAcopy	alpha=0.001, smooth, SD=4	12	396	884204	13988	0.00086	0.02941	0.00238
DNAcopy	alpha=0.005, smooth, SD=4	12	396	884204	13988	0.00086	0.02941	0.00238
DNAcopy & MergeLevels	alpha=0.01, smooth, w=0.00001	19	717	883883	13981	0.00136	0.02582	0.00237
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1 w=0.001	26	1059	883541	13974	0.00186	0.02396	0.00235
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1 w=0.001, ans=0.01	26	1059	883541	13974	0.00186	0.02396	0.00235
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=1 w=0.001, ans=0.1	26	1059	883541	13974	0.00186	0.02396	0.00235
DNAcopy & MergeLevels	alpha=0.1, smooth, SD=2	21	816	883784	13979	0.00150	0.02509	0.00234
DNAcopy	alpha=0.001, smooth, SD=1	16	582	884018	13984	0.00114	0.02676	0.00233
DNAcopy	alpha=0.005, smooth, SD=1	16	587	884013	13984	0.00114	0.02653	0.00229
DNAcopy	alpha=0.001, smooth, SD=3	13	454	884146	13987	0.00093	0.02784	0.00226
DNAcopy & MergeLevels	alpha=0.01, smooth, SD=1	16	593	884007	13984	0.00114	0.02627	0.00225
DNAcopy	alpha=0.005, smooth, SD=3	13	458	884142	13987	0.00093	0.02760	0.00222
FASeg	p=0.0001, smooth=7	18 16	694 603	883906 883997	13982 13984	0.00129 0.00114	0.02528 0.02585	0.00221 0.00218
DNAcopy DNAcopy & MergeLevels	alpha=0.005, smooth	15	568	883997 884032	13984	0.00114	0.02585	0.00218
FASeq	alpha= 0.005 , smooth, w= 0.00001 p= 0.1 , smooth= 10	28		883339	13965	0.00107	0.02373	0.00209
CGHseq	penalty= -0.01	37	1751	882849	13963	0.00264	0.02069	0.00184
DNAcopy & MergeLevels	alpha=0.05, smooth, w=0.00001	24	1119	883481	13976	0.00171	0.02100	0.00156
DNAcopy	alpha=0.01	21	961		13979	0.00150	0.02138	0.00155
DNAcopy	alpha=0.001	16	703	883897	13984	0.00114	0.02225	0.00152
DNAcopy	alpha=0.005	19	891	883709	13981	0.00136	0.02088	0.00136
DNAcopy	alpha=0.05	24	1169	883431	13976	0.00171	0.02012	0.00134
CGHseg	penalty=-0.01	5	187	884413	13995	0.00036	0.02604	0.00123
FASeg	p=0.1, smooth=5	33	1710	882890	13967	0.00236	0.01893	0.00119
CGHseg	penalty=-0.2	2	72	884528	13998	0.00014	0.02703	0.00084
DNAcopy	alpha=0.1	23		883298	13977	0.00164	0.01736	0.00055
BioHMM		15	849	883751	13985	0.00107	0.01736	0.00045
CGHseg	penalty=-0.005	42		882108	13958	0.00300	0.01657	0.00043
FASeg	p=0.01, smooth=5	8		884162	13992	0.00057	0.01794	0.00042
FASeg	p=0.001	4		884336	13996	0.00029	0.01493	-0.00009
FASeg	p=0.0001	4		884336	13996	0.00029	0.01493	-0.00009
FASeg	p=0.000001	4	264		13996	0.00029	0.01493	-0.00009
FASeg	p=0.0001, smooth=50	5	370	884230	13995	0.00036	0.01333	-0.00037
Wavelets		25	1769	882831	13975	0.00179	0.01394	-0.00059
HMM	0.01	9	717	883883	13991	0.00064	0.01240	-0.00073
FASeg	p=0.01	5	467	884133	13995	0.00036	0.01059	-0.00092

Table 4.8. Comparison of methods for detecting regions of copy number gain in 50 randomly selected cancer cell lines. Abbreviations for describing parameters are as follows: FASeg: p=significance threshold, smooth=smoothing range; DNAcopy: alpha=parameter alpha, smooth=outliers smoothed, SD=change-points differing by less than X standard deviations removed; MergeLevels: w=Wilcoxon threshold, ans=Ansari-Bradley threshold; CGHseg: penalty=penalty constant. Undefined parameters are default. TP=number of true positives (amplified oncogenes), FP=number of false positives (amplified non-oncogenes), TN=number of true negatives (non-amplified nononcogenes), FN=number of false negatives (non-amplified oncogenes). Numbers are calculated across all cell lines. Coverage=TP/(TP+FN), Accuracy=TP/(TP+FP). MCC = Matthew's Correlation Coefficient. Of the runs involving DNAcopy alone, those in which the data were not smoothed before segmentation performed worst. Higher values for the parameter alpha, which result in increased sensitivity, generally performed better due mainly to a higher coverage. For the purposes of the cross-species comparison, higher coverage, even at the expense of lower accuracy, is preferable since the mouse candidate cancer genes help to identify the targets of amplification in the human amplicons, and false positives are therefore likely to be ignored. Reducing the number of standard deviations below which change-points were removed resulted in a higher coverage of oncogenes. This may be because the highest peak of amplification, which often contains the critical cancer gene(s), is more likely to remain distinct from lower level copy number gains and the segment will have a higher mean copy number and will contain fewer amplified passengers. For higher values of alpha, merging the segments using default parameters also resulted in higher coverage. However, upon inspection of the results, it appeared that some oncogenes were lost upon merging, while some were gained. All of the oncogenes that were unique to the run without merging were still amplified in the run with merging, and vice versa, but they did not reach the copy number threshold of greater than or equal to 2.7. This is because merging increases the mean copy number of some segments and decreases the mean copy number of others, in line with the copy numbers of other segments in the genome. This demonstrates why it is useful to use a range of copy number thresholds in the comparative analysis. Changing the Ansari-Bradley threshold from 0.1 to 0.01 made no difference to the results, but lowering the value for the Wilcoxon threshold increased the MCC score. Using a lower value for the Wilcoxon threshold means that a higher proportion of segments will not be considered significantly different from one another and will therefore be merged. However, more detailed analysis suggests that lowering the value may not produce sensible results. For example, using a value of 1.0×10^{-5} rather than 1.0×10^{-4} , the segment of copy number 3.00 that contains *CCND1* in the neck squamous cell carcinoma cell line SCC-15 is merged with a segment of copy number 1.78 to give an overall copy number of 1.85. While the oncogene is still amplified, merging of this kind removes the peaks in amplification, which are most likely to harbour the critical targets of amplification. Similarly, a segment of copy number 0.12 on chromosome 4 of the bone cancer cell line CAL-72 is merged with other segments to give a copy number of 0.47. This segment is likely to be a homozygous deletion but is merged with segments that are more likely to represent heterozygous deletions.

Comparison of the FASeg runs showed that using the default smoothing span of 25 rather than a lower value resulted in lower accuracy and coverage and, therefore, a lower MCC score. When a significance threshold of P=0.0001 was used, a smoothing span of 10 rather than 50 not only identified more oncogenes (18 rather than 4) but also had tighter amplicon boundaries that still retained the oncogene. For example, lung cancer cell line LC-2-ad and pancreatic cancer cell line HuP-T4 contained amplicons that encompassed the oncogenes MYC and POU5F1, respectively. Using smoothing spans of 50 and 10, the number of SNPs within the amplicon containing MYC was calculated as 17 and 15, respectively, while the number within the amplicon containing *POU5F1* was 102 and 94, respectively. Increasing the significance threshold generally decreased the MCC score. Using a smoothing span of 7, a significance threshold of *P*=0.001 yielded 64 amplicons, while a significance threshold of P=0.0001 yielded 43 amplicons. 35 amplicons were identical, while the rest were either missing from the latter run or were shared but spanned a larger region when the threshold was higher. For example, an amplicon in the lung cancer cell line ChaGo-K-1 spanned 1.39 Mb and had a mean copy number ratio of 3.97 using a threshold of P=0.0001, and 919.42 kb with a mean copy number ratio of 4.16 using a threshold of P=0.001. Likewise, the amplicon encompassing MYC in lung cancer cell line LC-2-ad was also larger (6.74 Mb rather than 4.77 Mb) using a threshold of P=0.0001 rather than P=0.001 and had a lower mean copy number (5.06 rather than 5.49). The amplicons that were missing from the run with a higher significance threshold may still be present, but the mean copy number may not reach the copy number threshold of 2.7 because a larger region, including less amplified or non-amplified SNPs, is defined as the region of copy number change and this dilutes the mean copy number ratio for the entire segment. Overall, using a significance threshold of P=0.01 and a smoothing span of 7 appeared to give the best results, with the highest MCC score and highest accuracy and coverage. It is worth noting, however, that the parameter value for the smoothing span is well below that recommended in Yu et al. (2007). The results obtained using the top scoring runs from FASeg and DNAcopy plus MergeLevels were compared. 20 cancer genes were identified by both algorithms. 11 were unique to the FASeg output, and 5 were unique to the DNAcopy and MergeLevels output. In most cases, the missing genes were still amplified, but were below the copy number threshold of 2.7. This analysis indicates that the choice of method and parameters can make a considerable difference to the output and involves finding a suitable balance between accuracy and coverage.

4.7 Global comparison of mouse candidate cancer genes and human CNVs

The global comparison method of Section 4.5.1.1 was applied to human CNVs (see Section 4.2.3) and the gene lists from Section 4.2.1. Rather than using copy number thresholds, CNVs were separated into deletions and duplications, which were specified in the original downloaded file. As with previous analyses, the number of deletions/duplications within which each gene resided was counted. For each number of deletions/duplications, the number of mouse candidates was compared to the distribution of randomised genes using the Z-test. The results are depicted in Figure 4.21. None of the gene lists showed over-representation within deletions or duplications. The only positive association was observed for 7 known oncogenes (namely DDIT3, NSD1, IRF4, 2 genes encoding Histone H4, NUT and PDE4DIP) that were within 32 or more duplications. The association increased as the number of duplications increased, to a maximum of $P=2.07 \times 10^{-4}$ for 5 known oncogenes in 93 or more duplications. This suggests that some oncogenes are amplified in the normal population, and these individuals may have a predisposition to cancer. However, in general, genes involved in cancer were not found within CNVs. In fact, genes nearest to CISs (P<0.001 and P<0.05) and genes with insertions in coding regions were slightly under-represented in deletions, and genes within translated and transcribed regions were highly under-represented in both duplications and deletions. Many of the genes that are involved in oncogenesis are also involved in other important cellular functions, and this may explain why candidate oncogenes are rarely deleted in healthy individuals. Duplication of tumour suppressor genes could also lead to oncogene repression, producing a similar outcome, while deletion of tumour suppressor genes could lead to tumourigenesis. The results show that cells do not tolerate changes in copy number in genes that are important in tumourigenesis.

For each gene list, the number of genes residing within CNV deletions and within regions of copy number loss (less than or equal to a ratio of 0.6) in human cancer cell lines was counted. A 2-tailed Fisher Exact Test was performed to determine whether there was any association between genes found in deletions in normal individuals and deletions in cancer cell lines. The same analysis was performed using CNV duplications and regions of copy number gain (greater than or equal to 2.7). The *P*-values are provided in Table 4.9. In accordance with the results obtained in the global analysis, there was an under-

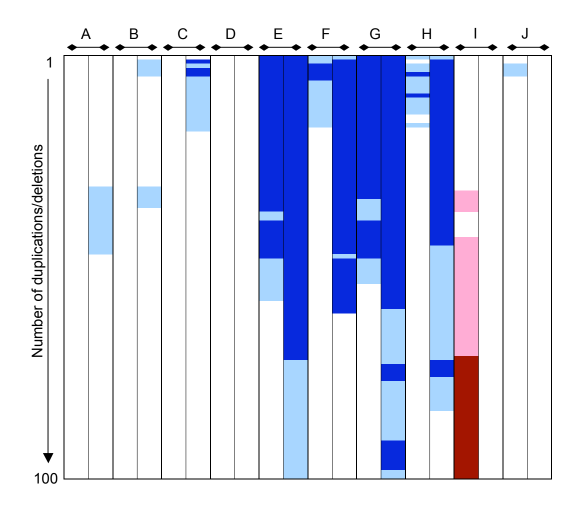


Figure 4.21. Under- and over-representation of human orthologues of candidate cancer genes in regions of copy number variation (CNV). (A) Genes nearest to CISs with P<0.001. (B) Genes nearest to CISs with P<0.05. (C) Genes with insertions within the coding region. (D) Genes with insertions but no singletons in the coding region. (E) Genes with insertions within the translated region. (F) Genes with insertions but no singletons in the translated region. (G) Genes with insertions in the transcribed region. (I) Genes with insertions but no singletons in the translated region. (I) Known oncogenes. (J) Known tumour suppressor genes. For each gene list, the left-hand column represents the significance of the association between the genes and CNV duplications, with rows representing the number of duplications, increasing in increments of 1 to a maximum of 100. Each box in the right-hand column represents the significance of the association between the genes and CNV deletions. P<0.01, dark blue for under-representation and dark red for over-representation; P<0.05, light blue for under-representation and pink for over-representation.

Gene list	Deletions	Amplicons
ORF only	3.76E-04	0.442
ORF only (no singletons)	6.12E-02	0.290
Translated region only	2.70E-10	0.780
Translated region only (no singletons)	7.02E-06	0.747
Transcribed region only	2.77E-14	0.082
Transcribed region only (no singletons)	6.68E-09	0.178
CIS nearest P<0.05	2.07E-04	5.12E-03
CIS nearest P<0001	2.09E-04	0.229

Table 4.9. *P*-values for the co-occurrence between genes from each gene list within CNVs and regions of copy number change in human cancer cell lines. "Deletions" gives the *P*-values for the co-occurrence of genes in CNV deletions and deletions of copy number less than or equal to 0.6 in human cancers, while "Amplicons" gives the *P*-values for the co-occurrence of genes in CNV duplications and amplicons of copy number greater than or equal to 2.7 in human cancers. *P*-values were calculated using a 2-tailed Fisher Exact Test. All significant *P*-values in "Deletions" represent an underrepresentation of genes in both CNVs and cancer deletions, while the significant *P*-value in "Amplicons" represents an over-representation of genes in CNVs and cancer amplicons.

representation in all lists of genes that co-occurred in both CNV deletions and deletions in human cancers. There was no association between genes in CNV duplications and copy number gains in human cancers, except for genes nearest to CISs with P<0.05, for which more genes than expected co-occurred in CNVs and amplicons. Again, this suggests that amplification of these genes in the general population may confer a predisposition to the development of cancer.

4.8 Discussion

The most significant finding from this chapter is that retroviral insertional mutagenesis is relevant to the discovery of cancer genes in regions of copy number change in human cancers. As anticipated, the overlap is stronger between candidate oncogenes and regions of copy number gain than between candidate tumour suppressor genes and regions of copy number loss. This partly reflects the fact that retroviral insertional mutagenesis predominantly identifies oncogenes due to the major mechanisms by which the retrovirus mutates genes and the requirement for both copies of a tumour suppressor gene to be It may, however, facilitate the identification of mutated (see Section 3.4). haploinsufficient tumour suppressor genes, for which the deletion of one gene copy can contribute to cancer. The other reason for the weaker association between tumour suppressor genes and deletions is that all genes that contained at least one insertion within the transcribed, translated or coding region were included in the analysis. Firstly, this can result in the inclusion of oncogenes that are activated by intragenic truncating mutations (see Section 3.4) and, secondly, many of the insertions may have occurred randomly and may not contribute to oncogenesis. However, the kernel convolution-based method for identifying CISs (de Ridder et al., 2006, see Section 2.10.2) is biased towards oncogenes because insertions within many parts of a tumour suppressor gene may cause its inactivation and therefore insertions may not cluster into tight CISs. For this reason, including all genes provides a more comprehensive list of candidates for a role in tumour suppression.

Significantly, CIS genes were over-represented in amplicons from both haematopoietic and lymphoid cell lines and lines derived from solid tumours. This demonstrates that retroviral insertional mutagenesis is relevant to the discovery of cancer genes in cancers other than lymphomas. This is also proven in the identification of individual candidates, since many were amplified or deleted in a range of cancer types, and some, including *MEIS1, MMP13* and *ACCN1*, were amplified or deleted in cancer types in which they had previously been implicated. While this study does not include any functional validation, the candidates include a considerable number of known and implicated cancer genes, demonstrating that the method is effective. In general, the discussion of individual genes has focussed on those for which there is some evidence, albeit sometimes limited, that gives cause for presenting the genes as potential oncogenes or tumour suppressor genes. However, the genes listed in Tables 4.4, 4.6 and 4.7 provide a large number of novel candidates that may be of interest to the cancer community. Interestingly, candidate cancer genes were under-represented in CNVs in apparently healthy individuals, further suggesting that amplification and/or deletion of these genes can have a detrimental effect on the cell and, in turn, on the individual.

Despite the promising results, there are a number of potential limitations associated with the analysis. Firstly, all of the human cancers were cell lines, rather than primary tumours. Cancer cells cultured in vitro lack the microenvironment of the tumour from which they are derived. While this means that they may not be fully representative of the original tumour, the homogeneity of cell lines can be an advantage since it prevents contamination by stromal cells and potential dilution of the copy number changes identified by CGH. It is, however, possible that the phenotype and genotype of cancer cell lines may differ from those of the original tumour due to genomic instability. Gene expression profiling of lung tumours and cell lines has demonstrated that, in culture, adenocarcinomas progress towards poorly differentiated phenotypes with expression profiles similar to those for squamous cell and small cell lung carcinomas (Virtanen et al., 2002). However, comparisons of human breast and lung cancer cell lines and their corresponding tumours demonstrated an extremely high correlation for both genotype and phenotype, concluding that cell lines from both cancer types are suitable model systems for the original tumours (Wistuba et al., 1998; Wistuba et al., 1999). In addition, gene expression profiles for the NCI60 cell lines, which are the most commonly used cancer cell lines in cancer research and constitute a proportion of the cell lines used in this chapter, also showed that most were representative of their corresponding tumour types (Wang *et al.*, 2006b). Therefore, the use of cancer cell lines is warranted in this analysis, especially as the study is generally concerned with the number of copy number changes affecting a gene, rather than the tissue specificity.

A second potential drawback is that the ploidy of the cancer cell lines is not known. None of the methods used for detecting copy number changes within CGH data can determine the ploidy, and yet an uploidy is a common characteristic of cancers. Attempts were made to determine the ploidy of cell lines based on the copy numbers of merged segments since, for example, a triploid cell line should only have copy number gains of 1.33, 1.67, 2.00, 2.33, 2.67, and so on, while a tetraploid should have copy number gains of 1.25, 1.5, 1.75, 2, 2.25, and so on. However, the mean copy number ratios for segments are not accurate enough to reliably assign cancers to a particular state. Irrespective of the ploidy, a copy number ratio of 3 indicates that there is a 3-fold increase in the number of copies. In this study, it is assumed that the balance of genes is more important than the actual number, i.e. a 3-fold increase in the number of copies of an oncogene is expected to have the same effect whether the baseline copy number is 2 or 4 genes. In addition, this study is concerned less with the exact copy number of genes, and more with whether genes are amplified or deleted, and the use of a set of copy number thresholds, rather than just one for amplification and one for deletion, ensures that as many candidates as possible are identified.

The analysis does not determine whether an amplified or deleted gene is significantly recurrent. However, genes that are only amplified or deleted in a single cell line may be biologically relevant, as demonstrated for MEIS1, and as many different cancer types were used in the analysis, tissue-specific amplicons and deletions may not be significantly recurrent across all cell lines. A gene for which there is no evidence of a role in cancer may not be a convincing candidate if it is amplified or deleted in a single cell line, but the presence of retroviral insertions within the mouse orthologue provides further support. For all candidates, the number of amplicons or deletions containing the gene and the number of additional genes in the minimal amplified or deleted region are provided to help in assessing the contribution of a gene to tumourigenesis. In Chapter 5, efforts are made to make it easier to identify the most promising candidates by ranking genes and assigning a P-value based on the number of samples in which they are amplified or deleted. In an attempt to filter out less promising candidates, any genes that were coamplified with oncogenes or other mouse candidates were removed from the analysis, and yet co-amplified genes may co-operate in tumourigenesis (see Section 1.3.3.3). Nevertheless, given the number of candidates identified, it was considered more important to remove false positives, even at the expense of some "real" cancer genes.

As demonstrated in Section 4.5.1.4, some mouse candidates do not have human orthologues and are therefore excluded from the analysis. In some cases, the human orthologue may not have been identified, while in others, there may not be an orthologue in the human genome. However, the results of the analysis in Section 4.5.1.4 suggest that the proportion of human orthologues may be higher for "real" mouse candidates than for incorrectly assigned candidates. Any discrepancy in the number of mouse genes and the number of human orthologues does not affect the global comparison of Section 4.5.1, since the randomisation takes only mouse genes with human orthologues. This also prevents any introduction of bias resulting from the fact that only protein-coding genes have human orthologues, and that cancer genes are likely to be predominantly proteincoding. Another possible method for comparing the human and mouse data would be to map the insertion sites across to the human genome and then to assign the insertions to human genes. This could be achieved using the Ensembl Compara API, which enables the retrieval of genomic alignments between mouse and human. This would avoid the problem of lack of orthologues but there are many gaps in the alignment, which would prevent the precise mapping of a considerable proportion of insertions. To demonstrate, prior to mapping the retroviral insertions of Chapter 2 and 3 to the NCBI m36 mouse assembly, insertions were mapped to NCBI m34. Only 64.3% of insertions were successfully mapped across to the human genome (NCBI 35) using the Ensembl Compara API. A further drawback of mapping insertions could be that if there really is no human orthologue for a given mouse candidate gene, or there is a break in synteny between mouse and human, the insertions mapped to the human genome will be assigned to an incorrect gene.

The analysis is also limited by the resolution of the data. Efforts have been made to choose suitable boundaries for the ends of amplicons and deletions, but without increasing the density of the SNPs it is impossible to determine whether genes beyond the first or last amplified or deleted SNP are indeed amplified or deleted. It is also possible that small amplicons and deletions may be missed, while the high levels of noise in the data may also lead to regions of copy number change being missed or falsely identified. Encouragingly, the most successful methods for detecting changes produced similar outputs, and the fact that known and implicated oncogenes and tumour suppressor genes were identified, often in cancer types in which they have previously been shown to be disrupted, was also reassuring. However, in Chapter 5, a higher density SNP array is

used, and is compared to the 10K array to determine whether it represents a significant improvement.

Chapter 5 Identifying human cancer genes in highresolution copy number data

5.1 Introduction

Advances in comparative genomic hybridisation (CGH)-based technology have led to the development of higher resolution platforms that can identify smaller amplicons and deletions in cancers, and can more accurately define the breakpoints of regions of copy number change. Higher resolution SNP array CGH platforms have been generated by increasing the density of SNPs from across the genome that are represented on the array. This chapter describes comparative analyses between mouse candidate cancer genes identified by retroviral insertional mutagenesis and copy number data for 598 human cancer cell lines generated by the Wellcome Trust Sanger Institute (WTSI) Cancer Genome Project using high-resolution SNP array CGH. Section 5.2 describes the datasets used in these analyses, and the methods and results are described in Section 5.3. In Section 5.4, the results are compared to those obtained with the 10K CGH data described in Chapter 4 to determine whether there is a significant advantage in using the higher resolution data for integrative analyses. While the analyses in Chapter 4 involved lists of mouse candidate cancer genes that were generated by the Netherlands Cancer Institute, the analyses within this chapter involve candidates identified from the work described in Chapter 2 of this thesis, and different methods have been used to identify interesting candidates. Therefore, in order to directly compare both platforms, the analyses described in Section 5.3 are repeated using the 10K CGH data. Finally, Section 5.5 describes the identification of amplified and deleted human orthologues of mouse candidate cancer genes that co-occur with TP53 and/or CDKN2A mutations, or co-occur with one another, in the human cancer cell lines. These results are then compared to cooccurring CIS genes identified in mouse lymphomas in Section 3.5.2 in an attempt to identify cross-species conservation of co-operation between cancer genes. This chapter represents the culmination of work to characterise the mouse candidate cancer genes described in Chapters 2 and 3, and to demonstrate their relevance to human tumourigenesis. It also provides a clear illustration of how integrative data analysis can facilitate the identification of human cancer gene candidates, which can then be functionally validated in the laboratory.

5.2 Description and processing of the datasets

5.2.1 High-resolution copy number data

Copy number data were generated by the Wellcome Trust Sanger Institute Cancer Genome Project for 598 human cancer cell lines from 29 different tissues (see Table 5.1 and, for more detail, Appendix Table 4.2) using the Affymetrix Genome-Wide Human SNP Array 6.0, which comprises 1.8 million genetic markers for measuring copy number change, of which more than 906,600 are SNPs and more than 946,000 are probes for detecting copy number variation. The genetic markers were mapped to the NCBI 36 human genome assembly. The intensity values were processed into copy number ratios using the method described for the 10K data in Section 4.2.2. This is the point at which I received the data. The analysis only considers autosomes, for which the total number of markers is 1,773,325. The number of copy number markers per chromosome and the distances between adjacent markers are shown in Table 5.2 and Figure 5.1, respectively. The average distance between markers is 1.65 (\pm 41.69) kb, which compares very favourably with the mean distances of 258.50 (\pm 634.21) kb and 292.82 (\pm 683.49) kb for the two 10K arrays used in Chapter 4.

For each cell line, the R packages DNAcopy (Olshen et al., 2004), with default parameters plus removal of change-points less than 2 standard deviations (SD) apart, and MergeLevels (Willenbrock and Fridlyand, 2005) with default parameters, were used to segment each autosome into regions of uniform copy number and to merge segments that were not significantly different across the genome, respectively. DNAcopy and MergeLevels are described in Section 4.3. In this analysis, change-points less than 2 SD, rather than 3 SD, apart were removed because the comparison of methods in Section 4.6 demonstrated that reducing the number of standard deviations should result in the identification of a higher proportion of critical cancer genes. The mean number of segments per cell line was $675.33 (\pm 428.30)$, which is an average of 24.92 segments per chromosome. Given that DNAcopy and MergeLevels were originally developed for BAC array CGH and, therefore, for dealing with a considerably smaller set of copy number values than the current dataset, it is likely that the genomes are over-segmented. Decreasing the DNAcopy parameter α results in fewer change-points but requires an increased number of permutations, which is unfeasible for a dataset of this size, for which

	Number of
Site of origin	cell lines
Haematopoietic and lymphoid	103
Lung	103
Central nervous system	49
Large intestine	36
Skin	36
Breast	35
Autonomic ganglia	29
Bone	29
Stomach	19
Kidney	19
Ovary	18
Upper aerodigestive tract	18
Soft tissue	17
Oesophagus	15
Pancreas	12
Urinary tract	12
Cervix	11
Thyroid	9
Endometrium	8
Biliary tract	6
Liver	6
Pleura	5
Testis	3
Placenta	2
Prostate	2
Adrenal gland	1
Eye	1
Gastrointestinal tract	1
Small intestine	1
Vulva	1
Total	598

Table 5.1. Tissues of origin of human cancer cell lines used in high-resolution copynumber analysis.

Chromosome	Number of markers
1	145591
2	152881
3	127049
4	119457
5	115131
6	112395
7	100581
8	97736
9	81856
10	93272
11	89214
12	86990
13	65757
14	56782
15	53389
16	53920
17	46469
18	51802
19	30236
20	43457
21	24984
_22	24376
Total	1773325

Table 5.2. Number of copy number probes per human autosome.

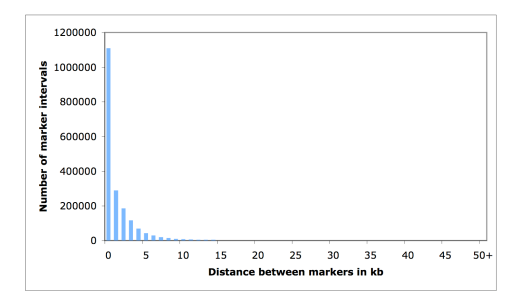


Figure 5.1. Distance between adjacent copy number probes across the human genome.

the CPU time is already very large. However, this analysis does not attempt to elucidate the events that have led to the observed changes in copy number across individual genomes, but focuses instead on individual genes that are amplified or deleted across a significant number of cell lines. Here, the use of DNAcopy and MergeLevels provides a means of filtering out anomalies at individual SNPs, rather than accurately defining entire regions of copy number change.

Since markers are closely spaced, the problems associated with defining amplicon and deletion boundaries that were described in Section 4.5.1.2 are unlikely to arise in this analysis. Therefore, boundaries were simply defined as the halfway point between the first/last amplified or deleted SNP and the preceding/proceeding SNP in the genome. None of the human cancer cell lines selected for use in this study had a shared common ancestor (see Section 4.2.2 for further details).

As in Chapter 4, the terms "gain" and "amplicon" are used interchangeably (see p.162). In the proceeding analyses, an amplification was defined as a gain of copy number greater than or equal to 1.7 and a deletion was defined as a loss of copy number less than or equal to 0.6, or less than or equal to 0.3. A threshold of 1.7 was chosen because this was the lowest copy number at which an over-representation of human orthologues of mouse candidate cancer genes from the insertional mutagenesis screen was observed (P=0.00846for genes amplified in 2 or more cell lines). Likewise, copy number 0.6 was the highest threshold at which an over-representation of orthologues in deleted regions was observed (P=0.0289 for genes deleted in 10 or more cell lines). Copy number 0.3 was the highest threshold at which an over-representation of orthologues was observed in 1 or more cell lines (P=0.00467), and these may represent homozygous deletions. The method used to generate the above *P*-values is described in Section 5.3.1. The average number of gains of copy number greater than or equal to 1.7 was 34.03 (±36.57) per cell line. The average size of these amplicons was 299.10 (±1667.93) kb and an average of 2.99 (±14.50) genes was found in each amplicon. The average number of losses of copy number less than or equal to 0.6 per cell line was 204.10 (±194.36). These losses were on average 196.87 (±3058.58) kb in size, encompassing 2.61 (±32.98) genes. Deletions were therefore smaller on average than amplicons, suggesting that deletion of gene copies may have a more detrimental effect on a cell than an increase in gene copies. Figure 5.2 shows the distribution of the number of amplicons and deletions in this collection of cell lines and the distribution of the length of aberrations.

Chapter 5

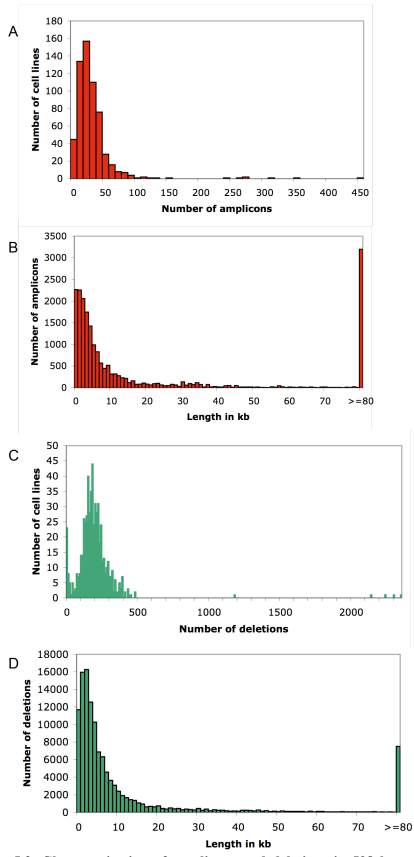


Figure 5.2. Characterisation of amplicons and deletions in 598 human cancer cell lines analysed using high-resolution SNP array CGH. (A) Number of amplicons per cell line. (B) Length of amplicons. (C) Number of deletions per cell line. (D) Length of deletions.

A 2-tailed Fisher Exact Test was used to identify types of cancer that were over- or underrepresented among cell lines containing amplicons of copy number greater than or equal to 1.7 and deletions of copy number less than or equal to 0.6. 362 cell lines contained at least one amplicon, while 542 contained at least one deletion. Cell lines derived from cancers of the oesophagus were over-represented among those containing amplicons (P=6.79x10⁻⁴) while cell lines derived from haematopoietic and lymphoid cancers were under-represented (P=3.87x10⁻³). This is consistent with the results obtained in the 10K analysis described in Section 4.4. Most cell lines contained at least one deletion, and consequently there was no significant difference between the numbers of each cancer type containing deletions.

5.2.2 Additional datasets

The dataset of copy number variants (CNVs) from Redon *et al.* (2006) is described in Section 4.2.3. However, in this chapter, 1,390 CNVs mapping to autosomes on the NCBI 36 human genome assembly, rather than NCBI 35, were used. These are available for download from http://www.sanger.ac.uk/humgen/cnv/data/cnv_data/. Known cancer genes from the Cancer Gene Census are described in Section 2.2.3. The mouse candidate cancer genes used in this chapter were the 439 genes identified in the murine leukaemia virus (MuLV) insertional mutagenesis screen described in Chapter 2. 384 of the 439 candidate cancer genes had human orthologues in Ensembl v48 (see Section 3.2.2). Mouse candidate genes are referred to here as CIS genes. Other datasets referred to in this chapter are described in Section 3.2.1.

5.3 Comparative analysis of human high-resolution CGH data versus mouse insertional mutagenesis data

5.3.1 Global comparison

The number of CIS genes with human orthologues was counted within amplicons above copy number thresholds ranging from 1.1 to 5.0 with increments of 0.1, and within deletions below copy number thresholds ranging from 0.9 to 0.1 with increments of 0.1. The number of non-CIS genes with human orthologues in amplicons and deletions was also calculated, and a 2-tailed Fisher Exact Test was used to determine whether, for each copy number threshold, the number of CIS genes was significantly different to that

expected by chance. The number of cell lines in which the genes were amplified or deleted above or below a given threshold was also counted. This global comparison is similar to that described in Section 4.5.1, except that the Fisher Exact Test, rather than the randomisation approach, was used to generate *P*-values. This gives an accurate comparison of the number of CIS genes to the exact number of non-CIS genes, rather than to an estimated number. Copy number increments of 0.1, rather than the smaller set of thresholds from Chapter 4, were used to maximise the amount of information provided by the comparison. Figures 5.3A-C show the pattern of over-represented CIS genes at varying thresholds of copy and cell line number for all cell lines, as well as for those derived specifically from haematopoietic and lymphoid tissues or from solid tumours.

For copy number thresholds of 1.7 and above, there was an over-representation of CIS genes, demonstrating that a significant proportion are amplified in human cancers and may play an important role in human tumourigenesis. The most significant result was a *P*-value of 3.23×10^{-7} for genes within 1 or more cell lines at a copy number greater than or equal to 3.4. In general, the significance increased with increasing copy number. This may reflect the fact that regions of higher copy number are more likely to be tumourigenic, and will therefore contain a higher proportion of candidate cancer genes than amplicons with a lower copy number. In addition, regions amplified to a higher copy number are likely to be more localised, containing fewer genes, and the highest peak in amplification is most likely to harbour the critical gene. The significance generally decreased with increasing numbers of cell lines. This is mainly because fewer CIS genes are amplified across larger numbers of cell lines, and thus there may not be enough power for a significant *P*-value. It may also indicate that regions that are amplified across a large number of cell lines from different types of cancer are less likely to be involved in tumourigenesis. As for the candidate genes in Section 4.5.1.3, the CIS genes were overrepresented in amplicons in haematopoietic and lymphoid cancer cell lines but with less significance than for the whole set of samples, and in cell lines derived from solid tumours. The *P*-value for genes within 1 or more solid tumour cell lines at a copy number greater than or equal to $3.4 \text{ was } 6.29 \times 10^{-7}$, which is roughly double that observed for the full set of cell lines, but is still highly significant.

An over-representation of CIS genes was also identified in deletions but with lower significance than for genes in amplicons. The most significant result was a *P*-value of 0.00467 for genes within 1 or more cell lines at a copy number less than or equal to 0.3.

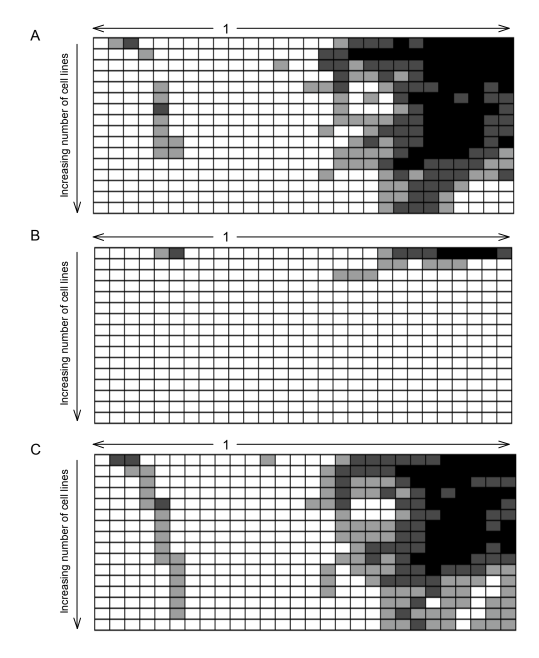


Figure 5.3. Over-representation of CIS genes in amplicons and deletions of varying copy number threshold and number of cell lines across all cell lines (A), haematopoietic and lymphoid cancer cell lines (B), and cell lines derived from solid tumours (C). Each box represents the significance of the association between CIS genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Copy number thresholds below 1 represent deletions, and range from 0.1 to 0.9 with 0.1 increments. Copy number thresholds above 1 represent amplicons, and range from 1.1 to 2.9 with 0.1 increments. The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines. For example, the box in the bottom right-hand corner represents the *P*-value for the over-representation of CIS genes that occur in amplicons of copy number greater than or equal to 2.9 in at least 16 cancer cell lines.

All other *P*-values were greater than 0.01. As mentioned in Section 4.5.1.3, it is expected that the overlap with deletions will be smaller than with amplicons because most of the CIS genes are likely to function as oncogenes. Deletions in haematopoietic and lymphoid cell lines were only slightly over-represented at copy numbers less than or equal to 0.6 and 0.5, while deletions in cell lines derived from solid tumours showed a similar pattern of over-representation to the full set of cell lines.

5.3.2 Identifying individual cancer gene candidates

5.3.2.1 Methods

The human orthologues of all mouse genes were extracted from Ensembl v48 and the number of amplicons containing each gene was calculated. Non-CIS genes were ranked according to the number of amplicons in which they resided, and a *P*-value was calculated for each CIS gene by counting the number of non-CIS genes within a higher number of amplicons and dividing it by the total number of non-CIS genes. The same procedure was used to calculate *P*-values for deleted CIS genes. These analyses were performed using the full set of cancer cell lines, as well as tissue-specific sets for cancers that were represented by 10 or more cell lines. Where genes contained regions of variable copy number within one cell line, the highest copy number was chosen to represent that gene, and the maximum copy number for that gene across all cell lines was determined. A *P*-value for the maximum peak of amplification of each CIS genes using the same method as used for the number of amplicons and deletions.

As for the 10K data in Section 4.5.2, minimal amplified and deleted regions were identified by calculating the coordinates of the smallest overlap of regions that contained the CIS gene. Other genes within the region were identified using the coordinates of human genes in Ensembl v45. For each gene within a minimal amplified region, the total number of cell lines in which that gene was amplified was calculated, and the maximum copy number across all lines was determined. A similar procedure was applied to genes within a minimal deleted region, except that the number of deletions was calculated, and the minimum copy number was determined. The minimal amplified or deleted region within which a CIS gene resides may not necessarily be a minimal amplified or deleted region from across the entire genome. The coordinates of minimal amplified and deleted

regions across the genome were therefore determined and were compared to the coordinates of CIS genes to determine whether the CIS genes resided in these regions. To avoid confusion, these regions are known as MCRs (minimal common regions). 8,694 MCRs were identified among amplicons, while 35,213 were identified among deletions of copy number less than or equal to 0.6.

The position of a CIS relative to a CIS gene was calculated using the genomic coordinates and orientation of the mouse gene, extracted from Ensembl v45, and the coordinates of the CIS in the output from the kernel convolution-based method for CIS identification (de Ridder *et al.*, 2006; see Section 2.10.2). The coordinates of the CIS are given as the position of the highest peak in insertion density within a 30 kb kernel.

Global comparisons of the number of amplicons/deletions and maximum copy number in CIS genes versus non-CIS genes were calculated using the Mann Whitney U test, wherein all the values for CIS genes were compared to all the values for non-CIS genes to determine whether the values for CIS genes tended to be greater. A 2-tailed Fisher Exact Test was used to determine whether there was any significant difference between the number of cell lines of different tissue types containing amplicons and deletions. Unless otherwise stated, all other *P*-values were generated using a 1-sided Fisher Exact Test to determine whether genes were over-represented.

5.3.2.2 Candidate cancer genes within amplicons

There were 9,681 mouse genes with human orthologues within the amplicons of human cancer cell lines. Of these, 232 were CIS genes, which is more than expected by chance (P=0.00447). 27 CIS genes were found in statistically significant recurrent amplicons (P<0.05). This included a significant over-representation of genes that were designated dominant cancer genes in the Cancer Gene Census, compared with CIS genes that were not recurrently amplified (P=2.85x10⁻⁴). 2,730 mouse genes with human orthologues were found within amplicons in haematopoietic and lymphoid cancer cell lines. CIS genes were over-represented among these genes (71 genes, P=0.0408) but, surprisingly, not as significantly as those amplified in all tumours. This further demonstrates that genes identified by insertional mutagenesis in the mouse may be relevant to the identification of cancer genes in a range of human tumours, not just in those originating in lymphoid tissue. This is at odds with the findings in Section 3.2.2, where the genes

showed no association with candidate breast and colon cancer genes from Sjöblom *et al.* (2006). However, the breast and colon candidates were identified by exon resequencing and therefore represent genes mutated by point mutations and indels in cancer. The most common effects of insertional mutagenesis, i.e. gene upregulation by promoter or enhancer insertion, more closely resemble those resulting from changes in copy number. It is therefore more likely that the human orthologues of mouse oncogenes identified by insertional mutagenesis will be disrupted by copy number changes than by small intragenic substitutions and indels. Interestingly, there was no over-representation of genes with mutations in COSMIC among CIS genes that were recurrently amplified, suggesting that copy number changes and small intragenic mutations are not positively associated within candidate cancer genes. All genes in recurrent amplicons across all cell lines, and specifically in haematopoietic and lymphoid cell lines, with a *P*-value of less than 0.1 are shown in Table 5.3.

Of the 27 statistically significant genes, there were 16 where the minimal amplified region contained only that gene. Among the remaining genes, FYN binding protein gene (FYB), myeloid cell leukaemia sequence 1 (MCL1) and acidic nuclear phosphoprotein 32 family, member E (ANP32e) co-occurred with known oncogenes, while all other genes co-occurred with genes that were amplified in more cell lines and, in some cases, to a higher maximum copy number. FYB co-occurred with the oncogene leukaemia inhibitory factor, LIFR, and all 6 of the additional genes in the minimal amplified region were amplified in a higher number of cell lines than FYB, suggesting that FYB is not the likely target for amplification in this region. Likewise, MCL1 and ANP32e co-occurred with oncogenes ALL1 fused gene from chromosome 1q (AF1O) and aryl hydrocarbon receptor nuclear translocator (ARNT). AF1Q is a fusion partner of MLL that is involved in leukaemogenesis (Tse et al., 1995), and high expression of AF1Q is associated with poor prognosis in paediatric acute leukaemia (Tse et al., 2004). It has also been shown to be overexpressed in thyroid oncocytic tumours, which are a type of tumour characterised by the presence of abundant mitochondria (Jacques et al., 2005), and in breast cancer cells, where it was associated with enhanced proliferation and metastastic potential (Chang et al., 2008; Li et al., 2006a). ARNT forms a fusion protein with ETV6 in acute myeloblastic leukaemia (Salomon-Nguyen et al., 2000) and encodes a component of the transcription factor Hypoxia-inducible factor 1 (HIF1), which is implicated in tumour growth and angiogenesis (for review, see Semenza, 2002). However, the antiapoptotic gene MCL1

		Position of CIS	Cancer							Number of		Gene in	genes in minimal	Number of oncogenes in minimal	in more	to higher
CIS gene	Mouse Ensembl ID	relative to gene	gene	COSMIC	Mullighan				p53 BS		P-value	MCR?	region		cell lines?	copy?
Wwox	ENSMUSG0000004637	Inside	-	-	-	CNV	Nanog	Oct4	-	69	0.0041	Y				
Etv6	ENSMUSG0000030199	Inside	Dominant	-	-	-	-	-	-	54	0.0043	Y				
Мус	ENSMUSG0000022346	Upstream	Dominant	-		-	Nanog	-	p53	55	0.0043	Y				
Mycn	ENSMUSG0000037169	Inside		COSMIC	Mullighan	-	Nanog	Oct4	-	25	0.0101	Y		0		
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	25	0.0101	Y				
Ccnd1	ENSMUSG0000070348	Upstream	Dominant	-	-	-	-	-	-	23	0.0111	Y				
Ikzf3	ENSMUSG0000018168	Inside	-	COSMIC	-	-	-	Oct4	-	22	0.0121	Y	-	0		
Sla	ENSMUSG0000022372	Inside	-	-	-	-	-	-	-	20	0.0132		1	0	Y	Y
Lgals9	ENSMUSG0000001123	Upstream	-	-	-			-	-	17	0.0154	Y		0		
Pml	ENSMUSG0000036986	Inside	Dominant		-	CNV	-	Oct4	-	14	0.0212	Y		-		
Itpr2	ENSMUSG0000030287	Upstream	-	COSMIC	-	-	-	-	-	14	0.0212	Y	-	0		
Fyb	ENSMUSG0000022148	Inside	-	-	-	-	Nanog	Oct4	-	14	0.0212		6	1	Y	
D12Ertd553e	ENSMUSG0000020589	Downstream	-	-	-	-	Nanog	-	-	14	0.0212	Y		0		
Slc1a3	ENSMUSG0000005360	Downstream	-	-	-	-	-	-	-	13	0.0248		9	0	Y	Y
Capsl	ENSMUSG0000039676	Downstream	-	-	-	-	-	-	-	13	0.0248		18	0	Y	Y
Cugbp2	ENSMUSG0000002107	Inside	-	-	Mullighan	-	-	Oct4	-	13	0.0248	Y		0		
Sdk1	ENSMUSG0000039683	Downstream	-	-	-	CNV	Nanog	Oct4	-	13	0.0248	Y		0		
Trp53inp1	ENSMUSG0000028211	Upstream	-	-	-	-	-	-	-	12	0.0295		6		Y	
Ptp4a3	ENSMUSG0000059895	Inside	-	-	-	-	-	-	-	10	0.0365		30		Y	Y
Mcl1	ENSMUSG0000038612	Downstream	-	-	Mullighan	-	-	-	-	10	0.0365		20		Y	
Erg	ENSMUSG0000040732	Upstream	Dominant	-	-	-	-	-	-	10	0.0365	Y	0			
1600014C10Rik	ENSMUSG0000054676	Inside	-	-	-	-	-	-	-	9	0.0447	Y	0	0		
Pag1	ENSMUSG0000027508	Upstream	-	-	-	-	-	-	-	9	0.0447		15	0	Y	Y
Anp32e	ENSMUSG0000015749	Upstream	-	-	Mullighan	-	-	-	-	9	0.0447		54	2	Y	Y
Tpd52	ENSMUSG0000027506	Upstream	-	-	-	-	Nanog	-	-	9	0.0447		15	0	Y	Y
Evi1	ENSMUSG0000027684	Inside	Dominant	COSMIC	-	-	-	-	-	9	0.0447	Y		0		
Flt3	ENSMUSG0000042817	Inside	Dominant	COSMIC	-	-	-	-	-	9	0.0447	Y	0	0		
Dock8	ENSMUSG0000052085	Upstream	-	-	-	CNV	-	-	-	8	0.0564	Y	0	0		
Ccnd3	ENSMUSG0000034165	Inside	Dominant	-	-	-	-	-	-	8	0.0564	Y	1	0		
Bcl11b	ENSMUSG0000048251	Inside	Dominant	COSMIC	-	-	-	-	-	7	0.0709	Y	0	0		
Supt3h	ENSMUSG0000038954	Upstream	-	-	-	CNV	-	-	-	7	0.0709	Y	0	0		
Cldn10a	ENSMUSG0000022132	Inside	-	-	-	-	-	-	-	7	0.0709	Y	0	0		
Rorc	ENSMUSG0000028150	Inside	-	COSMIC	Mullighan	-	-	-	-	7	0.0709		8	0	Y	Y
Zfp217	ENSMUSG0000052056	Upstream	-	COSMIC		-	Nanog	-	-	7	0.0709		4	0	Y	Y
Kit	ENSMUSG0000005672	Downstream	Dominant	COSMIC	-	-		-	-	6	0.0891	Y	0	0		
Ubac2	ENSMUSG0000041765	Inside	-	-	-	-	-	-	-	6	0.0891		8	0	Y	Y
Med13	ENSMUSG0000034297	Upstream	-	-	-	-	-	-	-	6	0.0891		16	2	Y	Y
Cd48	ENSMUSG0000015355	Upstream	-	-	Mullighan	-	-	-	-	6	0.0891	Y	0	0		
Thra	ENSMUSG0000058756	Inside	-	-	-	-	-	-	-	6	0.0891		3	0	Y	Y
Tmem49	ENSMUSG0000018171	Inside	-	-	-	-	-	-	-	6	0.0891		9	1	Y	Y
St6galnac5	ENSMUSG0000039037	Inside	-	-	-	-	-	-	-	6	0.0891	Y	ō	ō		
Myb	ENSMUSG0000019982	Downstream	-	-	Mullighan	-	-	-	-	6	0.0891	Ý	Ō	ō		
Mad1l1	ENSMUSG0000029554	Inside	-	-		-	-	Oct4	-	6	0.0891	Y	0	0		

В

		Position of CIS	Cancer							Number of		Gene in	genes in minimal	Number of oncogenes in minimal	in more	to higher
CIS gene	Mouse Ensembl ID	relative to gene	gene	COSMIC	Mullighan				p53 BS			MCR?	region		cell lines?	copy?
Wwox	ENSMUSG0000004637	Inside	-	-	-	CNV	Nanog	Oct4	-	9	0.0024	Y	0	0		
Мус	ENSMUSG0000022346	Upstream	Dominant	-	-	-	Nanog	-	p53	7	0.0025	Y	0	0		
Cugbp2	ENSMUSG0000002107	Inside	-	-	Mullighan	-	-	Oct4	-	5	0.0052	Y	0	0		
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	5	0.0052	Y	0	0		
Pag1	ENSMUSG0000027508	Upstream	-	-	-	-	-	-	-	4	0.0196		25	0		
Trp53inp1	ENSMUSG0000028211	Upstream	-	-	-	-	-	-	-	4	0.0196		20	0	Y	Y
Tpd52	ENSMUSG0000027506	Upstream	-	-	-	-	Nanog	-	-	4	0.0196		25	0		
Nedd4l	ENSMUSG0000024589	Upstream	-	-	-	CNV	Nanog	-	p53	3	0.0314		9	1		
Ptp4a3	ENSMUSG0000059895	Inside	-	-	-	-	-	-	-	3	0.0314		30	0	Y	Y
Sla	ENSMUSG0000022372	Inside	-	-	-	-	-	-	-	3	0.0314		5	0	Y	
Pml	ENSMUSG0000036986	Inside	Dominant	COSMIC	-	CNV	-	Oct4	-	3	0.0314	Y	0	0		
Bcl11b	ENSMUSG0000048251	Inside	Dominant	COSMIC	-	-	-	-	-	3	0.0314	Y	0	0		
Mcl1	ENSMUSG0000038612	Downstream	-	-	Mullighan	-	-	-	-	3	0.0314		20	2		
Rorc	ENSMUSG0000028150	Inside	-	COSMIC	Mullighan	-	-	-	-	2	0.0478		8	0	Y	
Evi1	ENSMUSG0000027684	Inside	Dominant	COSMIC	-	-	-	-	-	2	0.0478	Y	0	0		
Mbd2	ENSMUSG0000024513	Upstream	-	-	-	-	-	-	-	2	0.0478		34	2	Y	Y
Anp32e	ENSMUSG0000015749	Upstream	-	-	Mullighan	-	-	-	-	2	0.0478		63	2	Y	Y
Ikzf3	ENSMUSG0000018168	Inside	-	COSMIC	-	-	-	Oct4	-	2	0.0478	Y	0	0		
Vpreb2	ENSMUSG0000059280	Upstream	-	-	Mullighan	-	-	-	-	2	0.0478		3	0		

Table 5.3. Lists of CIS genes that are in recurrent amplicons across all cell lines (A) and across haematopoietic and lymphoid cancer cell lines only (B). Cancer gene = known cancer gene in Cancer Gene Census; COSMIC = gene contains somatic mutations in COSMIC database; Mullighan = gene within amplicon in Mullighan *et al.* (2007) dataset of acute lymphoblastic leukaemias; CNV = gene within CNV identified in Redon *et al.* (2006); "[Nanog, Oct4, p53] BS" = gene contains binding site for Nanog, Oct4 and p53, respectively. "minimal region" = minimal amplified region containing CIS gene; "MCR" = minimal amplified region from across genome, not centred on CIS gene; "Number of genes in minimal region" = number of genes other than the CIS gene within the minimal amplified region.

was shown to be amplified and overexpressed in drug-resistant cancer cell lines (Yasui et al., 2004) and in a resistant subline of BL41 Burkitt lymphoma cells (Vrana et al., 2002), and it is overexpressed in a range of leukaemias (Nagy et al., 2003). Therefore, the presence of known oncogenes within a minimal amplified region does not necessarily exclude other genes within the region as candidates for a role in tumourigenesis. As discussed in Section 1.3.3.3, genes within an amplicon may co-operate in tumour development. The protein phosphatase 2 inhibitor ANP32e is, however, a less convincing candidate, since high expression of the gene is associated with a better survival rate in patients with follicular lymphoma (Bjorck et al., 2005) and it is downregulated in a progressive, compared to a regressive, murine fibrosarcoma cancer cell line (Hayashi et al., 2005). A recurrent amplicon containing MCL1 and ANP32e was identified in the Mullighan et al. (2007) dataset of acute lymphoblastic leukaemias (ALLs; see Table 3.1) and amplification of MCL1 and ANP32e is significantly recurrent in the subset of haematopoietic and lymphoid cell lines. These observations are consistent with the theory that LIFR, AF1O, ARNT, and possibly MCL1, are the critical cancer genes in the amplicon, since all are implicated in leukaemogenesis.

A role in tumourigenesis cannot be ruled out for the 7 amplified genes that co-occurred with genes that were amplified in a greater number of cell lines. For example, the Srclike adaptor gene *Sla*, known as *SLAP* in humans, is upregulated in *FLI1*-transformed erythroblasts (Lebigot *et al.*, 2003), while protein tyrosine phosphatase type IVA, member 3 (*PTP4A3*) is involved in promoting metastasis (for review, see Bessette *et al.*, 2007).

Of the 16 genes that were identified as the only gene in the minimal amplified region, WW domain-containing oxidoreductase (*WWOX*), Aiolos (*IKZF3*) and CUG triplet repeat, RNA binding protein 2 (*CUGBP2*) might be considered more likely to play a role in tumour suppression. Their presence within amplicons therefore suggests that they reside in unstable regions of the genome. It is possible that duplication within the gene may be a mechanism for disrupting the gene, or that the copy number values across a group of markers are erroneous, such that the gene appears to be the target of amplification when in fact it is not. However, as mentioned in Section 3.4.3, *IKZF3* can both activate and repress lineage-specific cells in lymphocytes and may therefore play a dual role in tumourigenesis, and it has been found to be upregulated in chronic lymphocytic leukaemia (Duhamel *et al.*, 2008). Overexpression of *CUGBP2* induces

apoptosis of colon cancer cells exposed to radiation (Natarajan et al., 2008) by inhibiting expression of MCL1, which is described above (Subramaniam et al., 2008). All of the amplicons in the CUGBP2 gene were localised to exon 2, suggesting either an error with the 3 markers in this region, or that duplication of this exon may lead to gene disruption (Figure 5.4A). Likewise, the minimal amplified region of *IKZF3* was focused on exon 3, but many amplicons also spanned the entire gene (Figure 5.4B). The presence of amplified *WWOX* may reflect the fact that it resides in the fragile site FRA16D, which is prone to rearrangement in human cancer (see Section 4.5.2.3). Almost all of the amplicons were clustered into 2 distinct regions, one of which contained amplicons spanning up to 19 markers and the other contained amplicons spanning 2 or 3 markers. Both of these regions were within introns, suggesting that they may not affect gene expression. In addition, recurrent amplification of WWOX was observed across all cell types, and was significant in 15 of the 19 cancer types for which there were more than 10 cell lines, which suggests that it may reflect some sort of global effect rather than tumourigenicity. As expected for tumour suppressor genes, *Ikzf3*, *Cugbp2* and *Wwox* are all disrupted by intragenic CISs.

As mentioned previously, a significant number of known oncogenes were also discovered in this analysis, demonstrating that the method does successfully identify candidate cancer genes. 10 known oncogenes were also among the 35 CIS genes for which the maximum copy number was significantly higher than for non-CIS genes (P < 0.05). All genes for which the maximum copy number has a P-value of less than 0.05 are shown in Table 5.4. After accounting for known oncogenes, the remaining candidates that were recurrently amplified with a P-value of <0.05 included ITPR2, SDK1, NP 001035167.2 and FAM49A (known as D12Ertd553e in the mouse), all of which were the only gene in the minimal amplified region. FAM49A co-occurred with MYCN in some cell lines, but since this gene was also mutated by insertional mutagenesis, a role in tumourigenesis for the amplified gene cannot be ruled out, and it is within the boundaries of a recurrent amplicon, also containing MYCN, that was identified in ALLs by Mullighan et al. (2007). *ITPR2* encodes an inositol 1,4,5-trisphosphate receptor that plays an essential role in calcium signalling. Although a role in tumourigenesis has not been confirmed, *ITPR2* is co-amplified with KRAS2 and KRAG in a range of human tumours (Heighway et al., 1996). In the current study, significant recurrent amplification of ITPR2 was observed in cell lines derived from cancers of the pancreas, colon, ovary, cervix and endometrium. KRAS2 has been implicated in the development of all of these cancer types. However, the

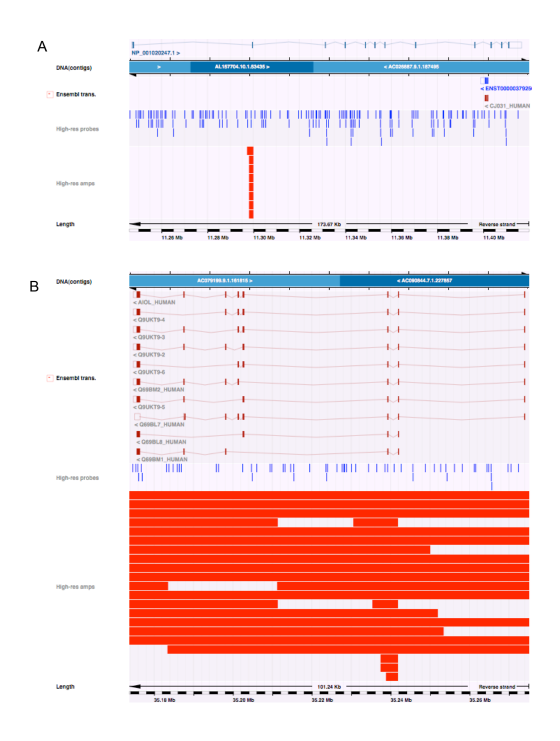


Figure 5.4. The minimal amplified regions within putative tumour suppressor genes *CUGBP2* (A) and *IKZF3* (B) are localised around specific exons. The Ensembl transcripts for *CUGBP2* and *IKZF3* are shown in blue and red, respectively. Copy number markers are shown as blue vertical lines (labelled High-res probes) and amplicons are shown as red rectangles (labelled High-res amps).

		Cancer	Number of	Maximum	
CIS gene	Mouse Ensembl ID	gene	amplicons	copy number	P-value
Tqfbr3	ENSMUSG0000029287		5	30.23	0.0030
Fli1	ENSMUSG0000016087	Dominant	1	24.70	0.0054
Mad1l1	ENSMUSG0000029554		6	22.19	0.0069
Cldn10a	ENSMUSG0000022132		7	21.05	0.0091
Ppp1r16b	ENSMUSG0000037754		2	21.05	0.0091
Ptpre	ENSMUSG0000041836		1	14.73	0.0154
A2AN91_MOUSE	ENSMUSG0000038578		1	14.37	0.0161
Mgat4a	ENSMUSG0000026110		2	14.29	0.0161
Stard3nl	ENSMUSG0000003062		3	13.59	0.0169
Anxa2	ENSMUSG0000032231		1	12.12	0.0190
Dym	ENSMUSG0000035765		4	10.95	0.0216
C330024D12Rik	ENSMUSG0000030553		1	10.95	0.0216
Ccnd3	ENSMUSG0000034165	Dominant	8	10.95	0.0216
Ikzf3	ENSMUSG0000018168		22	8.01	0.0281
Dock8	ENSMUSG0000052085		8	7.42	0.0295
Fgfr2	ENSMUSG0000030849	Dominant	5	6.82	0.0308
4932417H02Rik	ENSMUSG00000025583		2	6.76	0.0308
Мус	ENSMUSG0000022346	Dominant	55	6.66	0.0309
Etv6	ENSMUSG0000030199	Dominant	54	6.60	0.0310
Cugbp2	ENSMUSG0000002107		13	6.28	0.0335
Fgd2	ENSMUSG0000024013		3	6.28	0.0335
Mycn	ENSMUSG0000037169	Dominant	25	6.24	0.0338
D12Ertd553e	ENSMUSG0000020589		14	6.23	0.0339
Rara	ENSMUSG0000037992	Dominant	5	5.88	0.0351
Flt3	ENSMUSG0000042817	Dominant	9	5.80	0.0352
Evi1	ENSMUSG0000027684	Dominant	9	5.37	0.0367
Erg	ENSMUSG0000040732	Dominant	10	5.24	0.0369
1110036003Rik	ENSMUSG0000006931		3	5.10	0.0401
Med13	ENSMUSG0000034297		6	4.91	0.0408
Rcsd1	ENSMUSG0000040723		3	4.80	0.0414
Slamf6	ENSMUSG0000015314		5	4.70	0.0438
Recql5	ENSMUSG0000020752		4	4.62	0.0449
Thra	ENSMUSG0000058756		6	4.62	0.0449
Sla	ENSMUSG0000022372		20	4.58	0.0455
Cyb5	ENSMUSG0000024646		2	4.41	0.0475

Table 5.4. A list of CIS genes for which the maximum copy number across all cell lines is significantly higher than expected by chance.

identification of insertions within *ltpr2*, and a previous study showing that *ltpr2* is targeted by Hepatitis B virus insertional mutagenesis in hepatocellular carcinomas (Paterlini-Brechot et al., 2003), suggests that amplification of ITPR2 may also contribute to cancer development. In addition, 5 out of 136 human cancer samples tested have a somatic misssense mutation in ITPR2 in the COSMIC database (Forbes et al., 2006). SDK1, or sphingosine-dependent protein kinase 1, has the same amino acid sequence as the kinase domain of PKC8 (Hamaguchi et al., 2003) and specifically phosphorylates certain isoforms of 14-3-3 that regulate signal transduction and have been implicated as potential oncogenes (Megidish et al., 1998; for review on 14-3-3 proteins, see Tzivion et al., 2006). However, activation of SDK1 leads to apoptosis (Suzuki et al., 2004), suggesting that it has a tumour suppressive, rather than an oncogenic, role in cancer. 5 of the amplicons overlapping SDK1 were very long, spanning a region of at least 4.4 Mb. The remaining amplicons were small and occurred in intronic regions, suggesting that they may not disrupt the gene. LGALS9 (galectin-9) appears to play dual roles in cancer, since it is associated with antimetastatic potential in breast cancer and oral squamous cell carcinoma cell lines (Irie et al., 2005; Kasamatsu et al., 2005), but it also stimulates phosphorylation of Tim-3, which is implicated in the survival of melanoma cells (Wiener et al., 2007). The amplicons were identified in gene NP 001035167.2, which is a paralogue of LGALS9. A CNV locus from Redon et al. (2006) spans the entire NP 001035167.2 gene and includes 61 gains in copy number and 5 losses from 270 HapMap individuals. The amplicons in NP 001035167.2 roughly overlap with the CNV (Figure 5.5). Therefore, while mouse *Lgals9*, and potentially its human orthologue LGALS9, may contribute to tumourigenesis, the human paralogue NP 001035167.2 may not, since it is commonly amplified or deleted in normal individuals as well as in human cancer cell lines. Incidentally, CIS genes are under-represented in CNVs (P=0.0217), which provides support for the fact that they contribute to cancer and are therefore unlikely to be disrupted in healthy individuals.

Of the 71 CIS genes in haematopoietic and lymphoid tumours, 19 were found in statistically recurrent amplicons (P < 0.05). 5 of the genes with a P-value of less than 0.05 (i.e. *NEDD4L*, *BCL11B*, *RORC*, *MBD2* and *VPREB2*) were not significantly amplified in the full set of tumours, suggesting that they are specifically associated with cancers of haematopoietic and lymphoid tissue. However, *NEDD4L* and *MBD2* both co-occurred with known oncogene mucosa associated lymphoid tissue lymphoma translocation gene 1 (*MALT1*), while *MBD2* also co-occurred with the B-cell leukaemia/lymphoma 2 gene

Chapter 5



Figure 5.5. Amplicons and deletions in the *LGALS9* paralogue *NP_001035167.2* overlap with a copy number variant (CNV) from Redon *et al.* (2006). The CNV and amplicons and deletions are shown as black, red and green rectangles, respectively. All amplicons, but only around half of all deletions, are shown. Copy number markers are shown in blue.

BCL2. Both *MALT1* and *BCL2* are implicated in lymphomagenesis. The retinoic acid receptor-related orphan receptor C, *RORC*, and the immunoglobulin omega chain precursor, *VPREB2*, have not been previously implicated in cancer, but mutations in *Rorc* were associated with abnormalities in the development of lymphoid organs in immunodeficient mice (Seymour *et al.*, 2006), while *VPREB2* is selectively expressed in pre-B lymphocytes (Kudo and Melchers, 1987; Okabe *et al.*, 1992) and contributes to B-cell development (Dul *et al.*, 1996; Mundt *et al.*, 2001; Shimizu *et al.*, 2002). *RORC* was identified in the recurrent ALL amplicon of Mullighan *et al.* (2007) that also contained *ANP32e* and *MCL1*, while *VPREB2* resides in a recurrent ALL amplicon that is telomeric of *BCR*. However, in this analysis, neither resided within an MCR in haematopoietic and lymphoid cell lines. B-cell leukaemia/lymphoma 11B gene *BCL11B* does reside in an MCR, and is the only gene within it. *BCL11B* encodes a Kruppel-like zinc finger protein that is involved in thymopoiesis and is required for the survival of human T-cell leukaemia and lymphoma cell lines, suggesting an antiapoptotic role in these cancers (Grabarczyk *et al.*, 2007).

Interestingly, 9 of the 27 genes significantly amplified across all cancer types were not amplified in any haematopoietic and lymphoid cell lines, while a further 4 were amplified but not significantly. This may seem surprising since the genes play a role in lymphomagenesis in the mouse, and therefore might be expected to do the same in the However, it is possible that they are simply not disrupted by human disease. amplification in haematopoietic and lymphoid cell lines, which tend to show lower levels of copy number change than cell lines derived from solid tumours. Amplification of ets variant gene 6 (ETV6) was significantly under-represented in haematopoietic and lymphoid lines compared with other cancer types ($P=3.59 \times 10^{-5}$). ETV6 contributes to human leukaemia through the formation of gene fusions that are not associated with an increase in copy number. There is no evidence to suggest that overexpression of ETV6 is implicated in tumourigenesis. It resides on, but is not believed to be a critical gene in, an amplicon that is frequently found in breast tumours and osteosarcomas (Gisselsson et al., 2002; Yao et al., 2006). 41 identical amplicons spanning 2 intronic markers were identified, suggesting that the copy number at these markers may be erroneous. MYCN is also under-represented in haematopoietic and lymphoid cell lines and is over-represented in cell lines derived from tumours of the autonomic ganglia ($P=9.12 \times 10^{-24}$). This is consistent with the role of amplified MYCN in the development of neuroblastomas. It is possible that genes that are activated by insertional mutagenesis, and contribute to

lymphomagenesis, in the mouse might not contribute to the formation of spontaneous mouse or human lymphomas because they are not normally expressed in lymphocytes. However, in the case of *MYCN*, activation by translocation has been demonstrated in non-Hodgkin's lymphoma (Finnegan *et al.*, 1995), and *MYCN* appears to co-operate with the ETS family gene *TEL2* in B-cell lymphomagenesis (Cardone *et al.*, 2005). As mentioned previously, it is also found within a recurrent amplicon in ALL (Mullighan *et al.*, 2007).

Other genes that were significantly over-represented in a particular tissue type were CCND1 (oesophagus, $P=6.51 \times 10^{-6}$), TMEM49 (breast, $P=1.37 \times 10^{-4}$), NCOA3 (breast, $P=1.85 \times 10^{-4}$) and RCBTB2 (large intestine, $P=2.01 \times 10^{-4}$). Amplification and overexpression of CCND1, or Cyclin D1, has been demonstrated in 32% of human oesophageal squamous cell carcinomas (Jiang et al., 1993) and 64% of oesophageal adenocarcinomas (Arber et al., 1996), while overexpression of nuclear receptor coactivator 3, NCOA3, is associated with poor prognosis in breast tumours (Zhao et al., 2003). All 3 amplicons containing NCOA3 were in breast cancer cell lines. This corresponded to a significant recurrence across breast cancers (P=0.0259) but not across all cell lines (P=0.249), suggesting that NCOA3 may contribute to tumourigenesis, but only in the breast. However, of the 3 genes that co-occurred with NCOA3, 2 (PRKCBP1 and EYA2) were amplified to a higher copy number, and Prkcbp1 is in fact a CIS gene. 6 cell lines contained a TMEM49 amplification, of which 4 were derived from breast tumours. Again, this corresponded to a significant recurrence across breast cancers (P=0.0117) but not across all cell lines (P=0.0891). TMEM49 has not been previously implicated in cancer but it falls within a common region of amplification in breast cancers on chromosome 17q23 that is associated with poor prognosis. Other genes in the minimal amplified region, including PPM1D, APPBP2, RPS6KB1 and BCAS3, have been implicated in breast cancer development (for review, see Sinclair et al., 2003) and of these, *RPS6KB1* and *BCAS3* were amplified to a higher copy number than *TMEM49* in other cell lines. However, the identification of insertions within *Tmem49* suggests that this gene may also be important in tumourigenesis. All 3 amplicons containing RCBTB2 were within colon cancer cell lines, which corresponded to a significant recurrence across colon cancers (P=0.0107) but not across all cell lines (P=0.249). RCBTB2 co-occurs with known tumour suppressor gene RB1, and is in fact a candidate tumour suppressor gene in a region on chromosome 13q14 that frequently shows loss of heterozygosity in prostate cancer (Latil et al., 2002). The insertions assigned to Rcbtb2 are upstream in the sense and antisense orientation, suggesting that the gene plays an oncogenic role, but all

insertions are within a longer Ensembl EST gene transcript that is not annotated as an Ensembl gene transcript and could therefore be intragenic, inactivating insertions (Figure 5.6, page 249). *RCBTB2* is not within an MCR, suggesting that it may not contribute to colon tumourigenesis.

Other genes were also amplified in a significant number of cell lines of a particular tissue type but not across all cell lines. Among these were genes encoding suppressor of Ty 3 homolog (SUPT3H), claudin-10 (CLDN10) and MAD1 mitotic arrest deficient-like 1 (MAD1L1), which were significantly amplified in soft tissue (P=0.00566), colon (P=0.0107) and lung (P=0.0267) cancer cell lines, respectively. All 3 genes were within MCRs. SUPT3H is a transcription factor that forms part of a multiprotein complex that mediates transcriptional activation (Brand et al., 1999). Importantly, it is required for transcription and cell proliferation induced by the MYC oncoprotein (Liu et al., 2008). Claudins are components of tight junctions and have been implicated in tumour progression (Kominsky, 2006). *CLDN10* is overexpressed in papillary thyroid carcinoma (Aldred et al., 2004) and overexpression in hepatocellular carcinoma cells promotes cancer cell survival, motility and invasiveness, leading to malignancy (Ip et al., 2007). Claudin-1 plays an important role in cellular transformation and metastasis in colon cancer (Dhawan et al., 2005; Resnick et al., 2005), and overexpression of claudin-7 and claudin-12 is also implicated (Darido et al., 2008; Grone et al., 2007). These findings suggest that amplification of claudin-10 may also contribute to colon cancer, which has not been previously shown. MAD1L1 is a mitotic checkpoint gene that has also been shown to harbour somatic missense mutations in lung cancer cell lines (Nomoto et al., 1999) and has been presented as a putative tumour suppressor gene (Tsukasaki et al., 2001). However, consistent with the results of the analysis described herein, a region on chromosome 7p22.3 that is centred on *MAD1L1* was found to be the most frequently observed copy number change in small-cell lung cancer cell lines (Coe et al., 2006), suggesting that it may play an oncogenic role that requires further investigation. The maximum copy numbers of *CLDN10* and *MAD1L1* were also significantly higher than for non-CIS genes (P < 0.05, see Table 5.4).

The number of recurrent amplifications was significantly higher across CIS genes than across non-CIS genes (P=0.00396). The median number of amplifications was 1 for both samples, and the maximum number was larger for non-CIS genes, at 115 recurrent amplifications, compared with 69 for CIS genes (specifically *MYC*). However, the mean

number was 2.77 for CIS genes compared with 2.30 for non-CIS genes. The maximum peak of amplification was also higher among CIS genes (P=0.00135). The minimum and maximum copy numbers were 1.253 (*UBE1L*) and 30.230 (*TGFBR3*) for CIS genes, and 1.227 and 67.960 for non-CIS genes. The median and mean peaks in amplification were 1.832 and 2.692 for CIS genes, and 1.721 and 2.405 for non-CIS genes. These results further suggest that a significant proportion of CIS genes may contribute to human tumourigenesis through the mechanism of amplification.

5.3.2.3 Candidate cancer genes within deletions

16,973 human genes with mouse orthologues, including all but one of the CIS genes, were identified in deletions of copy number less than or equal to 0.6 in human tumours. However, only 24 CIS genes were found in a significant number of deletions (P<0.05). Table 5.5A shows all CIS genes with a P-value of less than 0.1. Unlike those in amplicons, CIS genes in deletions were not significantly over-represented among known oncogenes (P=0.177) compared with other CIS genes. There was also no over-representation of CIS genes with mutations in COSMIC (P=0.951).

5 genes (CCND2, ETV6, LGALS9, SDK1 and WWOX) were both significantly amplified and significantly deleted. This is a larger overlap than expected by chance (P=0.0196). As discussed in the previous section, deletions within ETV6, WWOX and SDK1 are predicted to contribute to tumourigenesis, while the LGALS9 paralogue NP001035167.2 shows both gains and losses in copy number in the normal population. The deletions in NP001035167.2 (Figure 5.5) and ETV6 were in exactly the same location as the corresponding amplicons, further indicating that the copy number values may be erroneous or due to CNVs. This is also somewhat true of the deletions in WWOX and SDK1, but WWOX and, to a lesser extent, SDK1 also contained many deletions that spanned other regions, or the entire gene. However, only 2 MuLV insertions were found within Sdk1, and the coordinates of the CIS are 161.05 kb downstream of the gene, therefore shedding doubt on a tumour suppressive role for the gene in mouse lymphomas. CCND2 is a known oncogene that is amplified and overexpressed in a range of human cancers, including malignant gliomas (Buschges et al., 1999), B-cell neoplasms (Werner et al., 1997) and testicular germ cell tumours (Rodriguez et al., 2003). However, its proximity to $p27^{KIP1}$ on the short arm of chromosome 12 means that it is frequently

															Other	
													Number of	Number of	genes	
													genes in	TSGs in		Other genes
		Position of CIS	Cancer							Number of		Gene in	minimal	minimal	more cell	
CIS gene	Mouse Ensembl ID	relative to gene	gene	COSMIC	Mullighar				5 p53 BS		P-value	MCR?	region	region	lines?	lower copy?
Wwox	ENSMUSG0000004637	Inside	-	-	-		Nanog	Oct4	-	237	0.0000	Y	0	0		
Lgals9	ENSMUSG0000001123	Upstream	-	-	Mullighan	CNV	-	-	-	146		Y	0	0		
Etv6	ENSMUSG0000030199	Inside	Dominant	-	Mullighan	-	-	-	-	120	0.0049	Y	0	0		
Sdk1	ENSMUSG0000039683	Downstream	-	-	Mullighan	CNV	Nanog	Oct4	-	83	0.0084	Y	0	0		
Metrnl	ENSMUSG0000039208	Upstream	-	-	-	-	-	-	-	72		Y	0	0		
Zfp438	ENSMUSG0000050945	Upstream	-	-	-	-	-	-	-	67	0.0125	Y		0		
Midn	ENSMUSG0000035621	Inside	-	-	Mullighan			-	-	66	0.0136	Y		0	Y	
Arid3a	ENSMUSG0000019564	Inside	-	-	Mullighan	CNV	-	-	-	65	0.0151		6	0	Y	
Ptbp1	ENSMUSG0000006498	Upstream	-	-	Mullighan	-	-	-	-	65	0.0151		8	0	Y	
Mknk2	ENSMUSG0000020190	Upstream	-	-	-	-	-	-	-	56	0.0197		8	0	Y	
Mobkl2a	ENSMUSG0000003348	Downstream	-	-	-	-	-	-	-	56	0.0197		2	0	Y	
Ets1	ENSMUSG0000032035	Upstream/Downstream	-	COSMIC	-	-	-	-	-	55	0.0202	Y		0		
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	53	0.0219	Y		0		
Acot11	ENSMUSG0000034853	Upstream	-	-	-	-	-	-	-	52	0.0222	Y		0		
Gadd45b	ENSMUSG0000015312	Inside	-	-	-	-	-	-	-	52			15	0	Y	
Dym	ENSMUSG0000035765	Inside	-	-	-	-	-	-	-	50	0.0240	Y		0		
Notch1	ENSMUSG0000026923	Inside	Dominant	COSMIC	-	CNV	-	-	p53	50	0.0240	Y		0		
Gna15	ENSMUSG0000034792	Inside	-	-	-	-	-	-	-	47	0.0281		10	0	Y	
Tbxa2r	ENSMUSG0000034881	Upstream	-	-	-	-	-	-	-	46	0.0293		22	0	Y	
Nedd4l	ENSMUSG0000024589	Upstream	-	-	-	CNV	Nanog	-	p53	43	0.0323	Y		0		
Cyb5	ENSMUSG0000024646	Downstream	-	-	-	-	Nanog	-	-	42	0.0351	Y		0		
Dock8	ENSMUSG0000052085	Upstream	-	-	Mullighan	CNV	-	-	-	42		Y		0		
Ntn1	ENSMUSG0000020902	Downstream	-	-	Mullighan	-	Nanog	-	-	42	0.0351	Y		0		
Cbfa2t3	ENSMUSG0000006362	Upstream	Dominant	-	-	-	-	-	-	39	0.0455	Y		0		
Rtn4rl1	ENSMUSG0000045287	Downstream	-	-	Mullighan	-	-	-	-	37	0.0502		28	1	Y	Y
Slc43a2	ENSMUSG0000038178	Inside	-	-	Mullighan	-	-	-	-	37	0.0502		12	0	Y	Y
Smg6	ENSMUSG0000038290	Inside	-	COSMIC	Mullighan	-	-	-	-	37	0.0502		4	0	Y	
Ovca2	ENSMUSG0000038268	Downstream	-	-	Mullighan	-	-	-	-	36	0.0538		28	1	Y	Y
Ski	ENSMUSG0000029050	Upstream	-	-	-	-	-	-	-	36	0.0538	Y		0		
Tcf25	ENSMUSG0000001472	Downstream	-	-	-	CNV	-	-	-	36	0.0538		3	0	Y	
Ttll10	ENSMUSG0000029074	Upstream	-	-	-	-	-	-	-	36		Y	27	0	Y	
Vps13d	ENSMUSG0000020220	Inside	-	-	-	-	-	-	-	36		Y	0	0		
Arrdc5	ENSMUSG0000073380	Inside	-	-	-	-	-	-	-	35	0.0569		27	0	Y	Y
Rnf166	ENSMUSG0000014470	Inside	-	-	-	-	-	-	-	35	0.0569		18	0	Y	
Mbd2	ENSMUSG0000024513	Upstream	-	-	-	-	-	-	-	34	0.0601	Y	0	0		
Pik3cd	ENSMUSG0000039936	Upstream	-	-	-	-	-	Oct4	-	34	0.0601	Y		0	Y	
Prdm16	ENSMUSG0000039410	Inside	Dominant		-	CNV	-	-	-	33	0.0637	Y	0	0		
Foxp1	ENSMUSG0000030067	Inside	-	COSMIC	-	-	Nanog	-	-	32	0.0667	Y	0	0		
BC008155	ENSMUSG0000057411	Inside	-	-	-	CNV	-	-	-	29	0.0772	Y	4	0	Y	
1700081D17Rik	ENSMUSG0000022085	Downstream	-	-	-	CNV	-	-	-	28	0.0824	Y		0		
Kit	ENSMUSG0000005672	Downstream	Dominant	COSMIC	-	-	-	-	-	28	0.0824	Y	0	0		
Erg	ENSMUSG0000040732	Upstream	Dominant	-	Mullighan	-	-	-	-	26	0.0927	Y	0	0		
Park7	ENSMUSG0000028964	Upstream	-	-	-	-	-	-	-	25	0.0986		2	0	Y	

В

		Position of CIS	Cancer							Number of		Gene in	Number of genes in minimal	TSGs in minimal	more cell	
CIS gene	Mouse Ensembl ID	relative to gene	gene		Mullighan				p53 BS		P-value	MCR?	region	region	lines?	lower copy?
Wwox	ENSMUSG0000004637	Inside	-	-	-		Nanog	Oct4	-	41	0.0006	Y	0	0		
Lgals9	ENSMUSG0000001123	Upstream	-	-		CNV		1	-	22	0.0045	Y	0	0		
Sdk1	ENSMUSG0000039683	Downstream	-	-	Mullighan	CNV	Nanog	Oct4	-	19	0.0088	Y	0	0		
Etv6	ENSMUSG0000030199	Inside	Dominant	-	Mullighan	-	-	-	-	18	0.0094	Y	0	0		
Ntn1	ENSMUSG0000020902	Downstream	-	-	Mullighan	-	Nanog	-	-	10	0.0227	Y	0	0		
Acot11	ENSMUSG0000034853	Upstream	-	-	-	-	-	-	-	9	0.0259	Y	0	0		
Zfp438	ENSMUSG0000050945	Upstream	-	-	-	-	-	-	-	7	0.0343	Y		0		
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	7	0.0343	Y	0	0		
Kit	ENSMUSG0000005672	Downstream	Dominant	COSMIC	-	-	-	-	-	7	0.0343	Y	0	0		
Smg6	ENSMUSG0000038290	Inside	-	COSMIC	Mullighan	-	-	-	-	6	0.0518		36	1	Y	Y
Ovca2	ENSMUSG0000038268	Downstream	-	-	Mullighan	-	-	-	-	6	0.0518		36	1	Y	Y
SIc43a2	ENSMUSG0000038178	Inside	-	-	Mullighan	-	-	-	-	6	0.0518		36	1	Y	Y
Rtn4rl1	ENSMUSG0000045287	Downstream	-	-	Mullighan	-	-	-	-	6	0.0518		36	1	Y	Y
Foxp1	ENSMUSG0000030067	Inside	-	COSMIC		-	Nanog	-	-	6	0.0518	Y	0	0		
Tbc1d1	ENSMUSG0000029174	Downstream	-	-	-	-	Nanog	-	-	6	0.0518	Y	0	0		
Notch1	ENSMUSG0000026923	Inside	Dominant	COSMIC	-	CNV		-	p53	6	0.0518	Y	0	0		
Sema4d	ENSMUSG0000021451	Inside	-	-	-	-	-	-	-	5	0.0609	Y	Ó	Ó		
Pml	ENSMUSG0000036986	Inside	Dominant	COSMIC	-	CNV	-	Oct4	-	5	0.0609	Y	0	0		
Rcbtb2	ENSMUSG0000022106	Upstream	-	COSMIC	Mullighan	-	-	-	-	5	0.0609	Y	Ó	Ó		
Fut8	ENSMUSG0000021065	Upstream	-	-		-	-	-	-	5	0.0609	Y	0	0		
Lrrfip1	ENSMUSG0000026305	Inside	-	-	Mullighan	-	Nanog	-	-	4	0.0840	Ý	1	ō		
Pik3r5	ENSMUSG0000020901	Upstream	-	-	Mullighan	-		-	-	4	0.0840		10	0	Y	Y
2310016C08Rik	ENSMUSG0000043421	Downstream	-	-	-	-	-	-	-	4	0.0840		43	ō	Ý	Ŷ
Gse1	ENSMUSG0000031822	Upstream/Inside	-	COSMIC	-	-	-	-	-	4	0.0840	Y	0	0		
Tspan14	ENSMUSG0000037824	Inside	-	-	-	-	-	-	-	4	0.0840	Ý	ő	ő		
Ets1	ENSMUSG0000032035	Upstream/Downstream	-	COSMIC	-	-	-	-	-	4	0.0840	Ý	0	0		
Dock8	ENSMUSG00000052085	Upstream	-	-	Mullighan	CNIV		-			0.0840	Ý	ő	ő		
Vps13d	ENSMUSG00000020220	Inside	_	_	-	-		_			0.0840	Ý	0	0		
Gadd45a	ENSMUSG0000020220	Upstream	-		-	-	-	- Oct4		4	0.0840	'	47	0	Y	
Cauu45g Ccrk	ENSMUSG00000021455 ENSMUSG00000021483	Upstream	-	-	-	-	-	0014	-	4	0.0840		47	0	Y	
Metrnl	ENSMUSG00000021485 ENSMUSG00000039208	Upstream	-	-	-	-	-	-	-	4	0.0840	Y	47	0	T	
Mad1l1	ENSMUSG00000039208	Inside	-	-	- Mullighan	2	-	- Oct4	-	4	0.0840	Y	0	0		
riautit	EN31903G00000029554	mside	-	-	munighan	-	-	ULL4	-	4	0.0840	I	U	0		

Table 5.5. A list of CIS genes that are in recurrent deletions of copy number less than or equal to 0.6 across all cell lines (A) and across haematopoietic and lymphoid cancer cell lines (B). Cancer gene = known cancer gene in Cancer Gene Census; COSMIC = gene contains somatic mutations in COSMIC database; Mullighan = gene within deletion in Mullighan et al. (2007) dataset of acute lymphoblastic leukaemias; CNV = gene within CNV identified in Redon *et al.* (2006); "[Nanog, Oct4, p53] BS" = gene contains binding site for Nanog, Oct4 and p53, respectively. "minimal region" = minimal deleted region containing CIS gene; "MCR" = minimal deleted region from across genome, not centred on CIS gene; "Number of genes in minimal region" = number of genes other than the CIS gene within the minimal deleted region.

deleted along with $p27^{KIP1}$ in childhood acute lymphoblastic leukaemia (Komuro *et al.*, 1999). Only 2 of the deletions spanning *CCND2* were long enough to include $p27^{KIP1}$. Most were very small (~500 bp) and 18 co-occurred in the same region as 23 focal amplicons, suggesting that they may represent errors in copy number measurement.

For a number of the remaining genes that were recurrently deleted, there is supporting evidence in the literature suggesting that they are tumour suppressor genes. These include *OVCA2* and *MOBKL2A*, which were discussed in Section 3.4.3, plus *DOCK8*, which is deleted and under-expressed in human lung cancers (Takahashi *et al.*, 2006), and *CBFA2T3*, which is a putative breast tumour suppressor gene (Kochetkova *et al.*, 2002; Powell *et al.*, 2002). Interestingly, the human orthologue of *Smg6*, which was proposed to be a putative mouse tumour suppressor gene in Section 3.4.3, was also recurrently deleted, although *SMG6* was located in a minimal deleted region that included 4 other genes that were deleted in a greater number of cell lines. Of these candidates, only *Smg6* has a CIS within the gene. However, all of the genes contain insertions and, especially in the cases of *Ovca2* and *Mobkl2a*, which contain 9 and 6 insertions respectively, it appears that 2 nearby CISs may have been merged, resulting in a CIS location that does not reflect the true location of either CIS. This is one of the limitations of the kernel convolution-based method for identifying CISs, since CISs vary in size and the chosen kernel width may not be appropriate for all CISs.

Although *METRNL* and *ZNF438* (or *Zfp438* in the mouse) were frequently deleted and *ZNF438* has been shown to act as a transcriptional repressor (Zhong *et al.*, 2007), the distribution of insertions around the mouse genes suggests that they are unlikely to act as tumour suppressor genes, at least not in MuLV-induced lymphomagenesis. More promising candidate tumour suppressor genes include cytochrome b5 (*CYB5*), which is frequently deleted in uterine leiomyosarcoma (Cho *et al.*, 2005), and netrin-1 (*NTN1*), the expression of which is reduced in prostate tumours (Latil *et al.*, 2003). Loss of function of dymeclin (*DYM*) is implicated in the rare autosomal recessive Dyggve-Melchior Clausen syndrome (El Ghouzzi *et al.*, 2003), which is associated with mental retardation. No role in tumourigenesis has previously been observed, but the deletion of *DYM* and the presence of intragenic insertions within *Dym* implicates the gene as a potential tumour suppressor. Most of the deletions in *CYB5*, *NTN1* and *DYM* span the entire gene. Once again, only *Dym* contains an internal CIS, but both *Cyb5* and *Ntn1* contain intragenic insertions, respectively).

The significantly deleted genes also included known and implicated oncogenes *NOTCH1* and *ETS1*. In the case of *NOTCH1*, the minimal deleted region was within intron 2, where deletions that result in the formation of an N-terminally truncated oncoprotein are commonly observed in cancer (see Section 3.4.2). While some of the deletions spanned the entire gene, many left the last 3-8 exons intact, which again may result in the production of the N-terminally truncated, intracellular oncogenic NOTCH-IC protein (Figure 5.7). *ETS1* is a member of the ets protein family, which also includes *ERG* and *ETV6*. The distribution of MuLV insertions in *Erg* and *Etv6*, which are described in Section 3.4.2, suggests that truncation and/or deletion of these genes is implicated in tumourigenesis. A search in the COSMIC database revealed that a nonsense mutation (replacing arginine at reside 211) that would result in the removal of the DNA-binding Ets domain while still retaining the SAM_PNT domain has been observed in *ETS1* spans the final 2 exons of the gene and would result in a similar protein product containing the SAM PNT domain but no Ets domain (Figure 5.8).

A number of additional candidates were identified among genes that showed significantly recurrent deletion at copy numbers less than or equal to 0.3. All CIS genes occurring within recurrent deletions with a *P*-value of less than 0.1 are shown in Table 5.6A. These included vacuolar protein sorting 13D (VPS13D), juxtaposed with another zinc finger protein 1 (JAZF1), regulator of chromosome condensation and BTB domain containing protein 2 (*RCBTB2*) and phosphatidylinositol-4,5-bisphosphate 3-kinase catalytic subunit delta isoform (*PIK3CD*). *VPS13D* is poorly characterised and is not known to play a role in cancer, while JAZF1 forms fusions with JJAZ1 and PHF1 in endometrial stromal tumours (Koontz et al., 2001; Micci et al., 2006) and deletion within JAZF1 has not been implicated in tumourigenesis. In both genes, there is an intronic region of deletion that is exactly the same across each cell line, suggesting that it may not contribute to cancer. In the previous section, *RCBTB2* was shown to be significantly amplified but evidence in the literature suggested that it was a tumour suppressor gene. The identification of recurrent deletions within RCBTB2 lends further support to this theory. PIK3CD is overexpressed in, and contributes to the survival and proliferation of, blast cells in patients with acute myeloid leukaemia (Sujobert et al., 2005), implicating it as an oncogene. However, PIK3CD also resides within a region on chromosome 1p36 that is frequently deleted in neuroblastomas and was identified as the most interesting candidate for further study (Caren et al., 2007). Importantly, deletions with a copy number of 0.6 or



Figure 5.6. All of the MuLV insertions assigned to the *Rcbtb2* gene are within a larger, unannotated, EST transcript. Insertions are shown as black vertical lines. The Ensembl gene and EST are shown in red and purple, respectively.

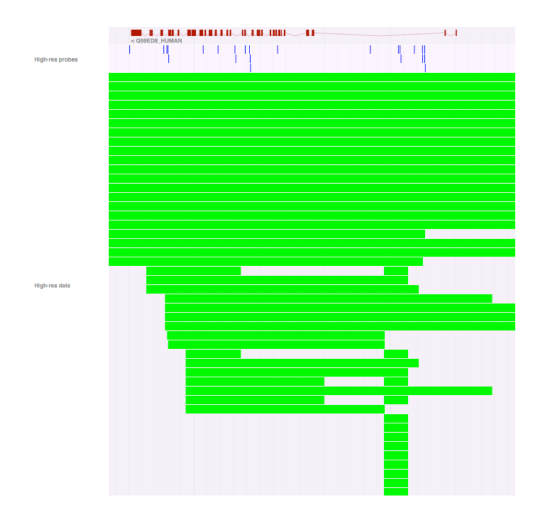


Figure 5.7. Intragenic deletions within *NOTCH1* result in the formation of the oncogenic NOTCH-IC protein. Although some deletions span the entire gene, some may result in a C-terminal truncation containing only the intracellular part of the protein. The *NOTCH1* gene is shown in red. Copy number markers are shown in blue. Deletions are shown in green.

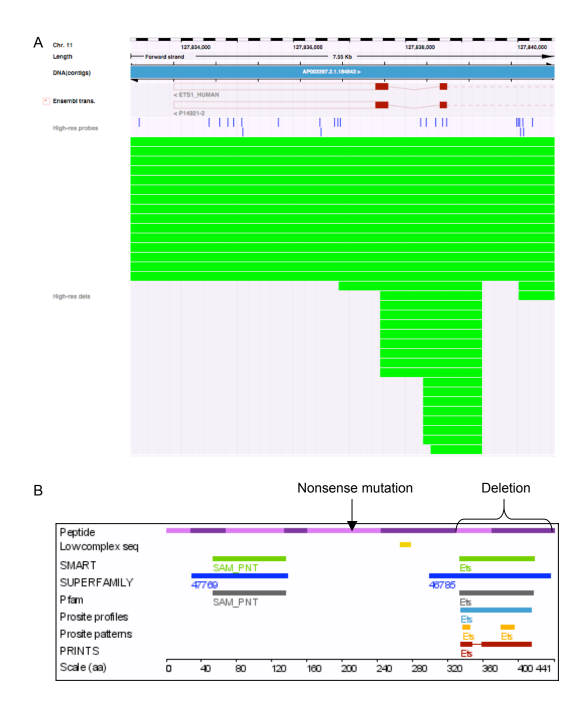


Figure 5.8. Mutations in the *ETS1* gene result in removal of the Ets domain. (A) Deletions within *ETS1* in human cancer cell lines. (B) The location of mutations in the context of the ETS1 protein. In Figure A, *ETS1* gene transcripts are shown in red, copy number markers are shown in blue and deletions are shown in green. Figure B shows the location of the SAM_PNT and Ets domains in the ETS1 protein (extracted from Ensembl geneview), and the position of the nonsense mutation in COSMIC and focal deletions in the human cancer cell lines.

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CIS gene	Mouse Ensembl ID	Position of CIS relative to gene	Cancer	COSMIC	Mullighan	CNV	Nanog BS	Oct4 BS	n53 BS	Number of deletions	P-value	Gene in MCR?	Number of genes in minimal region	Number of TSGs in minimal region	Other genes deleted in more cell lines?
Wwox	ENSMUSG0000004637	Inside	-	-	-		Nanog	Oct4	-	87	0.0011	Y	0	0	
Etv6	ENSMUSG00000030199	Inside	Dominant	-	Mullighan	-	-	-	-	60	0.0018	Y	0	0	
Zfp438	ENSMUSG00000050945	Upstream	-	-	-	-	-	-	-	33	0.0077	Ý	ō	0	
Lgals9	ENSMUSG0000001123	Upstream	-	-	Mullighan	CNV	-	-	-	31	0.0083		0	0	
Dym	ENSMUSG0000035765	Inside	-	-	-	-	-	-	-	19	0.0138	Y	ō	0	
Sdk1	ENSMUSG0000039683	Downstream	-	-	Mullighan	CNV	Nanog	Oct4	-	19	0.0138	Ý	ō	ō	
Acot11	ENSMUSG0000034853	Upstream	-	-	-	-	-	-	-	15	0.0176	Y	0	0	
Notch1	ENSMUSG0000026923	Inside	Dominant	COSMIC	-	CNV	-	-	p53	12	0.0237	Ý	ō	0	
Dock8	ENSMUSG0000052085	Upstream	-	-	Mullighan		-	-	-	11	0.0265	Y	0	0	
Ntn1	ENSMUSG00000020902	Downstream	-	-	Mullighan	-	Nanog	-	-	10	0.0282	Ý	Ő	0	
Vps13d	ENSMUSG0000020220	Inside	-	-	-	-	-	-	-	10	0.0282	Ý	ō	0	
Pik3cd	ENSMUSG0000039936	Upstream	-	-	-	-	-	Oct4	-	8	0.0382	Ý	1	0	Y
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	7	0.0422	Ý	0	0	
Jazf1	ENSMUSG0000063568	Inside	Dominant	-	Mullighan	CNV	-	-	-	7	0.0422	Y	0	0	
Rcbtb2	ENSMUSG00000022106	Upstream	-		Mullighan	-	-	-	-	, 7	0.0422	Ý	Ő	0	
Zfp608	ENSMUSG0000052713	Upstream	-		-	-	Nanog	-	-	7	0.0422	Ý	ō	0	
Zmiz1	ENSMUSG0000007817	Upstream/Inside	-	-	-	-	-	Oct4	-	6	0.0500	Y	0	0	
Ets1	ENSMUSG00000032035	Upstream/Downstream	-	COSMIC	-	-	-	-	-	6	0.0500	Ý	Ő	0	
Nedd4l	ENSMUSG0000024589	Upstream	-	-	-	CNV	Nanog	-	p53	5	0.0569	Ý	ō	0	
Lrrfip1	ENSMUSG0000026305	Inside	-	-	Mullighan	-	Nanog	-	-	5	0.0569	Ý	1	0	Y
Cldn10a	ENSMUSG0000022132	Inside	-	-	-	-	-	-	-	5	0.0569	Ý	0	0	-
Foxp1	ENSMUSG0000030067	Inside	-	COSMIC	-	-	Nanog	-	-	5	0.0569	Y	0	0	
Arhqef3	ENSMUSG00000021895	Inside	-	-	-	-	Nanog	-	-	5	0.0569	Ý	Ő	0	
Bcl11b	ENSMUSG00000048251	Inside	Dominant	COSMIC	-	-	-	-	-	5	0.0569	Ý	Ő	0	
Fqfr2	ENSMUSG0000030849	Downstream	Dominant		-	-	-	-	-	4	0.0714	Y	0	0	
Tspan2	ENSMUSG00000027858	Upstream	-	-	-	CNV	-	-	-	4	0.0714	Ý	0	0	
Tbc1d1	ENSMUSG0000029174	Downstream	-	-	-	-	Nanog	-	-	4	0.0714	· Y	0	0	
Flt3	ENSMUSG00000042817	Inside	Dominant	COSMIC	-	-	-	-	-	4	0.0714	Ý	0	0	
	ENSMUSG00000020589	Downstream	-	-	-	-	Nanog	-	-	4	0.0714	Ý	0	0	
Pml	ENSMUSG0000036986	Inside	Dominant	COSMIC	-	CNV		Oct4	-	4	0.0714	· Y	0	0	
Evi2b	ENSMUSG00000070354	Inside	-	-	-	-	-	-	-	4	0.0714		4	1	Y
Fut8	ENSMUSG00000021065	Upstream	_	_	_	-	_	_	-	4	0.0714	Y		0	
Vpreb2	ENSMUSG00000059280	Upstream	-	-	-	-	-	-	-	4	0.0714		1	0	
Ksr1	ENSMUSG00000018334	Downstream	-	COSMIC	-	CNV	-	-	-	3	0.0940	Y	0	0	
Tafbr3	ENSMUSG0000029287	Inside	-	-	-	-	-	Oct4	-	3	0.0940	· Y	0	0	
Sema4d	ENSMUSG0000021451	Inside	-	-	-	-	-	-	-	3	0.0940	Ý	0	0	
Cuqbp2	ENSMUSG0000002107	Inside	-	-	-	-	-	Oct4	-	3	0.0940	Ý	0	0	
Lef1	ENSMUSG00000027985	Upstream	-		Mullighan		Nanog	-	_	3	0.0940	Ý	0	0	
Runx1	ENSMUSG0000022952	Upstream	Dominant		-		-		_	3	0.0940	Y	0	0	
Kit	ENSMUSG00000022932	Downstream	Dominant		_		_		-	2	0.0940	· ·	0	0	
ML	LING/103000000000072	Downau cuill	Domiliant	CODINIC						J	0.0940		0	0	

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CIS gene	Mouse Ensembl ID	Position of CIS relative to gene	Cancer gene	COSMIC	Mullighar	1 CNV	Nanog BS	Oct4 B	5 p53 BS	Number of deletions	<i>P</i> -value	Gene in MCR?	Number of genes in minimal region	Number of TSGs in minimal region	Other genes deleted in more cell lines?
Wwox	ENSMUSG0000004637	Inside	-	-	-	CNV	Nanog	Oct4	-	16	0.0014	Y	0	0	
Etv6	ENSMUSG0000030199	Inside	Dominant	-	Mullighan	-	-	-	-	11	0.0067	Y	0	0	
Lgals9	ENSMUSG0000001123	Upstream	-	-	Mullighan	CNV	-	-	-	5	0.0194		0	0	
Rcbtb2	ENSMUSG0000022106	Upstream	-	COSMIC	Mullighan	-	-	-	-	4	0.0229	Y	0	0	
Zfp438	ENSMUSG0000050945	Upstream	-	-	-	-	-	-	-	3	0.0280	Y	0	0	
Ccnd2	ENSMUSG0000000184	Upstream	Dominant	-	-	-	-	-	-	3	0.0280	Y	0	0	
Notch1	ENSMUSG0000026923	Inside	Dominant	COSMIC	-	CNV	-	-	p53	3	0.0280	Y	0	0	
Dym	ENSMUSG0000035765	Inside	-	-	-	-	-	-	-	2	0.0379	Y	0	0	
Zmiz1	ENSMUSG0000007817	Upstream/Inside	-	-	-	-	-	Oct4	-	2	0.0379	Y	0	0	
Foxp1	ENSMUSG0000030067	Inside	-	COSMIC	-	-	Nanog	-	-	2	0.0379	Y	0	0	
Tbc1d1	ENSMUSG0000029174	Downstream	-	-	-	-	Nanog	-	-	2	0.0379	Y	0	0	
Vps13d	ENSMUSG0000020220	Inside	-	-	-	-	-	-	-	2	0.0379	Y	0	0	
Sdk1	ENSMUSG0000039683	Downstream	-	-	Mullighan	CNV	Nanog	Oct4	-	2	0.0379	Y	0	0	
Ntn1	ENSMUSG0000020902	Downstream	-	-	Mullighan	-	Nanog	-	-	2	0.0379	Y	0	0	
Fut8	ENSMUSG0000021065	Upstream	-	-		-		-	-	2	0.0379	Y	0	0	
Vpreb2	ENSMUSG0000059280	Upstream	-	-	-	-	-	-	-	2	0.0379		1	0	

Table 5.6. A list of CIS genes that are in recurrent deletions of copy number 0.3 or less across all cell lines (A) and across haematopoietic and lymphoid cancer cell lines only (B). Cancer gene = known cancer gene in Cancer Gene Census; COSMIC = gene contains somatic mutations in COSMIC database; Mullighan = gene within deletion in Mullighan et al. (2007) dataset of acute lymphoblastic leukaemias; CNV = gene within CNV identified in Redon *et al.* (2006); "[Nanog, Oct4, p53] BS" = gene contains binding site for Nanog, Oct4 and p53, respectively. "minimal region" = minimal deleted region containing CIS gene; "MCR" = minimal deleted region from across genome, not centred on CIS gene; "Number of genes in minimal region" = number of genes other than the CIS gene within the minimal deleted region.

less within *PIK3CD* were significantly over-represented among cancer cell lines derived from the autonomic ganglia in this study ($P=6.37 \times 10^{-8}$). The human orthologues of 11 other CIS genes on chromosome 1 were also over-represented among these cell lines, but *PIK3CD* was deleted in the highest number of cell lines (11), which lends support to the conclusions of Caren *et al.* (2007). However, while some of the insertions disrupting *Pik3cd* were within the gene, most were upstream in the antisense orientation, suggesting that *Pik3cd* is upregulated in the MuLV-induced mouse lymphomas. Deletions within *WWOX* that had a copy number of 0.3 or less were over-represented in lung cancer cell lines ($P=2.91 \times 10^{-4}$). Loss of *WWOX* is strongly associated with tumour histology and aggressiveness of non-small cell lung cancers (Donati *et al.*, 2007).

All of the haematopoietic and lymphoid tissue-specific candidates within deletions of copy number less than or equal to 0.6 were also found among candidates identified across all cell lines (Table 5.5B). Of the candidates with *P*-values of less than 0.05 in deletions of copy number 0.3 or less (Table 5.6B), 11 out of 16 haematopoietic and lymphoid candidates also had *P*-values of less than 0.05 across all cell lines. The remaining 5 had *P*-values of less than 0.1 and included *FOXP1*, which was also identified in the 10K analysis (Section 4.5.2.3) and was presented as a putative tumour suppressor gene in Section 3.4.3, and *ZMIZ1*, which may play both tumour suppressive and oncogenic roles (see Section 3.3).

A number of interesting candidates were also identified among those genes occurring in significantly recurrent deletions in specific tissue types. These included *FLI1*, *MYO18A*, *TGFBR3* and *SKI*, all of which were in MCRs across all cell lines. *FL11* and *MYO18A* contained recurrent deletions in bone cancer cell lines (P=0.0371 for both genes). Interestingly, a translocation that leads to the formation of an EWS-FL11 fusion protein is present in 95% of Ewing's sarcomas, which are tumours of the bone and soft tissue (Delattre *et al.*, 1994). However, *FL11* is an oncogene, and all but three of the deletions spanning this gene also encompassed other genes (including *ETS1*). *MYO18A* has not been implicated in cancer, but has been shown to be expressed in bone marrow stromal cells, where it may play a role in the maintenance of cell architecture (Furusawa *et al.*, 2000). TGF β receptor type III precursor *TGFBR3* is implicated as a tumour suppressor gene in a range of cancer types, including non-small cell lung cancer (Finger *et al.*, 2008), pancreatic cancer (Gordon *et al.*, 2008), prostate cancer (Turley *et al.*, 2007) and breast cancer (Dong *et al.*, 2007), but the identification of recurrent deletions in soft tissue

cancer cell lines (P=0.0270) suggests it is also important in these tumours. *SKI* plays an oncogenic role in some cancers, such as human melanomas (for review, see Reed *et al.*, 2005), but *Ski*-deficient heterozygous mice show an increased susceptibility to tumourigenesis (Shinagawa *et al.*, 2001) and reduced *SKI* expression in breast and lung cancer cells enhances tumour metastasis (Le Scolan *et al.*, 2008). The *P*-value for the number of deletions of *SKI* across all cell lines was 0.0538, but was lower for deletions in cancer cell lines of the kidney (P=0.0351), central nervous system (P=0.0166) and autonomic ganglia (P=0.0130).

The number of deletions was slightly higher across CIS genes than non-CIS genes for deletions with a copy number of 0.6 or less (P=0.0306), and significantly higher for deletions with a copy number of 0.3 or less (P=0.00340). For CIS genes in deletions of copy number less than or equal to 0.6, the median and mean number of deletions were 8 and 13.54, while for non-CIS genes these were 7 and 11.85, respectively. The maximum number of deletions among CIS genes was 237, occurring within *WWOX*. For CIS genes in deletions of and 1.449, respectively, compared with 0 and 1.303 for non-CIS genes. The maximum number among CIS genes was 87, again occurring within *WWOX*.

The work presented in this section demonstrates the overlap between mouse CIS genes and regions of copy number change in human cancer, and provides a selection of candidates that warrant further investigation.

5.4 Comparison between high-resolution and 10K CGH data

Figure 5.9 provides an illustration of the difference in resolution between the two SNP array CGH platforms. A number of deletions that are clearly visible in the high-resolution data are completely missed by the 10K data for B-cell lymphoma cell line DOHH-2 (Figures 5.9A and B). The breast cancer cell line HCC1143 shows considerable variation in copy number across the genome, but the large distances between adjacent SNPs, and therefore between segments of differing copy number, in the 10K data make it impossible to determine the copy numbers of genes in the intervening regions (Figure 5.9C). This problem is alleviated in the high-resolution data, where most genes contain, or are very close to, a copy number marker (Figure 5.9D).

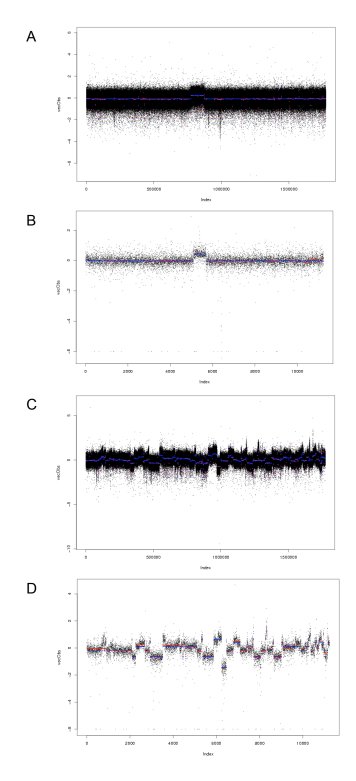


Figure 5.9. High-resolution and 10K SNP array CGH data for the entire genome of B-cell lymphoma cell line DOHH-2 (A and B, respectively) and breast cancer cell line HCC1143 (C and D, respectively). Black points are copy number values for individual SNPs, red lines are mean copy numbers for DNAcopy segments, blue lines are copy number values for merged segments. Markers are positioned according to their order in the genome rather than their exact coordinates. Copy numbers are provided as log_2 -ratios.

In order to directly compare the use of high-resolution and 10K CGH data for integrative analyses with the mouse CIS genes, the procedures described in Section 5.3 were applied to the 598 cell lines within the 10K dataset that were also within the high-resolution dataset. Figure 5.10 shows the distribution of the number of amplicons and deletions in these cell lines and the distribution of the lengths of aberrations as determined using the 10K SNP array. To ensure that the dataset was treated identically to the higher resolution data, the start and end coordinates of an amplicon or deletion were taken as the halfway point between the first or last amplified or deleted SNP in a segment and the nearest SNP in the adjacent segment, and amplicons and deletions were defined as regions with a copy number of 1.7 or more, and 0.6 or less, respectively. The average number of amplicons was $1.34 (\pm 1.92)$ per cell line. The amplicons were on average $17.0 (\pm 28.31)$ Mb in size and contained 173.14 (±293.84) genes. The average number of deletions was 4.45 (± 3.79) . These deletions were on average 20.97 (± 36.04) Mb in size, encompassing 198.05 (±336.49) genes. Amplicons and deletions were therefore considerably longer in the lower resolution dataset. As mentioned in Section 5.2.1, the higher resolution data are over-segmented, but the lower resolution data may be missing small regions of copy number change and some genes may be incorrectly assigned to amplicons and deletions because of the large distances between probes. The number of minimal common regions (MCRs) for amplicons and deletions were 300 and 741, respectively. These are ~30-fold and 50-fold less than the numbers obtained using the high-resolution data, again reflecting possible over-segmentation of the high-resolution data but also the increased ability to detect small regions of copy number change.

The number of amplicons in the 10K data that contained each CIS gene were counted and compared, using the Mann Whitney U test, to the values for the high-resolution data generated in Section 5.3.2. For the 10K data, the median and mean numbers of amplicons were 2 and 3.82, compared with 1 and 2.768 for the high-resolution data. Therefore, the values across all CIS genes were significantly lower in the high-resolution data $(P=6.77 \times 10^{-8})$. Likewise, the number of deletions of copy number less than or equal to 0.6 was significantly lower $(P=5.12 \times 10^{-5})$, with the median and mean measuring 11 and 16.54, respectively, in the 10K data, compared with 8 and 13.54 in the high-resolution data. However, a global analysis of CIS genes in amplicons and deletions in the 10K data, based on the analysis described in Section 5.3.1, demonstrated that the significance of the overlap between CIS genes and amplicons was much lower in the 10K data than in the high-resolution data, and there was no over-representation of CIS genes within

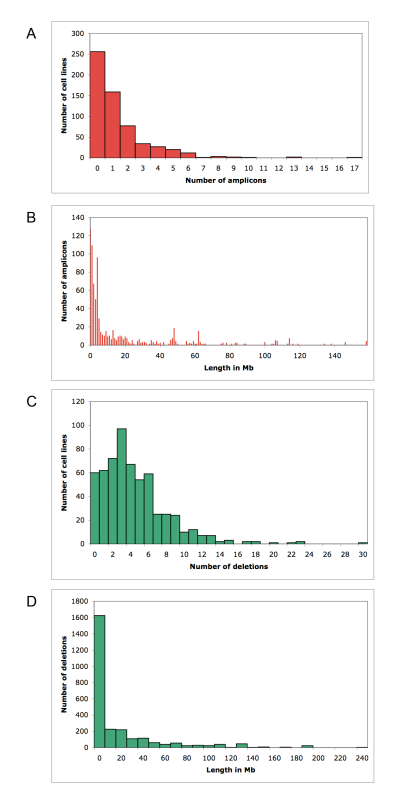


Figure 5.10. Characterisation of amplicons and deletions in 598 human cancer cell lines analysed using 10K SNP array CGH. (A) Number of amplicons per cell line. (B) Length of amplicons. (C) Number of deletions per cell line. (D) Length of deletions.

deletions (Figure 5.11). Therefore, while CIS genes were amplified and deleted in more cell lines, on average, in the 10K dataset than in the high-resolution dataset, non-CIS genes were also amplified and deleted in more cell lines. It is possible that due to the sparseness of the data, regions of copy number change have been incorrectly called or have been extended beyond their true boundaries.

The median and mean number of deletions of copy number 0.3 or less that contained each CIS gene was 0 and 0.444, respectively, for the 10K data, compared with 0 and 1.449 for the high-resolution data. Consequently, the values across all CIS genes were significantly higher in the high-resolution dataset (P=0.00144), which is in contrast to the results for amplicons and higher copy deletions. This may reflect the fact that homozygous deletions are likely to be small and are therefore missed by the lower resolution analysis. CIS genes were over-represented in deletions of this copy number in the high-resolution, dataset.

The superiority of the high-resolution dataset was demonstrated by applying the Matthew's Correlation Coefficient (MCC), which is described in Section 2.10.2 and also used in Section 4.6. In each cell line, the number of CIS genes in amplicons was counted and defined as the number of true positives. Non-CIS genes in amplicons were defined as false positives. CIS genes that did not occur in amplicons were defined as false negatives and non-CIS outside of amplicons were defined as true negatives. The number of true and false positives and negatives in each cell line were then added together to give the number across all cell lines. This prevents a CIS gene that is amplified in just one cell line from having the same weight as one that is amplified in multiple cell lines. The procedure was repeated using deletions of copy number less than 0.6 and 0.3. It was also repeated using known cancer genes from the Cancer Gene Census in place of CIS genes, with oncogenes being counted in amplicons, and tumour suppressor genes being counted in deletions. The results are presented in Table 5.7. Although it is not expected that all CIS genes or known cancer genes contribute to human cancer through a change in copy number, these tests do give an indication of the comparative reliability of the two datasets. In concurrence with the results above, the coverage of CIS genes in amplicons and deletions of copy number 0.6 or below was higher in the 10K data but the accuracy was higher in the high-resolution data, while both the coverage and accuracy of CIS genes in deletions of copy number 0.3 or less were higher in the high-resolution data. In all cases, a higher MCC score was obtained using the high-resolution dataset. The same

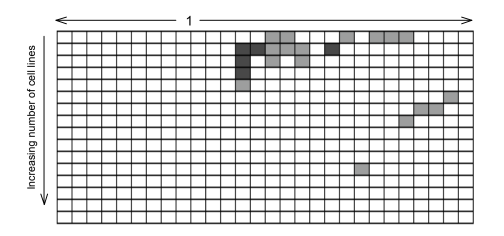


Figure 5.11. Over-representation of CIS genes in amplicons of varying copy number threshold and number of cell lines in the 10K dataset. Each box represents the significance of the association between CIS genes and amplicons/deletions at a given copy number threshold and cell line number. P<0.0001, black; P<0.001, dark grey, P<0.05, light grey. Copy number thresholds below 1 represent deletions, and range from 0.1 to 0.9 with 0.1 increments. Copy number thresholds above 1 represent amplicons, and range from 1.1 to 2.9 with 0.1 increments. The number of cell lines increases in increments of 1, up to a cut-off of 16 cell lines.

Region	Resolution	ТР	TN	FP	FN	Accuracy	Coverage	мсс
A	High	1060	9954616	39034	227974	0.026438	0.00463	0.001710
Amplicons	Low	1704	9922569	71081	227330	0.023411	0.00744	0.000576
Deletione (0.0	High	5184	9792549	201101	223850	0.025130	0.02263	0.002643
Deletions ≤ 0.6	Low	7939	9636527	357123	221095	0.021747	0.03466	-0.000855
Deletions $\leq = 0.3$	High	555	9971540	22110	228479	0.024487	0.00242	0.000663
Deletions <= 0.3	Low	210	9981105	12545	228824	0.016464	0.00092	-0.001419
В								
Region	Resolution	TP	TN	FP	FN	Accuracy	Coverage	МСС
Amplicons	High	1017	10032313	39077	150277	0.025365	0.006722	0.005491
Amplicons	Low	1199	9999804	71586	150095	0.016473	0.007925	0.001174

36527

35755

37385

37481

0.005560

0.005257

0.012751

0.015131

0.030445

0.050937

0.007671

0.005123

0.004440

0.004990

0.007052

0.006676

9979872 205138

363143

22376

12562

9821867

10162634

<u>1017244</u>8

High

Low

High

Low

Deletions <= 0.6

Deletions <= 0.3

1147

1919

289

193

Table 5.7. Comparison of the high- and low-resolution datasets based on the proportion of CIS genes (A) and known cancer genes (B) that are amplified and
deleted. TP = number of CIS/cancer genes in amplicons/deletions, TN = number of non-
CIS/non-cancer genes that are not in amplicons/deletions, FP = number of non-CIS/non-
cancer genes in amplicons/deletions, FN = number of CIS/cancer genes that are not in
amplicons/deletions. Accuracy is given by TP/(TP+FP); Coverage is given by
TP/(TP+FN). MCC = Matthew's Correlation Coefficient. Deletions $\leq 0.6/03 =$
deletions with a copy number of less than or equal to $0.6/0.3$. Amplicons are regions with
a copy number of 2.7 or above.

pattern was observed for known cancer genes, except for copy number 0.6 or less, where the MCC score was very slightly higher in the 10K dataset. These results demonstrate that at higher resolution, regions of copy number change are likely to be more defined, making it easier to identify the critical gene(s) that contribute to tumourigenesis. In addition, it is possible to identify smaller changes, particularly deletions, that are missed by lower resolution CGH. Importantly, it appears that the possible over-segmentation of the high-resolution data has not been detrimental to the analysis.

Finally, individual candidates identified in the 10K analysis were compared to those identified in the high-resolution analysis. The 10K dataset contained 20 genes in statistically significant recurrent amplicons (Table 5.8A). Only 2 of these genes were known oncogenes, compared with 9 in the high-resolution dataset, and both were identified in fewer cell lines in the 10K dataset. All of the remaining significantly recurring genes in the 10K dataset were amplified in a higher number of cell lines than those in the high-resolution dataset. However, 11 of these genes were not within an MCR in the 10K data, while a further 2 (ADRBK1 and GPR152) were co-amplified with CCND1 in 21 cell lines. In the high-resolution dataset, ADRBK1 and GPR152 were found in 5 and 4 cell lines, respectively, while CCND1 was found in 23 lines. Therefore, in this case, it is only possible to discern the critical cancer gene at higher resolution. NDRG3 and SLA2 were amplified in 24 and 23 cell lines, respectively, in the 10K dataset but just 1 and 0 cell lines, respectively, in the high-resolution dataset. Visual inspection of the SNPs from the 10K dataset in the context of Ensembl showed that there were none overlapping NDRG3 and SLA2, and therefore the copy number for these genes could not be accurately determined (Figure 5.12A). In order to directly compare the positions of copy number markers in the 10K and high-resolution datasets, the coordinates of the 10K SNPs, which were originally mapped to the NCBI 35 human genome assembly, were converted to NCBI 36 using the UCSC LiftOver tool (http://genome.ucsc.edu/cgibin/hgLiftOver). Copy number markers for both datasets were displayed in Ensembl contigview using a DAS track (see Section 2.6 for further details regarding DAS). Interestingly, the regions surrounding the closest 10K SNPs to NDRG3 and SLA2 were not amplified in the high-resolution dataset, suggesting that the sparseness of probes can lead to the miscalling of copy number changes across large regions. CAPSL and ZNF217 (known as Zfp217 in the mouse) were also amplified in a greater number of cell lines in the 10K dataset, but these too did not contain any 10K SNPs. In the case of ZNF217, the number of high-resolution amplicons overlapping the adjacent gene, BCAS1, was slightly

	Position of				Number of			Amplified in more
	CIS relative	Number of		Gene in	additional	Number of known	Other genes	
Gene	to gene	amplicons	P-value	MCR?	genes in MCR	oncogenes in MCR	in MCR	resolution?
Мус	Upstream	39	0.0008	Y	0	0		
Capsl	Downstream	25	0.0055	Y	0	0		Y
Slc1a3	Downstream	24	0.0089	N	0	0		Y
Fyb	Inside	24	0.0089	N	0	0		Y
Sla	Inside	21	0.0114	N	0	0		Y
Zfp217	Upstream	21	0.0114	Y	0	0		Y
Ptp4a3	Inside	20	0.0178	N	0	0		Y
Ndrg3	Downstream	20	0.0178	Y	1	0	Sla2	Y
Sla2	Inside	19	0.0227	Y	1	0	Ndrg3	Y
Bcl2l1	Inside	18	0.0298	N	0	0		Y
Ncoa3	Upstream	18	0.0298	N	0	0		Y
Prkcbp1	Upstream	17	0.0375	N	0	0		Y
Ccnd1	Upstream	17	0.0375	Y	2	0	Gpr152, Adrbk1	
Serinc3	Inside	17	0.0375	N	0	0		Y
Ppp1r16b	Inside	17	0.0375	N	0	0		Y
Gpr152	Downstream	17	0.0375	Y	2	1	Adrbk1, Ccnd1	Y
Stk4	Inside	17	0.0375	N	0	0		Y
Adrbk1	Inside	17	0.0375	Y	2	1	Gpr152, Ccnd1	Y
Cldn10a	Inside	16	0.0394	N	0	0		Y
Ubac2	Inside	14	0.0458	Y	0	0		Y

В

	Position of				Number of	Amplified in more
	CIS relative	Number of		Gene in	additional	cell lines than
Gene	to gene	amplicons	P-value	MCR?	genes in MCR	high-resolution?
Cyb5	Downstream	89	0.0030	N	0	Y
Nedd4l	Upstream	84	0.0050	N	0	Y
Mbd2	Upstream	80	0.0060	N	0	Y
Dym	Inside	76	0.0082	N	0	Y
Dock8	Upstream	76	0.0082	Y	0	Y
Rcbtb2	Upstream	71	0.0124	Y	0	Y
Lcp1	Inside	66	0.0177	N	0	Y
1190002H23Rik	Downstream	64	0.0227	N	0	Y
Aqp4	Downstream	64	0.0227	Y	0	Y
Elf1	Inside	63	0.0241	N	0	Y
1700081D17Rik	Downstream	61	0.0291	N	0	Y
Katnal1	Upstream	60	0.0311	Y	0	Y
Spata13	Inside	59	0.0350	N	0	Y
Ubac2	Inside	59	0.0350	N	0	Y
Flt3	Inside	58	0.0370	N	0	Y
Cldn10a	Inside	55	0.0404	N	0	Y
D18Ertd653e	Inside	52	0.0440	N	0	Y

С

	Position of				Number of			Amplified in more
	CIS relative	Number of		Gene in	additional	Number of known	Other genes in	cell lines than high
Gene	to gene	amplicons	P-value	MCR?	genes in MCR	oncogenes in MCR	MCR	resolution?
Cyb5	Downstream	9	0.0052	N	0	0		Y
Mbd2	Upstream	8	0.0061	N	0	0		Y
Nedd4l	Upstream	6	0.0093	N	0	0		Y
Rasgrp1	Upstream	6	0.0093	N	0	0		Y
Rcbtb2	Upstream	4	0.0170	Y	0	0		
Dock8	Upstream	4	0.0170	N	0	0		Y
							Vpreb2, BC030863, Cecr5,	
Bid	Upstream	3	0.0405	Y	4	0	Tuba8	Y
	·						Vpreb2, BC030863, Bid,	
Cecr5	Upstream	3	0.0405	Y	4	0	Tuba8	Y
Wwox	Inside	3	0.0405	Y	0	0	Vpreb2, Cecr5,	
BC030863	Downstream	3	0.0405	Y	4	0	Bid, Tuba8	Y
Ubac2	Inside	3	0.0405	N	0	0	.,	Y
							BC030863, Cecr5,	
Vpreb2	Upstream	3	0.0405	Y	4	0	Bid, Tuba8 BC030863, Cecr5,	Y
Tuba8	Downstream	3	0.0405	Y	4	0	Bid, Vpreb2	Y

Table 5.8. A list of CIS genes that are in recurrent amplicons (A), recurrent deletions of copy number 0.6 or less (B) and recurrent deletions of copy number 0.3 or less (C) in the 10K CGH dataset. "MCR" = minimal amplified or deleted region.

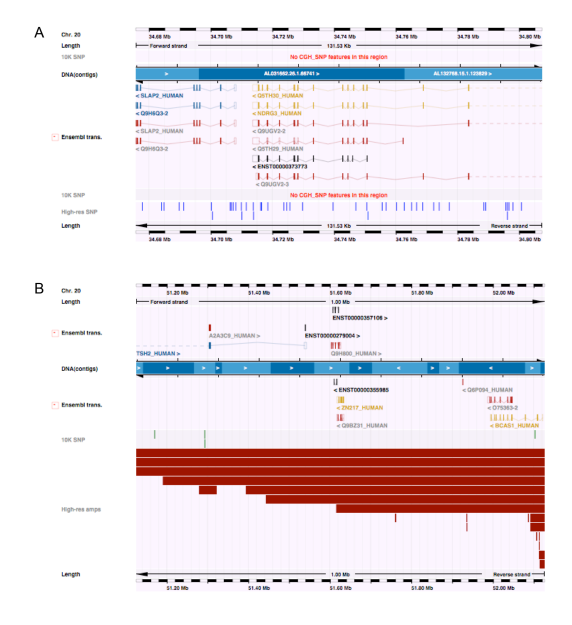


Figure 5.12. *SLA2* and *NDRG3* (A) and *ZNF217* (B) are amplified in a greater number of cell lines in the 10K dataset than in the high-resolution dataset but do not contain SNPs in the 10K dataset. In Figure A, copy number markers in the high-resolution dataset are shown in blue. In Figure B, copy number SNPs in the 10K dataset are shown in dark green. Amplicons identified in the high-resolution analysis are shown as red rectangles.

higher and may contribute to the amplicons incorrectly assigned to *ZNF217* in the 10K dataset (Figure 5.12B).

Among deletions of copy number less than or equal to 0.6, all of the 17 significantly recurrent genes in the 10K dataset (Table 5.8B) were deleted in a greater number of cell lines than in the high-resolution dataset. However, 13 of these genes were not within an MCR. Of the 4 that were, 3 (*RCBTB2, AQP4* and *KATNAL1*) did not contain any 10K SNPs, and therefore the copy numbers of these genes in the 10K dataset are not accurate. In addition, neither *Rcbtb2* nor *Aqp4* contained any intragenic MuLV insertions in mouse lymphomas, while *Katnal1* contained just 2 insertions. Only *DOCK8* contained SNPs, and it is not clear why the number of deletions was so much greater at lower resolution. Just 4 of the 24 significantly recurrent deleted genes in the high-resolution dataset were significantly recurrent in the 10K dataset. A number of promising candidates from the high-resolution analysis, including *WWOX*, *SDK1* and *CBFA2T3*, were deleted in fewer cell lines, and were not significant, in the 10K dataset.

For deletions of copy number 0.3 or less, 11 genes were deleted in a higher number of cell lines in the 10K dataset. 6 were not within an MCR and the remaining 5 resided in the same MCR (Table 5.8C). *WWOX* was deleted in fewer cell lines than in the high-resolution data, and *RCBTB2* was deleted in the same number of lines. *WWOX* and *RCBTB2* were the only significantly recurrent deleted genes from the high-resolution dataset that were represented among significant genes from the 10K dataset.

The lists of significantly recurrent amplified and deleted CIS genes differ considerably between the 10K and high-resolution datasets. These differences reflect the low density of copy number markers in the 10K analysis, which results in small deletions being missed and, therefore, putative tumour suppressor genes going undetected. They also result from genes being incorrectly flagged as promising candidates when copy number regions are miscalled or are extended beyond their true boundaries. However, as discussed in Sections 5.3.2.2 and 5.3.2.3, in the high-resolution analysis, some of the significantly amplified and deleted genes, including *ETV6*, *CUGBP2* and *CCND2*, contained small, repetitive, regions of copy number change across multiple cell lines that did not look likely to contribute to tumourigenesis. These tiny changes were not identified in the lower resolution analysis and therefore amplifications and deletions within these genes were not significantly recurrent. Therefore, for amplicons, which are

often large and encompass multiple genes, the most convincing candidate oncogenes may be those that are identified using both platforms. For deletions, which are often smaller, the identification of candidate tumour suppressor genes is more reliant on the highresolution dataset.

Having established that candidates may be incorrectly assigned to amplicons or deletions because of the low resolution of the 10K data, it is important to revisit the candidates identified in Chapter 4 to determine whether they are still valid. Among mouse candidates near to CISs with a P-value of less than 0.001, 17 of those presented as putative oncogenes in Table 4.4 were not included within the more conservative dataset used in this chapter and have therefore not been studied in relation to the high-resolution These included Meis1, Mmp13, Smad7, Lrrc5 and Taok3, all of which were data. discussed in Section 4.5.2.1.1. Of the remaining 37 candidates, all were found within at least one human amplicon, but only 11 had a *P*-value of less than 0.1 in the recurrence analysis. Apart from Zfp217, all of these genes were known oncogenes from the Cancer Gene Census. The P-values for Nfkb1, Slamf6 and Rreb1, which were presented as candidates in Section 4.5.2.1.1, were greater than 0.1. However, Nfkb1 was amplified in a single cell line in both datasets and, while *Slamf6* and *Rreb1* were amplified in fewer cell lines in the high-resolution data (5 and 2 cell lines, respectively), the minimal amplified regions contained just 3 and 2 genes, respectively, and included no known oncogenes. Hence there is no convincing evidence from the higher resolution data to suggest that these genes are not amplified in cancer.

Of the candidates that were identified in deletions in the 10K data, 39 were discussed in Section 4.5.2.3. 27 of these genes were not within the list of mouse candidate cancer genes used in this chapter. This is because the lists used in Chapter 4 included all genes containing insertions, whether the insertions clustered into a CIS or not, whereas only CIS genes were considered in the list generated in Chapter 2 and used in this chapter. Among the remaining 12 genes, 5 were recurrently deleted to copy number 0.6 and below and/or 0.2 and below with a *P*-value of less than 0.1. These were *Wwox*, *Foxp1*, *Sdk1*, *Bcl11b*, *Lef1* and *Mbd2*. The only implicated tumour suppressor gene that was no longer identified in the high-resolution analysis was *Gpr56*. Known and implicated oncogenes *Evi1*, *Myc*, *Fli1*, *Rasgrp1* and *Map3k8* were deleted, but not significantly, therefore providing further evidence that these deleted genes are most likely passengers, rather than causative genes, in human cancers.

5.5 Identification of co-operating cancer genes

5.5.1 Genotype-specific cancer genes

Most of the cancer cell lines used in this study are part of the Cancer Cell Line Project undertaken by the Wellcome Trust Sanger Institute Cancer Genome Project (http://www.sanger.ac.uk/genetics/CGP/CellLines/). The aim of the project is to systematically sequence all known cancer genes in all of the selected cell lines. 595 of the 598 cell lines used in this study have been analysed for somatic mutations in TP53 and CDKN2A. 311 have mutations in TP53, while 160 have mutations in CDKN2A. For each amplified or deleted CIS gene, a 2-tailed Fisher Exact Test was used to determine whether there was any significant difference between the number of cell lines with TP53 or CDKN2A mutations that contained the amplified or deleted gene compared with the number with TP53 or CDKN2A mutations that did not contain the amplified or deleted gene. A positive association between an amplified or deleted gene and cell lines bearing a TP53 or CDKN2A mutation suggests that the disrupted CIS gene may co-operate with inactivation of TP53 or CDKN2A in human tumourigenesis. Likewise, a negative association suggests that the genes never co-operate, possibly because they act in the same cancer pathway. This analysis is equivalent to the genotype-specific analysis described in Section 3.5.1 that was performed on tumours generated in mice deficient in p53, or p19 and/or p16 (p16 and p19 are collectively known as Cdkn2a). The identification of genes that co-operate in both species provides strong evidence that the co-operation is real, and that cancer pathways are conserved between species.

This analysis was performed on CIS genes amplified to copy number 1.7 or above and deleted to copy number 0.6, or 0.3, or below. While a small number of tests had *P*-values of less than 0.05, none of the tests were below the significance level adjusted to account for multiple testing (either with *q*-values, determined using the R package QVALUE (see Section 3.5.1), or using the Bonferroni correction, where the significance level of 0.05 was divided by the number of amplified or deleted genes). Nevertheless, associations with a *P*-value of less than 0.05, which are presented in Table 5.9, were investigated to determine whether there was any evidence in the literature to support the findings. *CCND1* amplification co-occurred with *TP53* mutation with a *P*-value of 0.0173. Overexpression of *CCND1* and mutant *TP53* have been shown to co-occur in human uterine endometrial carcinomas (Nikaido *et al.*, 1996). However, insertional mutagenesis of *Ccnd1* was associated with the loss of *Cdkn2a*, and not with *Trp53*, in mouse

Region	Genotype	CIS gene	Mouse Ensembl ID	P-value
Amplicons	<i>TP53</i> up	Ccnd1	ENSMUSG00000070348	0.0173
Amplicons	1755 up	Sdk1	ENSMUSG0000039683	0.0228
		BC008155	ENSMUSG0000057411	0.0009
		Dock8	ENSMUSG0000052085	0.0012
		B3gntl1	ENSMUSG0000046605	0.0016
		1300007F04Rik	ENSMUSG0000000686	0.0035
		Zfp608	ENSMUSG0000052713	0.0075
		Mobkl2a	ENSMUSG0000003348	0.0104
		Mknk2	ENSMUSG0000020190	0.0104
		Metrnl	ENSMUSG0000039208	0.0115
		Arrdc5	ENSMUSG0000073380	0.0126
	<i>TP53</i> up	Ubac2	ENSMUSG0000041765	0.0130
		Cldn10a	ENSMUSG0000022132	0.0173
		Gadd45b	ENSMUSG0000015312	0.0184
Deletions <= 0.6		Abcg1	ENSMUSG0000024030	0.0217
		Arid3a	ENSMUSG0000019564	0.0247
		Ptbp1	ENSMUSG0000006498	0.0247
		Mbd2	ENSMUSG0000024513	0.0332
		Il6st	ENSMUSG0000021756	0.0356
		Midn	ENSMUSG0000035621	0.0360
		Gna15	ENSMUSG0000034792	0.0448
		Ski	ENSMUSG0000029050	0.0091
	<i>TP53</i> down	Park7	ENSMUSG0000028964	0.0143
	1P55 00WI	Pml	ENSMUSG0000036986	0.0160
		Prdm16	ENSMUSG0000039410	0.0308
	CDKN2A up	Kdr	ENSMUSG0000062960	0.0109
		Acot11	ENSMUSG0000034853	0.0469
Deletions <= 0.3	CDKN2A up	Vpreb2	ENSMUSG00000059280	0.0051

Table 5.9. A list of amplified and deleted CIS genes that are over- or underrepresented in cell lines that contain a mutation in *TP53* or *CDKN2A*. "Amplicons" represents regions with a copy number greater than or equal to 1.7; "Deletions ≤ 0.6 " and "Deletions ≤ 0.3 " represent regions with a copy number less than or equal to 0.6 and 0.3, respectively. "*TP53* up" and "*TP53* down" represent CIS genes that are overrepresented and under-represented, respectively, in *TP53*-mutated cell lines; "*CDKN2A* up" represents CIS genes that are over-represented in *CDKN2A*-mutated cell lines. *P*values were generated using the 2-tailed Fisher Exact Test. lymphomas. A positive association was observed between deletion of KDR and mutation in CDKN2A, and mutagenesis of Kdr was also weakly associated with loss of Cdkn2a in mouse tumours (P=0.015). KDR is a receptor for vascular endothelial growth factor (VEGF) that plays a role in tumour angiogenesis and would generally be expected to be amplified, rather than deleted, in cancer cell lines, and none of the MuLV insertions in Kdr were intragenic. However, non-endothelial KDR expression is associated with increased survival of patients with urothelial bladder carcinomas (Gakiopoulou-Givalou *et al.*, 2003).

Deletion within DJ-1 (*PARK7*) was negatively associated with *TP53* mutation. However, *PARK7* is a putative oncogene, and it does not reside within an MCR. Deletions within *PARK7* are implicated in Parkinson's disease rather than cancer (Abou-Sleiman *et al.*, 2004). Also negatively associated with *TP53* mutation were deletions within the promyelocytic leukaemia gene *PML*. PML forms an oncogenic fusion protein with RAR α in acute promyelocytic leukaemia, but alone, *PML* functions as a tumour suppressor gene. PML and p53 were identified as independent prognostic markers in gallbladder carcinomas, suggesting that they do not co-operate in tumourigenesis (Chang *et al.*, 2007). *PML* is a direct target of p53 (de Stanchina *et al.*, 2004), which suggests that there is no selective advantage in mutating both genes and provides support for the observed association.

Among the deleted genes that were positively associated with *TP53* mutations were ATrich interactive domain 3A (*ARID3A*) and growth arrest and DNA-damage-inducible protein beta (*GADD45β*). *ARID3A* contains a putative binding site for p53, and has been proposed to play a role in growth suppression mediated by p53 (Ma *et al.*, 2003). The fact that *ARID3A* deletions co-occur with *TP53* mutation suggests that other mechanisms may also contribute to the activation of *ARID3A* since inactivation of *TP53* would otherwise be sufficient to inactivate *ARID3A*. Down-regulation of *GADD45β* was found to be associated with human hepatocellular carcinoma but was inversely correlated with the presence of mutant p53 (Qiu *et al.*, 2003). In addition, p53 is believed to be involved in regulating the expression of murine *Gadd45β* (Balliet *et al.*, 2001), suggesting that inactivation of *GADD45β* and *TP53* should be negatively, rather than positively, associated. However, p53 knockdown in a human glioma cell line showed that *GADD45α* could be induced by a p53-independent mechanism (Heminger *et al.*, 2006), and it is therefore possible that the same may be true of *GADD45β*, in which case inactivation of both genes may provide an additional growth advantage on the tumour. Neither *ARID3A* nor *GADD45* β resides within an MCR, which casts doubt on the observed associations.

There was no evidence in the literature to directly support the remaining associations. This analysis has unfortunately not yielded any strong candidates for co-operation with inactivated tumour suppressor genes *TP53* and *CDKN2A*, and for those that showed a slight association, there was very little correlation with the co-operating candidates identified in the insertional mutagenesis screen. A possible explanation for the lack of association is that for some of the cell lines in which a gene is amplified or deleted, that gene may not be a critical gene and may not be contributing to tumourigenesis. Thus any association in cell lines where the CIS gene is the critical gene could be concealed. Another confounding factor is that some of the cell lines have not been screened to completion and therefore, in some cases, a cell line that is labelled as not containing a mutation may in fact do so.

5.5.2 Co-occurrence of amplified and deleted candidate cancer genes

The aim of this analysis was to identify amplified and deleted CIS genes that co-occur across human cancer cell lines. Co-occurring cancer genes may co-operate in tumourigenesis and therefore represent attractive targets for combined therapies (see Section 1.2.7). This analysis is equivalent to the analysis performed on mouse tumours in Section 3.5.2. Any candidates that are disrupted in common in both mouse and human cancers are particularly strong candidates for a co-operating role in tumourigenesis, and co-occurring genes were therefore compared across species.

Co-occurring genes were identified by taking each pair of CIS genes and counting the number of cell lines in which they were both amplified, and the number in which they were uniquely amplified. A 2-tailed Fisher Exact Test was used to determine whether there was any significant difference between the observed number of co-amplifications, and the expected number.

a	b
С	d

- a = Number of tumours in which both genes were amplified
- b = Number of tumours in which the first gene was uniquely amplified
- c = Number of tumours in which the second gene was uniquely amplified
- d = Number of tumours in which neither gene was amplified

The test was repeated to identify co-occurring deleted CIS genes, and co-occurring amplified and deleted CIS genes. Deleted genes were defined as genes with a copy number of 0.3 or less, since regions of copy number 0.6 or less can be very large and all but one of the CIS genes were identified in at least one of these deletions (see Section 5.3.2.3). To account for multiple testing, q-values were calculated using the R package QVALUE (see Section 3.5.1). Most of the co-occurrences of amplified genes were on the same chromosome and the genes may therefore co-occur simply because they are located close to one another. Significant co-occurrences between CIS genes on different chromosomes are shown in Table 5.10. None of these co-occurrences were identified among the mouse lymphomas in Section 3.5.2.

There were 3 significant co-occurrences of amplified genes, of which 2 included sodiumcoupled neutral amino acid transporter 1 (*SLC38A1*). The minimal amplified region containing *SLC38A1* also contained 33 additional genes, of which 29 were amplified in a greater number of cell lines and 14 of these were amplified to higher copy number. This suggests that *SLC38A1* is not the critical gene with which *KDR* and/or *KIT* co-operate. Interestingly, both genes in the remaining gene pair of *TMEM49* and *NCOA3* were significantly over-represented in breast cancer cell lines (see Section 5.3.2.2) and all 3 of the cell lines in which they co-occurred were derived from breast cancers. However, *TMEM49* and *NCOA3* were co-amplified with other putative breast cancer genes on chromosomes 17q23 and 20q, respectively. While it is therefore not possible to conclude that *TMEM49* and *NCOA3* co-operate in breast cancer, it does appear that genes within the two amplicons do co-operate.

There were 8 significant co-occurrences of deleted genes in the human cancer cell lines. There is no evidence in the literature to support any of these associations, and some of the genes seem unlikely candidates for a role in tumour suppression. *LRRFIP1* is over-expressed in Burkitt's Lymphoma and other cancer cell lines (Rikiyama *et al.*, 2003) and represses expression of tumour necrosis factor alpha, which is involved in apoptosis (Suriano *et al.*, 2005). Likewise, *BCL9L* is overexpressed in acute lymphoblastic

А

		Gene 1 human	Gene 2 human		Gene 1	Gene 2			
Gene 1	Gene 2	coordinates	coordinates	Shared	unique	unique	Shared cell lines	P-value	q-value
Slc38a1	Kdr		4:55639416-	2	0		NCI-H1930 (lung),	5.60E-05	0.0082
		44949475	55686519				NCI-H1693 (lung)		
Slc38a1	Kit	12:44863110-	4:55218863-	2	0	4	NCI-H1930 (lung),	8.40E-05	0.0116
		44949475	55301636				NCI-H1693 (lung)		
Tmem49	Ncoa3	17:55139811-	20:45564053-	2	4	1	HCC2218 (breast),	2.51E-04	0.0278
		55273235	45719023				MCF7 (breast),		
							BT474 (breast)		

В

		Gene 1 human	Gene 2 human		Gene 1	Gene 2			
Gene 1	Gene 2	coordinates	coordinates	Shared	unique	unique	Shared cell lines	P-value	q-value
D12Ertd553e	Tspan2	2:16594890-	1:115392155-	4	0		0 no-11 (glioma), ATN-1 (leukaemia),	1.90E-10	1.54E-06
		16710580	115433638				COLO-205 (large intestine), NCI- H322M (lung)		
Dock8	Tspan2	9:263048-	1:115392155-	4	7		0 no-11 (glioma), ATN-1 (leukaemia),	6.26E-08	1.70E-04
		455255	115433638				COLO-205 (large intestine), NCI- H322M (lung)		
Dock8	D12Ertd553e	9:263048-	2:16594890-	4	7		0 no-11 (glioma), ATN-1 (leukaemia),	6.26E-08	1.70E-04
		455255	16710580				COLO-205 (large intestine), NCI- H322M (lung)		
Bcl9l	D12Ertd553e	11:118272059-	2:16594890-	2	0		2 ATN-1 (leukaemia), COLO-205	3.36E-05	0.0391
		118286823	16710580				(large intestine)		
Bcl9l	Tspan2	11:118272059-	1:115392155-	2	0		2 ATN-1 (leukaemia), COLO-205	3.36E-05	0.0391
		118286823	115433638				(large intestine)		
Pml	Mad1l1	15:72074067-	7:1821956-	2	2		0 NCI-H1417 (lung), NCI-H322M	3.36E-05	0.0391
		72127204	2239109				(lung)		
Bcl9l	Tbc1d1	11:118272059-	4:37569115-	2	0		2 ATN-1 (leukaemia), COLO-205	3.36E-05	0.0391
		118286823	37817189				(large intestine)		
Dock8	Lrrfip1	9:263048-	2:238200958-	3	8		2 no-11 (glioma), ATN-1 (leukaemia),	4.56E-05	0.0464
		455255	238353697				COLO-205 (large intestine), NCI-		
							H322M (lung)		

Table 5.10. A list of CIS genes that are co-amplified (A) or co-deleted (B) across a significant number of human cancer cell lines. The column "Shared" shows the number of cell lines in which the genes are co-amplified/co-deleted. "Gene 1 unique" and "Gene 2 unique" show the number of cell lines in which gene 1 and gene 2 are amplified/deleted without amplification/deletion of the other gene in the pair. *P*-values were calculated using the 2-tailed Fisher Exact Test, *q*-values were calculated using R package QVALUE.

leukaemias and a range of other tumour types (Katoh and Katoh, 2003), and TBC1D1 has been implicated as an oncogene in gastric cancer (Yang et al., 2007). FAM49A (or D12Ertd553e) is frequently co-amplified with MYCN in neuroblastomas and acute lymphoblastic leukaemias and so is also more likely to play an oncogenic role in cancer (see Section 5.3.2.2). However, DOCK8 and PML are putative tumour suppressor genes (see Sections 5.3.2.3 and 5.5.1, respectively) and TSPAN2 is methylated in breast cancer, suggesting that it may play a tumour suppressive role (Miyamoto et al., 2005), although it is also a potential marker of metastasis in tongue tumours (Carinci et al., 2005). 7 of the co-occurrences involved the leukaemia cell line ATN-1 and colon cancer cell line COLO-205, and 3 of these also involved glioma cell line no-11 and lung cancer cell line NCI-H322M. Each of these cell lines contains a very high number of deletions, i.e. 1228, 1431, 667 and 1284 in cell lines ATN-1, COLO-205, no-11 and NCI-H322M, respectively. They therefore appear to be extremely unstable and the observed cooccurrences may not be functionally important. This is supported by the diversity of cancer types. More interesting is the co-occurrence between deletions involving PML and MAD1L1 (MAD1). Both cell lines are from lung cancers. Lack of expression of PML has been demonstrated in human lung cancers (Gurrieri et al., 2004; Zhang et al., 2000; Zhao et al., 2006). In contrast, amplification of a region containing MAD1L1 is the most commonly observed event in small cell lung cancer cell lines (Coe et al., 2006) and amplified MAD1L1 was significantly over-represented in lung cancer cell lines in Section 5.3.2.2. However, an oncogenic role has not been proven, and *MAD1L1* is more widely implicated as a tumour suppressor gene, e.g. in human stomach cancer (Osaki et al., 2007). Interestingly, PML promotes MAD-mediated transcriptional repression (Khan et al_{2} , 2001), which may contribute to tumour suppression through the repression of MYC(Grandori *et al.*, 2000). However, as the genes are co-deleted, it appears that inactivation of *PML* may not be sufficient to completely abrogate the tumour suppressive functions of MAD. There were no significant co-occurrences of gene pairs in which one gene was amplified and the other was deleted.

All co-occurrences with a *P*-value of less than 0.05 were then compared to cooccurrences with a *P*-value of less than 0.05 within mouse lymphomas to identify candidates that might be conserved across species. It is possible that genes that co-occur within the same amplicon may also co-operate in tumourigenesis (see Section 1.3.3.3) and these were therefore included in the analysis. Gene pairs that co-occurred in both mouse and human are shown in Table 5.11.

1	١
r	1

				Mouse lymphomas										
		Gene 1	Gene 2	Gene 1 Gene 2							Gene 1	Gene 2		
Gene 1	Gene 2	coordinates	coordinates	Shared	unique	unique	P-value	q-value	Shared cell lines	Shared	unique	unique	P-value	q-value
Cldn10a	Rcbtb2	13:94883859-	13:47961104-	3	4	0	9.87E-07	2.91E-04	NCI-H630 (large intestine),	1	7	7	0.0484	0.1514
		95029907	48005317						NCI-H508 (large intestine),					
									COLO-205 (large intestine)					
Rhbdf2	Rnf157	17:71978571-	17:71651451-	2	0	0	5.60E-06	1.35E-03	CP66MEL (skin), MDA-MB-361	1	5	9	0.0445	0.1514
		72009103	71747985						(breast)					
Brd2	Rreb1	6:33044415-	6:7052829-	1	0	1	3.34E-03	0.254	MG-63 (bone)	2	6	29	0.0114	0.1514
		33057260	7197212											
Jup	Ccr7	17:37164382-	17:35963550-	1	2	2	0.0150	0.561	NCI-N87 (stomach)	3	9	50	0.0081	0.1368
		37196476	35975250											
Dym	Fgd2	18:44824172-	6:37081400-	1	3	2	0.0200	0.655	TE-206-1 (unknown)	3	13	58	0.0354	0.1514
		45241077	37489422											
Pecam1	Ncoa3	17:59752964-	20:45564053-	1	3	2	0.0200	0.655	MCF7 (breast)	2	16	17	0.0287	0.1514
		59794505	45719023											
Prkcbp1	Pecam1	20:45271266-	17:59752964-	1	3	3	0.0266	0.781	MCF7 (breast)	2	18	16	0.0319	0.1514
		45418881	59794505											
Ccnd1	Ppp1r10	11:69165054-	6:30676162-	1	22	0	0.0385	1	TE-6 (oesophagus)	2	69	3	0.0136	0.1514
		69178422	30692999											

В

		Human cancer cell lines										Mouse lymphomas					
		Gene 1	Gene 2		Gene 1	Gene 2					Gene 1	Gene 2					
Gene 1	Gene 2	coordinates	coordinates	Shared	unique	unique	P-value	q-value	Shared cell lines	Shared	unique	unique	P-value	q-value			
B230120H23Rik	Ikzf3	2:173648811-	17:35174724-	1	0	0	0.0017	0.0774	NCI-H1417 (lung)	2	10	31	0.0362	0.1514			
		173840979	35273967														
Tcte3	Lef1	6:169882140-	4:109188150-	1	0	2	0.0050	0.1241	NCI-H1417 (lung)	3	11	27	0.0025	0.0987			
		169893563	109309027														
Tbc1d1	Ubash3a	4:37569115-	21:42697088-	1	3	0	0.0067	0.1241	ATN-1 (leukaemia)	2	26	6	0.0092	0.1455			
		37817189	42740843														
Flt3	Ncoa3	13:27475411-	20:45564053-	1	3	0	0.0067	0.1241	NCI-H1417 (lung)	3	32	17	0.0151	0.1514			
		27572729	45719023														
Ubac2	Lef1	13:98651109-	4:109188150-	1	1	2	0.0100	0.1550	NCI-H1417 (lung)	4	27	27	0.0049	0.116			
		98836682	109309027														
Lef1	Mad1l1	4:109188150-	7:1821956-	1	2	1	0.0100	0.1550	NCI-H1417 (lung)	2	27	8	0.0180	0.1514			
		109309027	2239109						(),								

С

		Human cancer cell lines										Mouse lymphomas				
		Gene 1	Gene 2		Gene 1	Gene 2					Gene 1	Gene 2				
Gene 1	Gene 2	coordinates	coordinates	Shared u	unique	unique	P-value	q-value	Shared cell lines	Shared	unique	unique	P-value	q-value		
2010106G01Rik	Tgfbr3	15:48787031-	1:91918488-	1	0	2	0.0050	1 M	CF7 (breast)	1	6	6	0.0357	0.1514		
		48845202	92144147													
Spata13	D12Ertd553e	13:23632887-	2:16594890-	1	0	3	0.006689	1 C(DLO-205 (large intestine)	2	9	14	0.00626	0.1237		
		23779204	16710580						,							
Mylc2pl	Arid1a	7:101043326-	1:26895109-	1	3	1	0.0133	1 N	CI-SNU-5 (stomach)	1	2	11	0.0189	0.1514		
		101059296	26981188													
Flt3	Chst3	13:27475411-	10:73394126-	1	8	0	0.0151	1 C	OLO-205 (large intestine)	1	2	32	0.0313	0.1514		
		27572729	73443318		-	-		-								

Table 5.11. A list of genes that are co-amplified (A), co-deleted (B) or amplified and deleted (C) across human cancer cell lines and are also co-disrupted by MuLV in mouse lymphomas. The columns labelled "Shared" shows the number of cell lines in which the genes are co-amplified/co-deleted and the number of mouse lymphomas in which both genes are disrupted by MuLV. "Gene 1 unique" and "Gene 2 unique" show the number of cell lines in which gene 1 and gene 2 are amplified/deleted without amplification/deletion of the other gene in the pair, and the number of mouse lymphomas in which only one or other gene is disrupted by MuLV. In Figure C, Gene 1 is amplified and Gene 2 is deleted. *P*-values were calculated using the 2-tailed Fisher Exact Test, *q*-values were calculated using R package QVALUE.

8 co-amplified genes in human cancer cell lines were also co-mutated by MuLV in mouse lymphomas. Claudin-10 (*CLDN10*) and *RCBTB2* were co-amplified in 3 cancer cell lines derived from the large intestine but, as discussed in Section 5.3, *RCBTB2* is not a likely target of amplification. The same applies to *RHBDF2* and *RNF157*, which were co-amplified in 2 carcinoma cell lines derived from breast and skin. The minimal amplified region for both genes encompassed 196 genes, of which 4 were known oncogenes, 189 genes were amplified in a greater number of cell lines and 130 were amplified to a higher copy number.

The remaining co-amplifications occurred in a single human cancer cell line, but since they were also co-mutated in MuLV mutagenesis, they are worthy of further investigation. Ras-responsive element binding protein 1 (RREB1) and bromodomain containing 2 (BRD2) were co-amplified in a bone cancer cell line. Both have been previously implicated in cancer but 5 of the 23 genes within the minimal amplified region containing BRD2 were amplified in more cell lines and to greater copy number, suggesting that BRD2 is not a critical target gene. The co-occurrence of genes junction plakoglobin (JUP) and C-C chemokine receptor type 7 precursor (CCR7) is potentially more interesting since, while co-operation between these genes has not been previously observed, both have been implicated in stomach cancer, which is the origin of the cancer cell line in which a co-occurrence was identified. JUP is overexpressed in an amplicon on chromosome 17q that is frequently found in gastric cancer (Varis et al., 2002), while high CCR7 expression correlates with poorer surgical outcomes in gastric cancer patients (Ishigami et al., 2007). These genes showed the highest significance for MuLV cooccurrence but it is worth noting that the q-value was still low, at 0.137. Two of the remaining co-occurrences were in breast cancer cell line MCF7 and involved PECAM1 plus NCOA3 or PRKCBP1. Like TMEM49, mentioned above, PECAM1 resides on the breast-cancer-specific amplicon mapping to chromosome 17q23, while NCOA3 and *PRKCBP1* reside on the 20g breast cancer amplicon. These observations therefore lend further support to the theory that the 17q23 and 20q amplicons co-operate in breast tumourigenesis. The fact that MuLV insertions in PECAM1 also co-occurred with insertions in NCOA3 and PRKCBP1 suggests that PECAM1, rather than TMEM49, may be the critical cancer gene involved in a co-operation with both NCOA3 and PRKCBP1. However, caution must be applied since, while the P-values are less than 0.05, the qvalues for both tests are considerably higher. Finally, CCND1 and PPP1R10 co-occurred in an oesophageal cancer cell line. PPP1R10 inhibits the catalytic subunit of protein phosphatase 1 alpha (PPP1CA) (Kim *et al.*, 2003b). Along with *CCND1*, *PPP1CA* is overexpressed in an amplicon on chromosome 11q13 that is frequently observed in oral squamous cell carcinomas (Hsu *et al.*, 2006) but also in oesophageal squamous cell carcinomas (Huang *et al.*, 2006b). PPP1CA has recently been shown to contribute to oncogenic Ras- and p53-induced cell cycle arrest (Castro *et al.*, 2008). Therefore, it is possible that the amplification of *PPP1R10* counteracts the tumour suppressive effects of *PPP1CA*, which is amplified and overexpressed due to its proximity to *CCND1* and, potentially, other amplified cancer genes. However, this hypothesis clearly needs to be verified experimentally, especially since the overlap between genes is small.

Of the 6 pairs of co-deleted CIS genes, 5 were within the lung cancer cell line NCI-H1417 and 1 was within the leukaemia cell line ATN-1. As mentioned earlier, both cell lines contain a large number of deletions, which reduces the reliability of these associations. However, deletions within at least 3 of the genes have been observed in cancer. For example, heterozygous deletions in MAD1L1 increase the incidence of tumours in mice (Iwanaga et al., 2007), while recurrent deletions in LEF1 and IKZF3 were detected in acute lymphoblastic leukaemias in the study by Mullighan et al. (2007). The 4 co-occurrences of amplified and deleted gene pairs also included 2 that were from a highly unstable cell line, COLO-205. In addition, FLT3 was the only amplified gene for which the minimal amplified region did not contain other genes that were amplified in a greater number of cell lines and, for all associations, the q-value was 1. None of the associations between co-deleted or amplified and deleted genes have been previously observed in the literature, and in all cases, there was only 1 shared human cancer cell line. As all of the observed associations have a low significance, it has not been possible to identify any strong collaborations between cancer genes. However, the co-occurrence of the associations in both human and mouse provides additional evidence, and the gene pairs are therefore presented here as potential candidates for a co-operating role in tumourigenesis.

5.6 Discussion

The purpose of the work described in this chapter was to identify CIS genes that were significantly amplified or deleted in human cancer cell lines. As discussed in Chapter 4, mouse candidate cancer genes identified by retroviral insertional mutagenesis can help to narrow down the candidates in regions of copy number change in the human cancers. In

this chapter, the CIS genes identified in Chapter 2 were used. These are a more reliable set of candidates than those used in Chapter 4. The other major difference between the comparative analysis described in Chapters 4 and 5 was the resolution of the copy number data used. The high-resolution SNP 6.0 data has only recently become available. Due to time constraints and the lack of published methods for dealing with data of this size, the data were subjected to the same segmentation and merging algorithms as the lower resolution, 10K dataset. This may not be entirely appropriate but for the purposes of this study, it was deemed to be sufficient since only the copy numbers at CIS genes were relevant to the analysis, and only those CIS genes that were recurrently amplified or deleted were investigated. The efficacy of the analysis is demonstrated by the overrepresentation of known oncogenes in amplicons, and the identification of implicated tumour suppressor genes in deletions. The high-resolution analysis identified a higher proportion of CIS genes and known cancer genes than the 10K analysis, demonstrating its superiority. It was also shown that some of the most significant candidates identified in the 10K analysis did not appear to be within the true boundaries of regions of copy number change. However, the 10K data may be helpful in filtering out genes that have been incorrectly selected due to errors in the measurement of copy number at clusters of probes or in the calling of gains and losses in the high-resolution data. These problems could also be avoided by using the raw data, rather than the segmentation and merging algorithms, and averaging the copy number values across all markers within a gene. However, this could result in the loss of relevant data in genes where small deletions and amplifications are biologically important. For example, oncogenes NOTCH1 and ETS1 contained intragenic deletions that were helpful in defining the mechanisms of mutation.

Known oncogenes were over-represented among significantly amplified CIS genes, demonstrating that the method used to rank genes was a successful approach for finding promising candidates. Some genes may not be amplified or deleted in a large number of cell lines across all cancer types, but may be heavily implicated in tumourigenesis in a subset of cancers. To identify such tissue-specific candidates, it is essential to perform the analysis independently on each type of cancer. The chosen method ignores genes that are amplified or deleted in few cell lines. However, it is difficult to endorse such candidates unless evidence to support a role in tumourigenesis has been previously demonstrated, as it has for *MEIS1* and *NFKB1*, which were identified in Section 4.5.2.1.1. Nevertheless, a significance threshold of P < 0.05 is potentially too stringent. The *P*-value for each candidate is based on the number of amplifications compared with the

distribution of the number of amplifications for non-CIS genes. Since amplicons are often large and encompass multiple genes, it is highly probable that a number of non-CIS genes will be amplified for every CIS gene, even when the CIS gene is the target of amplification and other genes are passengers. It is therefore unwise to implement a strict cut-off when selecting promising candidates, but the method does provide a way to rank the candidates so that they can be compared against one another. Ranking the candidates according to the maximum copy number, rather than the number of amplicons, provides a method for identifying alternative candidates that, like *MEIS1* and *SLAMF6*, are amplified to high copy number but not across a large number of cell lines.

The search for collaborating cancer genes was somewhat disappointing. Most genes were amplified or deleted in only a small number of cell lines and the power of the analysis was therefore insufficient. In addition, the CIS gene may not be the main target of amplification in all of the cell lines containing a copy number change in that region, which would dilute the association. The lack of an overlap between observed associations and collaborations identified in mouse lymphomas may reflect the fact that pathways and collaborations can differ between tumour types, and associations may therefore be concealed in an analysis of co-occurrences across all human cancer cell lines. However, the numbers, and therefore the power of the analysis, would be too low if each tumour type was independently analysed. The results demonstrate the complexity of human cancer genomes, where different genes are mutated in different tumours and by different mechanisms. This analysis only considers CIS genes in regions of copy number However, genes can also be mutated by epigenetic changes, balanced change. translocations, point mutations and other small, intragenic mutations. Therefore, the analysis is perhaps too restrictive and requires the use of copy number data for a larger set of human cancers and/or datasets representing other types of cancer-associated mutations.

Importantly, CIS genes were over-represented in amplicons and deletions in human cancer cell lines of diverse origin, therefore demonstrating that retroviral insertional mutagenesis in the mouse is relevant to the identification of candidate genes in human cancers derived from a range of tissues. The CIS genes presented in this chapter are stronger candidates than those presented in the gene lists of Chapter 4, and, combined with the high-resolution copy number data for human cancers, they provide a resource of promising candidates for a role in both human and mouse cancer.

Chapter 6 Summary and Conclusions

In light of recent developments in high-throughput genome analysis, genome-wide mutation datasets can be generated for large numbers of cancer genomes at increasing speed and resolution and diminishing cost. However, extracting meaningful information from these datasets can be challenging, particularly as the human and mouse cancer genomes are highly complex, with hundreds of genes being implicated in cancer, and different cancers showing high variability in the spectrum of mutated cancer genes, and in the mechanisms of mutation. The main purpose of this project was to use genome-wide, cancer-associated mutation datasets from mouse and human to facilitate the identification of human genes involved in cancer development.

The principal mouse dataset used in this project was generated by insertional mutagenesis using the retrovirus murine leukaemia virus (MuLV). Retroviral insertional mutagenesis is an established approach for cancer gene discovery in the mouse, but the elucidation of the mouse genome sequence and advances in PCR-based methods for identifying insertion sites have greatly increased the efficiency and the size of the screens that can be performed. The main aim of Chapter 2 was to identify insertion sites and candidate cancer genes within 1,005 MuLV-induced mouse tumours. Following mapping with SSAHA2, the reads, and the insertion sites into which they were clustered, were filtered to remove contaminants. While the filtering procedure is rather conservative and may result in the removal of some true positives, it is essential in large-scale analyses where it would be impractical to manually check each read independently.

An important consideration when planning an insertional mutagenesis screen is how to maximise the coverage, i.e. the proportion of insertion sites successfully identified, whilst keeping control of costs. A comparison of the insertion sites identified in two separate PCR reactions performed on the same tumours demonstrated that the screen is not fully saturated. Without data for greater numbers of enzymes, and for the same enzymes with greater sequencing depth, it was not possible to accurately determine the conditions that would maximise coverage, but it is likely that both the number of enzymes and the depth of sequencing are important factors. Future studies can benefit from next-generation sequencing technologies, such as 454 sequencing (http://www.454.com), which,

compared with the traditional Sanger chain-termination methods, can sequence larger quantities of DNA at lower cost, thereby allowing for an increase in both the number of PCRs and the sequencing depth.

Insertions were assigned to genes by analysing the distribution of insertions around known cancer genes, which are assumed to be targets of mutation. The identification of common insertion sites was performed using two methods: Monte Carlo (MC) simulations (Suzuki et al., 2002) and a kernel convolution (KC)-based framework (de Ridder et al., 2006), and the candidate gene list generated using the KC method was chosen for subsequent analyses following a comparison based on the number of known cancer genes within the MC and KC gene lists. Some cancer-associated genes will almost certainly be missed by both methods. For example, some genes, particularly tumour suppressor genes, can be mutated in a variety of ways, and the insertions may not cluster into a sufficiently tight region to be detected as a single CIS. It is also possible that genes for which insertions are identified in just one or two tumours, and are therefore below the threshold for a significant CIS, do contribute to tumourigenesis. The paucity of insertions may reflect problems in mapping some of the insertions (see above) or may indicate that the gene is not frequently disrupted, e.g. because a specific set of cooperating cancer genes must also be mutated for the gene to contribute to tumourigenesis. However, for the purposes of this study, in which the mouse candidates were used to identify likely candidates within the human dataset, a smaller set of strong candidates is preferable to a larger set containing a high proportion of false positives. A smaller dataset of 73 mouse tumours generated using the Sleeping Beauty (SB) transposon was processed in a similar way to the MuLV dataset.

The main aim of Chapter 3 was to characterise the mouse candidate cancer genes identified in Chapter 2, both as a complete list and as individual candidates. A significant overlap between the mouse candidates and human cancer-associated genes within the Cancer Gene Census (Futreal *et al.*, 2004) and COSMIC database (Forbes *et al.*, 2006) demonstrated the relevance of insertional mutagenesis to human tumourigenesis. This is important on two counts, since it shows that genes disrupted by MuLV mutagenesis contribute to spontaneous cancer development, and that mouse cancer gene candidates are involved in human cancer. Interestingly, candidates were also over-represented among genes with Nanog and Oct4 binding sites, suggesting that a significant proportion may be involved in tumour cell self-renewal, which is consistent with the cancer stem cell

hypothesis discussed in Section 1.2.3.2. The overlap with regions of copy number change in human paediatric acute lymphoblastic leukaemias (Mullighan *et al.*, 2007) provided the first indication that a significant proportion of the mouse candidates may be amplified or deleted in human cancer, and may help to narrow down the candidates in regions of copy number change in human cancers. The analysis also identified miRNA genes among the mouse candidates, and uncovered an over-representation of targets for 3 miRNAs (mmumiR-449b, mmu-miR-449c and hsa-miR-565) that have not been previously implicated in tumourigenesis.

A comparison of the candidates generated by retroviral and transposon-mediated insertional mutagenesis suggested that while some genes are frequently disrupted by both mutagens, others are unique to one screen. This most likely reflects differences in the insertional bias and mechanisms of mutation of the two mutagens. It suggests that the screens are complementary and that the use of multiple mutagens should increase the yield of candidate cancer genes. The implementation of a larger *Sleeping Beauty* screen is therefore highly recommended, while the use of tissue-specific transposons would further increase the repertoire of candidates. It is, however, worth noting that some of the difference between the mutagens may reflect the incomplete saturation of the screens, which results in some insertion sites going undetected. Genes that are mutated by both MuLV and SB are very strong candidates for a role in tumourigenesis since they are unlikely to result from insertional bias, which differs between the two mutagens. The analysis revealed several MuLV- and SB-disrupted genes that warrant further investigation, including p116Rip, Zmiz1, ENSMUSG00000075105 and Osk. Preliminary work has already commenced into the functional validation of OSK, and results have demonstrated that knockdown of the gene in human HeLa cells causes chromosome lagging, which can lead to aneuploidy and cancer formation.

The chapter also focuses on the analysis of the distribution of insertions in and around cancer genes, and this was used to predict the likely mechanism of mutation of candidate genes. For example, *Ccr7* and *Jundm2* were predicted to be mutated by the same mechanism as *Mycn* and *Pim1*, wherein insertions are known to cause premature termination of gene transcription that results in the removal of mRNA-destabilising motifs and, therefore, a more stable gene transcript. This analysis is complicated by the number of ways in which the mutagen can disrupt a gene, and would be aided by using a transposon with a more limited repertoire of mutational mechanisms, e.g. the ability to

induce C-terminal truncations in one orientation only. For genes that are mutated by both MuLV and SB, determination of the mechanism of mutation is facilitated by the coanalysis of MuLV and SB insertions. For example, Notch1 contains both MuLV and SB insertions in 3 distinct locations of the gene that represent different types of oncogenic mutation, and the resulting gene products have been observed in human cancers. Elucidation of the mechanism of mutation may provide an insight into the role of a gene in cancer, and may facilitate the development of therapies targeted against specific mutants. The distribution of insertions can also help to distinguish oncogenes, in which insertions often form distinct clusters, from tumour suppressor genes, in which insertions are more likely to be scattered throughout the gene and may be more likely to include multiple insertions from the same tumour, potentially representing inactivation of both gene copies. As well as a number of known and implicated tumour suppressor genes, this analysis identified Osk, Smg6 and Foxp1 as potential tumour suppressor genes. Interestingly, compared with random insertions, those associated with candidate cancer genes were under-represented in a predicted set of gene-associated regulatory regions and, specifically, within predicted regulatory features associated with active genes. This supports the notion that these insertions are oncogenic and do not simply result from a bias towards insertion into the 5' ends of transcriptionally active genes. It also shows that disruption of regulatory regions is not a common mechanism of mutagenesis for MuLV.

The final part of this chapter was concerned with identifying genes that co-operate in cancer development. The aim of the first analysis was to identify genes that contain an over-representation of insertions in tumours from mice deficient in a particular tumour suppressor gene, therefore suggesting that the genes may collaborate with the loss of that tumour suppressor gene. A number of the significant associations were supported by evidence in the literature, suggesting that many of the unsupported associations may also be real. The same applies to the second analysis, in which pairs of genes that were comutated in a significant number of tumours were identified. Collaborating cancer genes are of particular interest because they can help to elucidate cancer pathways and may represent suitable targets for combined therapies, which may have a lower rate of resistance than therapies targeted to a single gene. Interesting collaborations included positive associations between the loss of *Cdkn2a* and MuLV-disrupted *Zeb2* and *Ccnd1*, and between MuLV-disrupted *Lck* and *A530013C23Rik*, *Stat5b* and *Csk*. The analyses also identified a number of mutually exclusive genes that co-occur less frequently than expected by chance. Such genes may act in the same pathway, and are therefore helpful

in elucidating cancer pathways. Understanding the pathway in which a gene acts may help in the development of cancer therapies since, in some cases, it may be more effective to target a gene that acts downstream of the mutated gene, rather than targeting the mutated gene itself.

In the remaining chapters, mouse candidate cancer genes identified by MuLV insertional mutagenesis were used in cross-species comparative analyses with copy number data from human cancer cell lines. The main aims of the analyses were to demonstrate that retroviral insertional mutagenesis is relevant to the discovery of cancer genes that are amplified or deleted in human cancer, and to help narrow down the candidates within regions of copy number change, which are often large and encompass many genes. In the course of this project, there have been rapid developments in genome-wide copy number analysis and therefore two datasets were used, the second being of much higher resolution and representing the current state of the art. However, even using the lower resolution dataset, it was clear that mouse candidate oncogenes were over-represented in regions of copy number gain and tumour suppressor genes, although less likely to be identified by insertional mutagenesis, were slightly over-represented in regions of copy number loss. The association was observed in cell lines derived from both haematopoietic and lymphoid cancers and solid tumours. Therefore, despite the fact that MuLV insertional mutagenesis generates mouse lymphomas and the resulting candidate cancer genes showed an over-representation of GO terms related to the activation and differentiation of B- and T-cells, the study is relevant to the identification of cancer genes in a diverse range of human cancers. Lymphomas are the most common cancer in mice. As mentioned in Section 1.4.1.2.2, Trp53 is mutated in many types of human cancer, but mutation in the mouse leads to the development of lymphomas or sarcomas (Jonkers and Therefore, simply because mutation in a certain gene leads to Berns, 2002). lymphomagenesis in the mouse, it does not necessarily follow that mutation of the orthologous human gene would result in the same type of cancer. In addition, insertional mutagenesis may result in genes being switched on in cells that would never normally express the gene, even through translocation. Comparison with human cancer-associated mutation data helps to demonstrate how, and whether, the MuLV-disrupted genes contribute to the formation of spontaneous human tumours.

The CGH data from both the 10K and SNP 6.0 SNP arrays were processed into regions of copy number change before being compared to the mouse dataset. Most of the available

methods were designed for processing BAC array CGH data, but the chosen method of DNAcopy (Olshen et al., 2004) plus MergeLevels (Willenbrock and Fridlyand, 2005), was found to be suitable for the 10K SNP array data. The limiting factor was the resolution of the data, since regions of copy number change may be missed or may be falsely predicted. For the higher resolution data, the analysis was limited by the lack of available methods specifically developed for high-density SNP arrays. However, comparative analyses using both datasets uncovered a significant number of known oncogenes in regions of copy number gain, therefore demonstrating the efficacy of the methods. In the lower resolution analysis, interesting candidate oncogenes that were disrupted in the mouse and amplified in human cancer included SLAMF6, MMP13, *RREB1* and *TAOK3*, while candidate tumour suppressor genes included *ARFRP2*, SCFD2, RBMS3, UTRN, ANK3 and ACCN1. Further candidates are provided in Section 4.5.2. Recurrent amplification of *hsa-miR-23a* and deletion of *hsa-miR-128b* were also demonstrated. In the high-resolution analysis, candidate oncogenes included ITPR2, FAM49A, BCL11B, TMEM49, SUPT3H and CLDN10. Candidate tumour suppressor genes included DYM, CYB5, NTN1 and SKI. As anticipated, the high-resolution data appeared to provide a better representation of the cancer genome than did the lower resolution data.

The development of high-resolution platforms for copy number analysis has helped to more accurately define regions of copy number change, but new or modified algorithms are required to deal with the data. The Wellcome Trust Sanger Institute Cancer Genome Project has very recently unveiled a Hidden Markov Model (HMM)-based method, in which data are segmented and are assigned to a state representing the copy number level, which is based on both SNP intensity values and allele ratios (Greenman et al., unpublished). This method analyses each sample independently, and takes account of the genotype at each SNP, making it possible to identify complex, yet fairly common, changes such as loss of heterozygosity without decrease in copy number, and decrease in copy number without loss of heterozygosity. Also in development is a modification of Christiaan Klijn's KC-SMART method (Klijn et al., 2008), in which non-discretised CGH data are input into the program and regions that are significantly aberrant across all tumours are detected, enabling the detection of recurrent amplicons and deletions. Both methods allow for deviation from the 3 states (gain, loss or no change) upon which many previous methods are dependent, therefore permitting more accurate analysis of heterogeneous and polyploid samples.

Chapter 6

Finally, analyses were performed to identify co-amplified and co-deleted candidate cancer genes, and candidate genes that were preferentially amplified or deleted in cell lines containing a somatic mutation in CDKN2A or TP53. There was evidence for cooperation between the breast-cancer-specific amplicons on chromosomes 17q23 and 20q. Other possible associations were also identified, e.g. deletion of PML was found to be both negatively associated with mutation of TP53 and positively associated with deletion of MAD1L1, while JUP and CCR7 were co-amplified. However, the P-values and qvalues for all associations were relatively high, and there was no clear overlap with codisrupted genes in the retroviral screen. The most likely explanation for the lack of significant associations identified by these analyses is that the dataset is simply not large enough. Although 598 cancer cell lines were used, many of these do not have highly unstable genomes, and therefore contain few regions of copy number change. In addition, the cell lines are derived from a variety of tissues and many amplicons and deletions appear to be cell-type-specific, but considering each cancer type independently further reduces the power of the analysis. Finally, there are a number of different mechanisms by which a gene may be disrupted in cancer, and therefore some of the genes that are codisrupted in the MuLV screen may be mutated by another mechanism, e.g. point mutation or hypermethylation, in human cancers.

Copy number changes are only one feature of cancer genomes and, therefore, a greater understanding of cancers and the pathways involved in cancer development could be achieved by integrating additional, complementary, cancer-associated mutation datasets. For example, a more in-depth comparison could be performed between the insertional mutagenesis data and mutations within the COSMIC database (Forbes *et al.*, 2006). In addition, some of the candidates identified in the insertional mutagenesis screen are known to be aberrantly methylated in cancer, e.g. *RASSF2*, *LAPTM5*, *PARK2* and *TSPAN2*, and a comparison with human epigenetic data may reveal further candidates. The human copy number data could also be compared to copy number data generated in the mouse, to identify regions that overlap in both species. In future, when datasets are available for large-scale, tissue-specific insertional mutagenesis screens, it will be possible to compare these with specific types of human cancer to identify tissue-specific cancer genes.

While informatics has taken on an increasingly important role in cancer genomics, it is still essential that candidate cancer genes identified in silico are experimentally validated. This thesis provides a number of strong candidates but their role is not yet proven. To demonstrate a possible involvement in tumourigenesis, the gene of interest can be knocked out or overexpressed in an animal model, such as zebrafish or mouse. Cooperation between cancer genes can be verified by knocking out or overexpressing both genes and comparing the phenotype to single mutants. The creation of a transgenic or endogenous mouse model is a lengthy procedure, and a less protracted method for validating oncogenes involves directly studying the effect on transgenic or chimeric founder mice. However, the use of animal models does not prove that the gene is important in human cancer. As discussed for QSK, the candidate gene can be knocked down in human cells using RNAi (RNA interference) and the effect on cell division and cell growth observed. For candidates that are amplified or deleted in human cancers, expression of those genes can be measured to determine whether there is any significant difference in the expression levels of cancers bearing the copy number change versus those that do not. Candidate genes can also be resequenced across multiple cancers to determine whether they contain somatic mutations in cancer.

The work described in this thesis provides a detailed analysis and characterisation of a large-scale retroviral insertional mutagenesis screen, and demonstrates the relevance of insertional mutagenesis to the discovery of human cancer genes. A selection of strong candidates for a role in tumourigenesis are presented, including mouse candidate cancer genes identified by both MuLV and SB insertional mutagenesis, co-operating mouse cancer genes, and human candidate cancer genes that are both disrupted by MuLV in the mouse and amplified or deleted in human cancers.

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Appendix A. Human Ensembl genes and their mouse orthologues for known cancer genes in the Cancer Gene Census.

Abbreviations:

Tissue type: E=epithelial, L=leukaemia/lymphoma, M=mesenchymal, O=other Dom/Rec: Dom=dominant, Rec=recessive Mutation type: A=amplification, D=large deletion, F=frameshift, Mis=missense, N=nonsense, S=splice site, T=translocation

Human Ensembl ID	Mouse Ensembl ID	Gene name	Somatic mutation	Germline mutation	Tissue type	Dom/ Rec	Mutation type
ENSG00000136754	ENSMUSG0000058835	ABI1	yes		L	Dom	T
ENSG00000097007	ENSMUSG0000026842	ABL1	yes		L	Dom	T, Mis
ENSG00000143322	ENSMUSG0000026596	ABL2	yes		L	Dom	Т
ENSG00000164398	ENSMUSG0000020333	ACSL6	yes		L	Dom	Т
ENSG00000143443	ENSMUSG0000068860	AF1q	yes		L	Dom	Т
ENSG00000172493	ENSMUSG0000029313	AFF1	yes		L	Dom	Т
ENSG00000144218	ENSMUSG0000037138	AFF3	yes		L	Dom	Т
ENSG0000072364	ENSMUSG0000049470	AFF4	yes		L	Dom	т
ENSG00000127914	ENSMUSG0000040407	AKAP9	yes		E	Dom	Ť
ENSG00000105221	ENSMUSG0000004056	AKT2	yes		E	Dom	A
ENSG00000105221	ENSMUSG0000073134	AKT2	yes		E	Dom	А
ENSG00000171094	ENSMUSG0000055471	ALK	yes		Ē	Dom	Т
ENSG00000134982	ENSMUSG0000005871	APC	yes	yes	Е, М, О	Rec	D, Mis, N, F, S
ENSG00000145819	ENSMUSG00000036452	ARHGAP26	yes	yes	1	Dom	T, F, S
ENSG00000196914	ENSMUSG00000059495	ARHGEF12	yes		L	Dom	т
ENSG00000143437	ENSMUSG00000015522	ARNT	yes		Ĺ	Dom	Ť
ENSG00000169696	ENSMUSG00000025142	ASPSCR1	yes		M	Dom	Ť
							Ť
ENSG00000123268	ENSMUSG0000055574	ATF1	yes		Е, М	Dom	
ENSG00000123268	ENSMUSG0000023027	ATF1	yes		Е, М	Dom	T
ENSG00000138363	ENSMUSG0000026192	ATIC	yes		L	Dom	Т
ENSG00000149311	ENSMUSG0000034218	ATM	yes	yes	L, O	Rec	D, Mis, N, F, S
ENSG00000142867	ENSMUSG0000028191	BCL10	yes		L	Dom	Т
ENSG00000119866	ENSMUSG0000000861	BCL11A	yes		L	Dom	Т
ENSG00000127152	ENSMUSG0000048251	BCL11B	yes		L	Dom	Т
ENSG00000171791	ENSMUSG0000057329	BCL2	yes		L	Dom	Т
ENSG00000069399	ENSMUSG0000053175	BCL3	yes		L	Dom	Т
ENSG00000113916	ENSMUSG0000022508	BCL6	yes		L	Dom	T, Mis
ENSG00000110987	ENSMUSG0000029438	BCL7A	yes		L	Dom	Т
ENSG00000116128	ENSMUSG0000038256	BCL9	yes		L	Dom	Т
ENSG00000186716	ENSMUSG0000009681	BCR	yes		L	Dom	Ť
ENSG00000023445	ENSMUSG00000032000	BIRC3	yes		L	Dom	Ť
ENSG00000197299	ENSMUSG0000030528	BLM	yes	VAC		Rec	Mis, N, F
				yes	L, E		
ENSG00000157764	ENSMUSG0000002413	BRAF	yes		E	Dom	Mis, T
ENSG00000012048	ENSMUSG0000017146	BRCA1	yes	yes	E	Rec	D, Mis, N, F, S
ENSG00000139618	ENSMUSG0000041147	BRCA2	yes	yes	L, E	Rec	D, Mis, N, F, S
ENSG00000141867	ENSMUSG0000024002	BRD4	yes		E	Dom	T
ENSG00000136492	ENSMUSG0000034329	BRIP1		yes	L, E	Rec	F, N, Mis
ENSG00000133639	ENSMUSG0000036478	BTG1	yes		L	Dom	Т
ENSG00000156970	ENSMUSG0000040084	BUB1B		yes	М	Rec	Mis, N, F, S
ENSG00000110619	ENSMUSG0000010755	CARS	yes		L	Dom	Т
ENSG00000137812	ENSMUSG0000027326	CASC5	yes		L	Dom	Т
ENSG00000129993	ENSMUSG0000006362	CBFA2T3	yes		L	Dom	Т
ENSG00000067955	ENSMUSG0000031885	CBFB	yes		L	Dom	Т
ENSG00000110395	ENSMUSG0000034342	CBL	yes		L	Dom	Т
ENSG00000100814	ENSMUSG0000071470	CCNB1IP1	yes		Μ	Dom	Т
ENSG00000110092	ENSMUSG0000070348	CCND1	yes		L, E	Dom	Т
ENSG00000118971	ENSMUSG0000000184	CCND2	yes		Ľ	Dom	Т
ENSG00000112576	ENSMUSG0000034165	CCND3	yes		Ĺ	Dom	T
ENSG00000163660	ENSMUSG00000027829	CCNL1	yes		Ē	Dom	Ť
ENSG00000128283	ENSMUSG00000049521	CDC42EP1	yes		L	Dom	Ť
				2005			
ENSG00000134371	ENSMUSG00000026361	CDC73	yes	yes	E, M E	Rec	Mis, N, F Mic N F S
ENSG00000039068	ENSMUSG0000000303	CDH1	yes	yes		Rec	Mis, N, F, S
ENSG00000140937	ENSMUSG00000031673	CDH11	yes	1000	M	Dom	T
ENSG00000135446	ENSMUSG0000006728	CDK4	100	yes	E	Dom	Mis
NSG00000105810	ENSMUSG0000040274	CDK6	yes		L	Dom	Т
ENSG00000147889	ENSMUSG00000044303	CDKN2A	yes	yes	L, E, M, O	Rec	D, S
NSG00000165556	ENSMUSG0000029646	CDX2	yes		L	Dom	Т
NSG00000183765	ENSMUSG0000029521	CHEK2		yes	E	Rec	F
ENSG00000109220	ENSMUSG0000029229	CHIC2	yes		L	Dom	Т
NSG00000128656	ENSMUSG0000056486	CHN1	yes		М	Dom	Т
ENSG00000179583	ENSMUSG0000022504	CIITA	yes		L	Dom	Т
ENSG00000172409	ENSMUSG0000027079	CLP1	yes		L	Dom	Т
ENSG00000141367	ENSMUSG0000047126	CLTC	yes		L	Dom	Т
ENSG00000169714	ENSMUSG0000030057	CNBP	yes		М	Dom	Т
NSG00000108821	ENSMUSG0000001506	COL1A1	yes		M	Dom	Ť
ENSG00000164919	ENSMUSG0000066491	COX6C	yes		M	Dom	T
ENSG00000164919	ENSMUSG00000014313	COX6C	yes		M	Dom	Ť
ENSG00000164919	ENSMUSG00000069096	COX6C	yes		M	Dom	Ť
ENSG00000118260	ENSMUSG00000025958	CREB1			M	Dom	Ť
ENSG00000118280		CREBI	yes				Ť
	ENSMUSG0000022521		yes		L	Dom	
ENSG00000105662	ENSMUSG0000003575	CRTC1	yes		E	Dom	T
ENSG00000168036	ENSMUSG0000006932	CTNNB1	yes		Е, М, О	Dom	H, Mis
ENSG00000144476	ENSMUSG0000044337	CXCR7	yes		M	Dom	T
ENSG00000138336	ENSMUSG0000075011	CXXC6	yes		L	Dom	Т
ENSG00000083799	ENSMUSG0000036712	CYLD	yes	yes	E	Rec	Mis, N, F, S
ENSG00000108091	ENSMUSG0000048701	D10S170	yes		E	Dom	Т
ENSG00000134574	ENSMUSG0000002109	DDB2		yes	E	Rec	Mis, N
	ENSMUSG0000025408	DDIT3	yes		M	Dom	Т

ENSG00000178105	ENSMUSG0000053289	DDX10	yes		L	Dom	т
ENSG00000110367	ENSMUSG0000059707	DDX6	yes		L	Dom	Т
ENSG00000110367	ENSMUSG0000032097	DDX6	yes		Ĺ	Dom	T
ENSG00000124795	ENSMUSG00000021377	DEK			Ĺ	Dom	Ť
			yes				
ENSG00000146648	ENSMUSG00000020122	EGFR	yes		Ε, Ο	Dom	A, O, Mis
ENSG00000156976	ENSMUSG0000022884	EIF4A2	yes		L	Dom	Т
ENSG00000156976	ENSMUSG0000022884	EIF4A2	yes		L	Dom	Т
ENSG00000102034	ENSMUSG0000031103	ELF4	yes		E	Dom	Т
ENSG00000105656	ENSMUSG0000070002	ELL	yes		Ĺ	Dom	T
							Ť
ENSG00000100393	ENSMUSG00000055024	EP300	yes		L, E	Rec	
ENSG0000085832	ENSMUSG0000028552	EPS15	yes		L	Dom	Т
ENSG00000141736	ENSMUSG0000062312	ERBB2	yes		E	Dom	A, Mis, O
ENSG00000104884	ENSMUSG0000030400	ERCC2		yes	E	Rec	Mis, N, F, S
ENSG00000163161	ENSMUSG0000024382	ERCC3		yes	E	Rec	Mis, S
ENSG00000175595	ENSMUSG0000022545	ERCC4		yes	Ē	Rec	Mis, N, F
	ENSMUSG0000026048	ERCC5					
ENSG00000134899				yes	E	Rec	Mis, N, F
ENSG00000157554	ENSMUSG0000040732	ERG	yes		M, E, L	Dom	Т
ENSG0000006468	ENSMUSG0000047643	ETV1	yes		М, Е	Dom	Т
ENSG0000006468	ENSMUSG0000004151	ETV1	yes		М, Е	Dom	Т
ENSG00000175832	ENSMUSG0000017724	ETV4	yes		M	Dom	Т
ENSG00000139083	ENSMUSG0000030199	ETV6	,		L, E, M	Dom	Ť
			yes				
ENSG0000085276	ENSMUSG0000027684	EVI1	yes		L	Dom	T
ENSG00000182944	ENSMUSG0000038649	EWSR1	yes		L, M	Dom	Т
ENSG00000182944	ENSMUSG0000070829	EWSR1	yes		L, M	Dom	Т
ENSG00000182944	ENSMUSG0000009079	EWSR1	yes		L, M	Dom	Т
ENSG00000182197	ENSMUSG0000061731	EXT1	,	VAC	<u>м</u>	Rec	Mis, N, F, S
				yes		_	
ENSG00000151348	ENSMUSG0000027198	EXT2		yes	M	Rec	Mis, N, F, S
ENSG00000187741	ENSMUSG0000032815	FANCA		yes	L	Rec	D, Mis, N, F, S
ENSG00000158169	ENSMUSG0000021461	FANCC		yes	L	Rec	D, Mis, N, F, S
ENSG00000144554	ENSMUSG0000034023	FANCD2		yes	Ĺ	Rec	D, Mis, N, F
		FANCE					
ENSG00000112039	ENSMUSG0000007570			yes	L	Rec	N, F, S
ENSG00000183161	ENSMUSG0000043480	FANCF		yes	L	Rec	N, F
ENSG00000165281	ENSMUSG0000028453	FANCG		yes	L	Rec	Mis, N, F, S
ENSG0000026103	ENSMUSG0000024778	FAS	yes		L, E, O	Rec	Mis
ENSG00000109670	ENSMUSG00000028086	FBXW7	,		E, 2, 0	Dom	Mis, N
			yes				
ENSG0000072694	ENSMUSG0000026656	FCGR2B	yes		L	Dom	Т
ENSG0000072694	ENSMUSG0000059498	FCGR2B	yes		L	Dom	Т
ENSG00000163497	ENSMUSG0000055197	FEV	yes		M	Dom	Т
ENSG0000077782	ENSMUSG0000031565	FGFR1	yes		L	Dom	Т
	ENSMUSG00000069135	FGFR1OP			Ĺ		Ť
ENSG00000112486			yes			Dom	
ENSG0000066468	ENSMUSG0000030849	FGFR2	yes		E	Dom	Mis
ENSG0000068078	ENSMUSG0000054252	FGFR3	yes		L, E	Dom	Mis, T
ENSG0000091483	ENSMUSG0000026526	FH		yes	Е, М	Rec	Mis, N, F
ENSG0000091483	ENSMUSG0000037498	FH		yes	E, M	Rec	Mis, N, F
				ycs			
ENSG00000145216	ENSMUSG0000029227	FIP1L1	yes		L	Dom	T
ENSG00000154803	ENSMUSG0000032633	FLCN		yes	Е, М	Rec?	Mis. N, F
ENSG00000122025	ENSMUSG0000042817	FLT3	yes		L	Dom	Mis, O
ENSG00000187239	ENSMUSG0000075415	FNBP1	yes		L	Dom	Т
ENSG00000150907	ENSMUSG0000044167	FOXO1A	yes		M	Dom	Ť
ENSG00000118689	ENSMUSG0000048756	FOXO3A	yes		L	Dom	T
ENSG00000070404	ENSMUSG0000020325	FSTL3	yes		L	Dom	Т
ENSG0000089280	ENSMUSG0000066554	FUS	yes		M, L	Dom	т
ENSG0000089280	ENSMUSG0000030795	FUS	yes		M, L	Dom	Т
ENSG00000119537	ENSMUSG0000009905	FVT1			L., 2	Dom	Ť
			yes				
ENSG0000007237	ENSMUSG0000033066	GAS7	yes		L	Dom	T _
ENSG00000102145	ENSMUSG0000031162	GATA1	yes		L	Dom	Mis, F
ENSG00000163655	ENSMUSG0000027823	GMPS	yes		L	Dom	Т
ENSG0000087460	ENSMUSG0000027523	GNAS	yes		E	Dom	Mis
ENSG0000066455	ENSMUSG0000021192	GOLGA5			E	Dom	Т
			yes				1
ENSG00000047932							0
	ENSMUSG0000019861	GOPC	yes		0	Dom	0
		GUPL	yes			Dom X-linked	T, D, Mis, N, F,
ENSG00000147257	ENSMUSG00000019861	GOPC GPC3	yes	yes	0		
	ENSMUSG00000055653	GPC3		yes	0	X-linked Rec	T, D, Mis, N, F, S
ENSG00000171723	ENSMUSG00000055653 ENSMUSG00000047454	GPC3 GPHN	yes	yes	O L	X-linked Rec Dom	T, D, Mis, N, F, S T
ENSG00000171723 ENSG00000127946	ENSMUSG00000055653 ENSMUSG00000047454 ENSMUSG00000039959	GPC3 GPHN HIP1	yes yes	yes	O L L	X-linked Rec Dom Dom	T, D, Mis, N, F, S T T
ENSG00000171723 ENSG00000127946 ENSG00000198339	ENSMUSG00000055653 ENSMUSG00000047454 ENSMUSG00000039959 ENSMUSG00000069305	GPC3 GPHN HIP1 HIST1H4I	yes yes yes	yes	O L L L	X-linked Rec Dom Dom Dom	T, D, Mis, N, F, S T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG00000108924	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000069305 ENSMUSG0000003949	GPC3 GPHN HIP1 HIST1H4I HLF	yes yes yes yes	yes	O L L L	X-linked Rec Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339	ENSMUSG00000055653 ENSMUSG00000047454 ENSMUSG00000039959 ENSMUSG00000069305	GPC3 GPHN HIP1 HIST1H4I	yes yes yes	yes	O L L L	X-linked Rec Dom Dom Dom	T, D, Mis, N, F, S T T T T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG00000108924	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000069305 ENSMUSG0000003949	GPC3 GPHN HIP1 HIST1H4I HLF	yes yes yes yes	yes	O L L L	X-linked Rec Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG00000108924 ENSG00000130675 ENSG00000149948	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG0000039959 ENSMUSG0000069305 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG00000056758	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2	yes yes yes yes yes yes	yes	O L L L M	X-linked Rec Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG00000108924 ENSG00000130675 ENSG00000149948 ENSG0000005073	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG0000039959 ENSMUSG0000003955 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG00000056758 ENSMUSG00000038210	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11	yes yes yes yes yes yes yes	yes	O L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG000018924 ENSG0000130675 ENSG0000149948 ENSG0000005073 ENSG00000106031	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG0000039959 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG00000056758 ENSMUSG0000038210 ENSMUSG0000038203	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13	yes yes yes yes yes yes yes yes	yes	0 L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T
ENSG00000171723 ENSG00000127946 ENSG0000198339 ENSG000018924 ENSG00000130675 ENSG00000149948 ENSG0000005073 ENSG00000106031 ENSG00000078399	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG00000069305 ENSMUSG0000001566 ENSMUSG00000056758 ENSMUSG0000038203 ENSMUSG0000038227	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9	yes yes yes yes yes yes yes yes yes	yes	0 L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG0000108924 ENSG0000130675 ENSG0000149948 ENSG0000005073 ENSG0000016031 ENSG0000078399 ENSG00000123388	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG0000039959 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG00000056758 ENSMUSG0000038210 ENSMUSG0000038203	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13	yes yes yes yes yes yes yes yes	yes	0 L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG0000108924 ENSG0000130675 ENSG0000149948 ENSG0000005073 ENSG0000016031 ENSG0000078399 ENSG00000123388	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG0000039959 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG0000001566 ENSMUSG0000038210 ENSMUSG00000382203 ENSMUSG0000038227 ENSMUSG00000038227	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9 HOXC11	yes yes yes yes yes yes yes yes yes yes	yes	0 L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T
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ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG000019834 ENSG0000130675 ENSG0000016031 ENSG0000005073 ENSG00000123388 ENSG00000123384 ENSG0000128713 ENSG0000128714 ENSG0000128714 ENSG000013522 ENSG0000137265 ENSG0000137265 ENSG00000137265 ENSG0000013814 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404	ENSMUSG0000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG00000039459 ENSMUSG0000003949 ENSMUSG0000003820 ENSMUSG0000038210 ENSMUSG0000038203 ENSMUSG00000038203 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001654 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000024789 ENSMUSG00000023652 ENSMUSG00000035672 ENSMUSG000000032652 ENSMUSG0000000078 ENSMUSG0000000078 ENSMUSG0000000078 ENSMUSG000000021843	GPC3 GPHN HIP1 HIST1H4I HLXB9 HMGA2 HOXA11 HOXA13 HOXA13 HOXC11 HOXC11 HOXC13 HOXD11 HOXD13 HRAS IKZF1 IL2 IL21R IKF1 IL2 IL21R IRF4 ITK JAZF1 KIT KIF6 KRAS KTN1	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L L L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T, Mis, O T T T, Mis, O Mis, N Mis T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG0000198324 ENSG0000149948 ENSG00000149948 ENSG0000005073 ENSG00000123388 ENSG0000123384 ENSG0000128713 ENSG0000128714 ENSG00000128714 ENSG000001785811 ENSG0000019471 ENSG0000019471 ENSG0000013522 ENSG0000013265 ENSG0000013265 ENSG0000013814 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000133703 ENSG00000126777 ENSG00000126777 ENSG0000012877	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000001566 ENSMUSG00000038203 ENSMUSG00000038203 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001854 ENSMUSG0000001854 ENSMUSG0000001356 ENSMUSG0000001356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG00000021356 ENSMUSG0000002672 ENSMUSG0000002672 ENSMUSG00000021843 ENSMUSG000000218454	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXC11 HOXC13 HOXC11 HOXC13 HOXD11 HOXD13 HRA5 IL2 IL2 IL2 IL2 IL2 IL2 IL2 IL2 IL2 IL2	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L L L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T, Mis, O T T T, Mis, O Mis, N Mis T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG0000019834 ENSG0000018924 ENSG0000018924 ENSG0000016031 ENSG0000016031 ENSG00000123388 ENSG00000123364 ENSG00000128713 ENSG00000128714 ENSG00000185811 ENSG00000185811 ENSG00000185811 ENSG0000013522 ENSG0000013522 ENSG0000013522 ENSG00000153814 ENSG00000153814 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000126777 ENSG00000126777 ENSG00000126777 ENSG000002834 ENSG000002834	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000003210 ENSMUSG00000038210 ENSMUSG00000038207 ENSMUSG00000038227 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001819 ENSMUSG00000018654 ENSMUSG00000018654 ENSMUSG00000013566 ENSMUSG0000002395 ENSMUSG0000002375 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000003568 ENSMUSG0000003568 ENSMUSG0000003656 ENSMUSG00000038266 ENSMUSG00000038265 ENSMUSG00000021843 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXC13 HOXC13 HOXC13 HOXD13 HOXD13 HAAS IKZF1 IL2 IL21R IRF4 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LASP1 LASP1	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG000019834 ENSG000018924 ENSG0000130675 ENSG0000016031 ENSG0000016031 ENSG00000123388 ENSG0000123364 ENSG0000128713 ENSG0000128713 ENSG0000128714 ENSG0000128714 ENSG000013522 ENSG000013522 ENSG000013522 ENSG000013522 ENSG000013523 ENSG00000153814 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000126777 ENSG0000012874 ENSG000002834 ENSG000002834 ENSG000002834 ENSG000002834	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG00000039459 ENSMUSG00000039459 ENSMUSG00000038210 ENSMUSG00000038210 ENSMUSG00000038210 ENSMUSG00000038227 ENSMUSG00000038227 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001854 ENSMUSG0000001854 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025495 ENSMUSG0000002395 ENSMUSG00000023752 ENSMUSG0000003566 ENSMUSG0000003765 ENSMUSG0000003366 ENSMUSG00000037792 ENSMUSG0000003366 ENSMUSG00000037792 ENSMUSG00000037692 ENSMUSG00000037792 ENSMUSG0000003366	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9 HOXC11 HOXC13 HOXD11 HOXC13 HOXD13 HRAS IKZF1 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LCK	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T, Mis, O T T T, Mis, O Mis, N Mis T T T T T T T T T T T T T T T T T T T
ENSG00000171723 ENSG00000127946 ENSG00000198339 ENSG0000019834 ENSG0000018924 ENSG0000018924 ENSG0000016031 ENSG0000016031 ENSG00000123388 ENSG00000123364 ENSG00000128713 ENSG00000128714 ENSG00000185811 ENSG00000185811 ENSG00000185811 ENSG0000013522 ENSG0000013522 ENSG0000013522 ENSG00000153814 ENSG00000153814 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000126777 ENSG00000126777 ENSG00000126777 ENSG000002834 ENSG000002834	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000003210 ENSMUSG00000038210 ENSMUSG00000038207 ENSMUSG00000038227 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001819 ENSMUSG00000018654 ENSMUSG00000018654 ENSMUSG00000013566 ENSMUSG0000002395 ENSMUSG0000002375 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000002378 ENSMUSG0000003568 ENSMUSG0000003568 ENSMUSG0000003656 ENSMUSG00000038266 ENSMUSG00000038265 ENSMUSG00000021843 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXC13 HOXC13 HOXC13 HOXD13 HOXD13 HAAS IKZF1 IL2 IL21R IRF4 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LASP1 LASP1	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG000019834 ENSG000018924 ENSG0000130675 ENSG0000016031 ENSG0000016031 ENSG00000123388 ENSG0000123364 ENSG0000128713 ENSG0000128713 ENSG0000128714 ENSG0000128714 ENSG000013522 ENSG000013522 ENSG000013522 ENSG000013522 ENSG000013523 ENSG00000153814 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000157404 ENSG00000126777 ENSG0000012874 ENSG000002834 ENSG000002834 ENSG000002834 ENSG000002834	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG00000039459 ENSMUSG00000039459 ENSMUSG00000038210 ENSMUSG00000038210 ENSMUSG00000038210 ENSMUSG00000038227 ENSMUSG00000038227 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001854 ENSMUSG0000001854 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000025495 ENSMUSG0000002395 ENSMUSG00000023752 ENSMUSG0000003566 ENSMUSG0000003765 ENSMUSG0000003366 ENSMUSG00000037792 ENSMUSG0000003366 ENSMUSG00000037792 ENSMUSG00000037692 ENSMUSG00000037792 ENSMUSG0000003366	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9 HOXC11 HOXC13 HOXD11 HOXC13 HOXD13 HRAS IKZF1 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LCK	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T, Mis, O T T T, Mis, O Mis, N Mis T T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000127946 ENSG000018339 ENSG000018924 ENSG0000130675 ENSG0000016031 ENSG00000078399 ENSG00000123388 ENSG0000123384 ENSG0000128713 ENSG0000128714 ENSG0000128714 ENSG0000128714 ENSG000019471 ENSG000019471 ENSG0000013522 ENSG0000013265 ENSG0000013265 ENSG0000013814 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000126777 ENSG000001287404 ENSG0000012834 ENSG000002834 ENSG000002834 ENSG0000012866 ENSG00000136167 ENSG00000183722	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000003201 ENSMUSG00000038203 ENSMUSG00000038203 ENSMUSG00000038203 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001854 ENSMUSG0000001854 ENSMUSG0000001855 ENSMUSG0000001855 ENSMUSG0000001855 ENSMUSG0000001855 ENSMUSG0000001855 ENSMUSG00000023750 ENSMUSG0000002375 ENSMUSG0000002375 ENSMUSG0000002365 ENSMUSG0000005672 ENSMUSG0000003866 ENSMUSG0000005672 ENSMUSG0000003866 ENSMUSG0000003866 ENSMUSG00000038366 ENSMUSG0000003866 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000038366 ENSMUSG00000021998	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXC11 HOXC13 HOXC13 HOXC11 HOXC13 HOXD13 HCXD11 HOXD13 HRA5 IKZF1 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LASP1 LASP1 LCK LCP1 LCF1 LCF1 LCF1	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L M M, O E, E, L L L L L M M	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG0000198924 ENSG0000018924 ENSG0000018924 ENSG0000018924 ENSG0000016031 ENSG00000123388 ENSG0000123364 ENSG0000128713 ENSG00000128714 ENSG00000128714 ENSG00000185811 ENSG00000185811 ENSG00000185811 ENSG0000013522 ENSG0000013522 ENSG0000013523 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000153703 ENSG00000126777 ENSG00000126777 ENSG000002834 ENSG000002834 ENSG00000182866 ENSG00000182866 ENSG00000183722 ENSG0000013572	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000003210 ENSMUSG00000038210 ENSMUSG00000038217 ENSMUSG00000038227 ENSMUSG0000001656 ENSMUSG0000001656 ENSMUSG0000001819 ENSMUSG00000018654 ENSMUSG00000018654 ENSMUSG00000027720 ENSMUSG00000018654 ENSMUSG00000027720 ENSMUSG00000027720 ENSMUSG00000027720 ENSMUSG0000002375 ENSMUSG00000023765 ENSMUSG0000002365 ENSMUSG00000025672 ENSMUSG00000003866 ENSMUSG00000021843 ENSMUSG0000003866 ENSMUSG00000021843 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000048332	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9 HOXC11 HOXC13 HOXD13 HRA5 IKZF1 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LCK LCP1 LHFP LIFR	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG000019834 ENSG0000130675 ENSG00000149948 ENSG0000005073 ENSG0000016031 ENSG00000123364 ENSG0000123364 ENSG0000128713 ENSG0000128713 ENSG0000128714 ENSG0000137265 ENSG000013522 ENSG0000137265 ENSG00000137265 ENSG00000137265 ENSG0000013814 ENSG0000013814 ENSG0000013703 ENSG00000128777 ENSG00000128777 ENSG00000128777 ENSG0000012834 ENSG0000012834 ENSG0000012834 ENSG0000012834 ENSG0000012834 ENSG0000012834 ENSG000012834 ENSG000012834 ENSG000012834 ENSG000012834 ENSG000012834 ENSG000012834 ENSG0000128722 ENSG000012872	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG00000039499 ENSMUSG0000003949 ENSMUSG00000038210 ENSMUSG0000038210 ENSMUSG0000038203 ENSMUSG00000038203 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG0000001655 ENSMUSG00000018549 ENSMUSG00000025499 ENSMUSG00000025499 ENSMUSG00000027720 ENSMUSG00000023459 ENSMUSG0000002395 ENSMUSG0000002395 ENSMUSG0000002375 ENSMUSG00000024789 ENSMUSG00000024789 ENSMUSG00000024789 ENSMUSG00000024789 ENSMUSG00000024789 ENSMUSG00000024783 ENSMUSG00000021843 ENSMUSG00000021843 ENSMUSG0000003366 ENSMUSG00000037792 ENSMUSG00000048532 ENSMUSG0000004832	GPC3 GPHN HIP1 HIST1H4I HLXB9 HMGA2 HOXA11 HOXA13 HOXA3 HOXC11 HOXC13 HOXD11 HOXC13 HCXC13 HOXD11 HOXC13 HC	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L L L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T
ENSG0000171723 ENSG0000127946 ENSG0000198339 ENSG0000198924 ENSG0000018924 ENSG0000018924 ENSG0000018924 ENSG0000016031 ENSG00000123388 ENSG0000123364 ENSG0000128713 ENSG00000128714 ENSG00000128714 ENSG00000185811 ENSG00000185811 ENSG00000185811 ENSG0000013522 ENSG0000013522 ENSG0000013523 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000153814 ENSG00000153703 ENSG00000126777 ENSG00000126777 ENSG000002834 ENSG000002834 ENSG00000182866 ENSG00000182866 ENSG00000183722 ENSG0000013572	ENSMUSG00000055653 ENSMUSG0000047454 ENSMUSG00000039959 ENSMUSG0000003949 ENSMUSG0000003949 ENSMUSG0000003210 ENSMUSG00000038210 ENSMUSG00000038217 ENSMUSG00000038227 ENSMUSG0000001656 ENSMUSG0000001656 ENSMUSG0000001819 ENSMUSG00000018654 ENSMUSG00000018654 ENSMUSG00000027720 ENSMUSG00000018654 ENSMUSG00000027720 ENSMUSG00000027720 ENSMUSG00000027720 ENSMUSG0000002375 ENSMUSG00000023765 ENSMUSG0000002365 ENSMUSG00000025672 ENSMUSG00000003866 ENSMUSG00000021843 ENSMUSG0000003866 ENSMUSG00000021843 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000021998 ENSMUSG00000048332	GPC3 GPHN HIP1 HIST1H4I HLF HLXB9 HMGA2 HOXA11 HOXA13 HOXA9 HOXC11 HOXC13 HOXD13 HRA5 IKZF1 IL2 IL21R IRF4 ITK JAK2 JAZF1 KIT KLF6 KRAS KTN1 LASP1 LCK LCP1 LHFP LIFR	yes yes yes yes yes yes yes yes yes yes	yes	O L L L L M L L L L L L L L L L L L L L	X-linked Rec Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	T, D, Mis, N, F, S T T T T T T T T T T T T T T T T T T

ENSG00000145012							
	ENSMUSG0000033306	LPP	yes		L, M	Dom	т
ENSG00000104903	ENSMUSG0000034041	LYL1	yes		L	Dom	Т
ENSG00000178573	ENSMUSG0000055435	MAF	yes		L	Dom	Т
ENSG00000204103	ENSMUSG00000074622	MAFB			Ĺ	Dom	Ť
			yes				
ENSG00000172175	ENSMUSG0000032688	MALT1	yes		L	Dom	T
ENSG00000184384	ENSMUSG0000031925	MAML2	yes		E	Dom	Т
ENSG0000065559	ENSMUSG0000033352	MAP2K4	yes		E	Rec	D, Mis, N
ENSG00000206115	ENSMUSG0000051636	MDS1	yes		Ē	Dom	Т
ENSG00000133895	ENSMUSG0000024947	MEN1	yes	yes	E	Rec	D, Mis, N, F, S
ENSG00000105976	ENSMUSG0000009376	MET	yes		E	Dom	Mis
ENSG00000162775	ENSMUSG0000048109	MKL1	yes		L	Dom	т
ENSG00000178053	ENSMUSG0000048416	MLF1	yes		L	Dom	т
ENSG0000076242	ENSMUSG0000032498	MLH1	yes	yes	Ε, Ο	Rec	D, Mis, N, F, S
ENSG00000118058	ENSMUSG0000002028	MLL	yes		L	Dom	т, о
ENSG00000130382	ENSMUSG0000024212	MLLT1	yes		L	Dom	Т
ENSG0000078403	ENSMUSG0000026743	MLLT10	yes		L	Dom	т
ENSG00000171843	ENSMUSG0000028496	MLLT3			Ĺ	Dom	Ť
			yes				
ENSG00000130396	ENSMUSG0000068036	MLLT4	yes		L	Dom	Т
ENSG00000108292	ENSMUSG0000038437	MLLT6	yes		L	Dom	Т
ENSG00000184481	ENSMUSG0000042903	MLLT7	yes		L	Dom	Т
ENSG00000169184	ENSMUSG0000070576	MN1	yes		L, O	Dom	Т
ENSG00000117400	ENSMUSG0000006389	MPL	yes	yes	L	Dom	Mis
ENSG0000095002	ENSMUSG0000024151	MSH2	yes	yes	E	Rec	D, Mis, N, F, S
ENSG00000116062	ENSMUSG0000005370	MSH6	yes	yes	E	Rec	Mis, N, F, S
ENSG00000153944	ENSMUSG0000069769	MSI2	yes	,	L	Dom	т
ENSG00000147065	ENSMUSG0000031207	MSN			Ĺ	Dom	Ť
			yes				
ENSG00000182712	ENSMUSG0000031200	MTCP1	yes		L	Dom	Т
ENSG00000185499	ENSMUSG0000042784	MUC1	yes		L	Dom	Т
ENSG00000132781	ENSMUSG0000028687	MUTYH		yes	E	Rec	Mis
ENSG00000136997	ENSMUSG00000022346	MYC	VAC	, 65	 L, Е	Dom	А, Т
			yes				
ENSG00000116990	ENSMUSG0000028654	MYCL1	yes		E	Dom	A
ENSG00000134323	ENSMUSG0000037169	MYCN	yes		0	Dom	A
ENSG00000133392	ENSMUSG0000018830	MYH11	yes		L	Dom	т
ENSG00000100345	ENSMUSG0000022443	MYH9			Ĺ	Dom	T
			yes				
ENSG0000083168	ENSMUSG0000031540	MYST3	yes		L	Dom	Т
ENSG00000156650	ENSMUSG0000021767	MYST4	yes		L	Dom	Т
ENSG00000104320	ENSMUSG0000028224	NBN		yes	L, E, M, O	Rec	Mis, N, F
ENSG00000140396	ENSMUSG0000005886	NCOA2	VAC	/	L , , , , , ,	Dom	Т
			yes				
ENSG00000138293	ENSMUSG0000021908	NCOA4	yes		E	Dom	T
ENSG00000138293	ENSMUSG0000056234	NCOA4	yes		E	Dom	т
							D, Mis, N, F, S,
ENSG00000196712	ENSMUSG0000020716	NF1	yes	yes	0	Rec	0
21000000190712	2103110300000020710	101 1	,00	,05	0	nee	
							D, Mis, N, F, S,
ENSG00000186575	ENSMUSG0000009073	NF2	yes	yes	0	Rec	0
ENSG0000077150	ENSMUSG0000025225	NFKB2	yes		L	Dom	т
ENSG00000100503	ENSMUSG0000021068	NIN	yes		L	Dom	т
ENSG00000147140	ENSMUSG0000067514	NONO	yes		E	Dom	Ť
ENSG00000147140	ENSMUSG0000031311	NONO	yes		E	Dom	T
ENSG00000148400	ENSMUSG0000026923	NOTCH1	yes		L	Dom	T, Mis, O
ENSG00000119508	ENSMUSG0000028341	NR4A3	yes		M	Dom	Т
ENSG0000009307	ENSMUSG0000072420	NRAS	yes		L, E	Dom	Mis
	ENSMUSG00000068823	NRAS					
ENSG0000009307			yes				
					L, E	Dom	Mis
ENSG0000009307	ENSMUSG00000047588	NRAS	yes		L, E L, E	Dom	Mis
ENSG0000009307 ENSG00000165671			yes				
ENSG00000165671	ENSMUSG00000047588 ENSMUSG00000021488	NRAS NSD1	yes yes		L, E L	Dom Dom	Mis T
ENSG00000165671 ENSG00000198400	ENSMUSG00000047588 ENSMUSG00000021488 ENSMUSG00000028072	NRAS NSD1 NTRK1	yes yes yes		L, E L E	Dom Dom Dom	Mis T T
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ENSG0000165671 ENSG0000198400 ENSG0000137497 ENSG0000110713 ENSG0000110713 ENSG00001205927 ENSG0000127083 ENSG00000127083 ENSG00000168092 ENSG00000135903 ENSG0000101015 ENSG00001035903 ENSG0000125618 ENSG00000125618 ENSG00000185630 ENSG00000185630 ENSG00000185630 ENSG00000185630 ENSG00000185630 ENSG00000185633 ENSG00000185633 ENSG00000185633 ENSG0000013211 ENSG0000013212 ENSG00000121879 ENSG00000121879 ENSG00000121879 ENSG00000121879 ENSG00000121879 ENSG00000121879 ENSG00000132170 ENSG00000140464 ENSG0000014024511 ENSG00000143294 ENSG00000142611	ENSMUSG0000047588 ENSMUSG0000028072 ENSMUSG0000028072 ENSMUSG0000006306 ENSMUSG00000063550 ENSMUSG00000041358 ENSMUSG00000041358 ENSMUSG00000041358 ENSMUSG00000048702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG0000004872 ENSMUSG0000004872 ENSMUSG00000026574 ENSMUSG00000026976 ENSMUSG00000026976 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG00000024620 ENSMUSG0000024620 ENSMUSG00000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG0000024620 ENSMUSG00000024046 ENSMUSG00000024046 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG00000024406 ENSMUSG0000026586 ENSMUSG0000026586 ENSMUSG00000026586	NRAS NSD1 NTRK1 NTRK3 NUMA1 NUP98 NUT OLIG2 OMD PAFAH1B2 PALB2 PALB2 PALB2 PAT21 PAX3 PAX5 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7	yes yes yes yes yes yes yes yes yes yes	yes	L, E E E, M L L E L M L L, M M L M E L E L L M L, L O L E, L E L M E E L M M, O E E O O E L E L M E E L M E E L M E E L E, L L E L M E E L E, L L	Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	Mis T T T T T T T T T T T T T T T T T Mis, O, T T T T Mis, F T Mis, F T T Mis, N T T T T T T T T T T T T T T T T T T T
ENSG0000165671 ENSG00001498400 ENSG0000149538 ENSG0000110713 ENSG0000110713 ENSG00001205927 ENSG0000127083 ENSG0000127083 ENSG0000168092 ENSG0000101015 ENSG000010105 ENSG0000105092 ENSG0000105092 ENSG00000125618 ENSG00000125618 ENSG00000125618 ENSG0000018601 ENSG0000018164 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000142611 ENSG00000142611 ENSG00000164985 ENSG00000164985	ENSMUSG0000047588 ENSMUSG0000028072 ENSMUSG0000028072 ENSMUSG00000059146 ENSMUSG0000006306 ENSMUSG00000039800 ENSMUSG00000013530 ENSMUSG00000048768 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG0000004872 ENSMUSG0000004872 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG0000002893 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024614 ENSMUSG00000024014 ENSMUSG0000002828 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024016 ENSMUSG00000024016 ENSMUSG00000024016 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024101	NRAS NSD1 NTRK1 NTRK3 NUMA1 NUP98 NUT OLIG2 OMD PAFAH1B2 PALB2 PALB2 PATZ1 PAX3 PAX5 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7	yes yes yes yes yes yes yes yes yes yes	yes yes	L, E E E, M L L E L M L L, M M L M E L E L L M L O L E, L E L M L O L E, L E L M E E L E L M E E L E L M E E L E, L L L	Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	Mis T T T T T T T F, N, Mis T T T T T T Mis, O, T T T T Mis, O, T T T Mis, F T Mis, N T T T T Mis, N, F, S T T T T T T T T
ENSG0000165671 ENSG0000198400 ENSG0000137497 ENSG0000110713 ENSG0000110713 ENSG00001205927 ENSG0000127083 ENSG0000127083 ENSG0000168092 ENSG000010105 ENSG0000135903 ENSG0000135903 ENSG00000125618 ENSG00000125618 ENSG00000125618 ENSG0000178104 ENSG0000178104 ENSG0000178104 ENSG0000178104 ENSG00001311 ENSG00001311 ENSG0000134853 ENSG0000134853 ENSG00000134853 ENSG00000134853 ENSG00000134853 ENSG00000134853 ENSG00000134853 ENSG00000134853 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000181690 ENSG00000140464 ENSG00000140464 ENSG0000014024511 ENSG00000142611 ENSG0000184985 ENSG0000164985 ENSG00000164985 ENSG00000164985 ENSG00000164985	ENSMUSG0000047588 ENSMUSG0000028072 ENSMUSG00000028072 ENSMUSG0000006306 ENSMUSG00000063550 ENSMUSG00000041358 ENSMUSG00000041358 ENSMUSG00000041358 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG0000002453 ENSMUSG0000002453 ENSMUSG0000002676 ENSMUSG000000252534 ENSMUSG000000252534 ENSMUSG00000025234 ENSMUSG00000025231 ENSMUSG00000025231 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000026098 ENSMUSG00000026098 ENSMUSG00000026098 ENSMUSG00000026098 ENSMUSG00000024014 ENSMUSG00000026098 ENSMUSG00000024014 ENSMUSG00000026098 ENSMUSG00000026098 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000026098 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG0000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014	NRAS NSD1 NTRK1 NTRK3 NUMA1 NUP98 NUT OLIG2 OMD PAFAH1B2 PALB2 PATZ1 PAX3 PAX5 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7	yes yes yes yes yes yes yes yes yes yes	yes yes yes	L, E E E E E L L E L M L L M L L E L E L E	Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	Mis T T T T T T T T T T T T T T T Mis, O, T T T T Mis, O, T T T Mis, F T T Mis, N T T T T T T T T T T T T T T T T T T T
ENSG0000165671 ENSG00001498400 ENSG0000149538 ENSG0000110713 ENSG0000110713 ENSG00001205927 ENSG0000127083 ENSG0000127083 ENSG0000168092 ENSG0000101015 ENSG000010105 ENSG0000105092 ENSG0000105092 ENSG00000125618 ENSG00000125618 ENSG00000125618 ENSG0000018601 ENSG0000018164 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000137193 ENSG00000142611 ENSG00000142611 ENSG00000164985 ENSG00000164985	ENSMUSG0000047588 ENSMUSG0000028072 ENSMUSG0000028072 ENSMUSG00000059146 ENSMUSG0000006306 ENSMUSG00000039800 ENSMUSG00000013530 ENSMUSG00000048768 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG00000044702 ENSMUSG0000004872 ENSMUSG0000004872 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG00000028736 ENSMUSG0000002893 ENSMUSG00000024620 ENSMUSG00000024620 ENSMUSG00000024614 ENSMUSG00000024014 ENSMUSG0000002828 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024014 ENSMUSG00000024016 ENSMUSG00000024016 ENSMUSG00000024016 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024106 ENSMUSG00000024101	NRAS NSD1 NTRK1 NTRK3 NUMA1 NUP98 NUT OLIG2 OMD PAFAH1B2 PALB2 PALB2 PATZ1 PAX3 PAX5 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7 PAX7	yes yes yes yes yes yes yes yes yes yes	yes yes	L, E E E, M L L E L M L L, M M L M E L E L L M L O L E, L E L M L O L E, L E L M E E L E L M E E L E L M E E L E, L L L	Dom Dom Dom Dom Dom Dom Dom Dom Dom Dom	Mis T T T T T T T F, N, Mis T T T T T T T Mis, O, T T T T Mis, O, T T T Mis, F T Mis, N T T T T Mis, N, F, S T T T

ENSG00000179295	ENSMUSG0000043733	PTPN11	yes		L	Dom	Mis
ENSG00000029725	ENSMUSG00000020817	RABEP1	yes		L	Dom	T
ENSG00000182185	ENSMUSG00000059060	RAD51L1	yes		M	Dom	Ť
ENSG00000204764	ENSMUSG00000040594	RANBP17	yes		L	Dom	Ť
ENSG00000138698	ENSMUSG00000028149	RAP1GDS1	yes		Ĺ	Dom	Ť
ENSG00000131759	ENSMUSG00000037992	RARA	yes		Ĺ	Dom	Ť
ENSG00000139687	ENSMUSG00000022105	RB1	yes	yes	L, E, M, O	Rec	D, Mis, N, F, S
ENSG00000162775	ENSMUSG00000048109	RBM15	yes	yes	L, L, II, U	Dom	Т
ENSG00000160957	ENSMUSG00000033762	RECQL4	ycs	VAC	M	Rec	, N, F, S
ENSG00000162924	ENSMUSG00000020275	REL	Vec	yes	L	Dom	N, F, S А
ENSG00000165731	ENSMUSG00000030110	RET	yes	1/05	Е, О		
ENSG00000168421	ENSMUSG00000029204	RHOH	yes	yes	L, U L	Dom Dom	T, Mis, N, F T
			yes				
ENSG00000047936	ENSMUSG00000019893	ROS1 RPL22	yes		0	Dom	T T
ENSG00000116251 ENSG00000163902	ENSMUSG0000028936 ENSMUSG00000030062	RPN1	yes		L	Dom Dom	T
ENSG00000159216	ENSMUSG00000022952	RUNX1	yes		L	Dom	Ť
ENSG00000079102	ENSMUSG0000006586	RUNX1	yes yes		L	Dom	Ť
210300000079102	211311030000000000000000000000000000000	ROMATI	ycs		-	Dom	Gene
ENSG00000126524	ENSMUSG0000025337	SBDS		yes	L	Rec	Conversion
ENSG00000117118	ENSMUSG0000009863	SDHB		yes	Ō	Rec	Mis, N, F
ENSG00000143252	ENSMUSG00000058076	SDHC		yes	õ	Rec	Mis, N, F
ENSG00000204370	ENSMUSG0000000171	SDHD		yes	õ	Rec	Mis, N, F, S
ENSG00000184702	ENSMUSG00000072214	SEPT5	yes	yes	L	Dom	Т
ENSG00000125354	ENSMUSG00000050379	SEPT6	yes		Ĺ	Dom	Ť
ENSG00000184640	ENSMUSG00000059248	SEPT9	yes		Ĺ	Dom	Ť
ENSG00000116560	ENSMUSG0000028820	SFPQ	yes		Ē	Dom	Ť
ENSG00000141985	ENSMUSG0000003200	SH3GL1	yes		L	Dom	T
ENSG0000008300	ENSMUSG0000023473	SLC26A6	yes		Ĺ	Dom	Ť
ENSG00000141646	ENSMUSG0000024515	SMAD4	yes	yes	E	Rec	D, Mis, N, F
ENSG00000099956	ENSMUSG0000000902	SMARCB1	yes	yes	M	Rec	D, N, F, S
ENSG00000128602	ENSMUSG0000001761	SMO	yes	,	E	Dom	Mis
ENSG00000185338	ENSMUSG00000038037	SOCS1	yes		Ĺ	Rec	F, O
ENSG00000128487	ENSMUSG00000042331	SPECC1	yes		Ĺ	Dom	Т
ENSG00000141380	ENSMUSG0000037013	SS18	yes		M	Dom	T
ENSG00000184402	ENSMUSG00000039086	SS18L1	yes		M	Dom	Ť
ENSG00000126752	ENSMUSG00000035371	SSX1	yes		M	Dom	T
ENSG00000126752	ENSMUSG00000071816	SSX1	yes		M	Dom	Ť
ENSG00000126752	ENSMUSG00000023165	SSX1	yes		M	Dom	T
ENSG00000126752	ENSMUSG00000068218	SSX1	yes		M	Dom	Ť
ENSG00000126752	ENSMUSG0000062814	SSX1	yes		M	Dom	Ť
ENSG00000126752	ENSMUSG00000068219	SSX1	yes		M	Dom	T
ENSG00000187754	ENSMUSG00000035371	SSX2	yes		M	Dom	Ť
ENSG00000187754	ENSMUSG0000071816	SSX2	yes		M	Dom	Т
ENSG00000187754	ENSMUSG00000023165	SSX2	yes		M	Dom	Ť
ENSG00000187754	ENSMUSG0000068218	SSX2	yes		M	Dom	Т
ENSG00000187754	ENSMUSG0000062814	SSX2	yes		M	Dom	Ť
ENSG00000187754	ENSMUSG0000068219	SSX2	yes		M	Dom	Ť
ENSG00000204645	ENSMUSG0000035371	SSX4	yes		М	Dom	т
ENSG00000204645	ENSMUSG0000071816	SSX4	yes		M	Dom	Ť
ENSG00000204645	ENSMUSG0000023165	SSX4	yes		М	Dom	т
ENSG00000204645	ENSMUSG0000068218	SSX4	yes		M	Dom	Ť
ENSG00000204645	ENSMUSG0000062814	SSX4	yes		М	Dom	т
ENSG00000204645	ENSMUSG0000068219	SSX4	yes		M	Dom	Ť
ENSG00000123473	ENSMUSG0000028718	STIL	yes		L	Dom	т
ENSG00000118046	ENSMUSG0000003068	STK11	yes	yes	Е, М, О	Rec	D, Mis, N, F, S
ENSG00000107882	ENSMUSG0000025231	SUFU	yes	yes	0	Rec	D, F, S
ENSG00000178691	ENSMUSG0000017548	SUZ12	yes	,	М	Dom	Т
ENSG00000165025	ENSMUSG0000021457	SYK	yes		L	Dom	Т
ENSG00000172660	ENSMUSG0000020680	TAF15	yes		L, M	Dom	Т
ENSG00000162367	ENSMUSG0000028717	TAL1	yes		Ľ	Dom	Т
ENSG00000186051	ENSMUSG0000028417	TAL2	yes		L	Dom	Т
ENSG00000187735	ENSMUSG0000033813	TCEA1	yes		E	Dom	Т
ENSG00000135100	ENSMUSG0000029556	TCF1	yes	yes	E	Rec	Mis, F
ENSG00000140262	ENSMUSG0000032228	TCF12	yes		М	Dom	Т
ENSG0000071564	ENSMUSG0000020167	TCF3	yes		L	Dom	Т
ENSG00000100721	ENSMUSG0000041359	TCL1A	yes		L	Dom	Т
ENSG00000135605	ENSMUSG0000029217	TEC	yes		М	Dom	Т
ENSG0000068323	ENSMUSG0000000134	TFE3	yes		E	Dom	Т
ENSG00000112561	ENSMUSG0000023990	TFEB	yes		E,M	Dom	T
ENSG00000114354	ENSMUSG0000022757	TFG	yes		E, L	Dom	Т
ENSG00000105619	ENSMUSG0000006335	TFPT	yes		L	Dom	Т
ENSG0000072274	ENSMUSG0000022797	TFRC	yes		L	Dom	T
ENSG00000054118	ENSMUSG0000072862	THRAP3	yes		М	Dom	T
ENSG0000054118	ENSMUSG0000043962	THRAP3	yes		м	Dom	Т
ENSG00000107807	ENSMUSG0000025215	TLX1	yes		L	Dom	Т
ENSG00000164438	ENSMUSG0000040610	TLX3	yes		L	Dom	T
ENSG00000184012	ENSMUSG0000000385	TMPRSS2	yes		E	Dom	Т
ENSG00000048462	ENSMUSG0000022496	TNFRSF17	yes		L	Dom	T
ENSG00000198900	ENSMUSG00000070544	TOP1	yes		L	Dom	T Mic N E
ENSG00000141510	ENSMUSG00000059552	TP53	yes	yes	L, E, M, O	Rec	Mis, N, F
ENSG00000167460	ENSMUSG00000031799	TPM4	yes		L	Dom	T
ENSG00000047410	ENSMUSG00000006005	TPR	yes		E	Dom	T T
ENSG00000122779	ENSMUSG00000029833	TRIM24	yes		L E	Dom	T
ENSG00000197323	ENSMUSG00000033014	TRIM33	yes			Dom	T
ENSG00000100815	ENSMUSG00000021188	TRIP11	yes	Voc	L	Dom	
ENSG00000165699	ENSMUSG00000026812	TSC1		yes	E, O E, O	Rec	D, Mis, N, F, S
ENSG00000103197	ENSMUSG0000002496	TSC2 TSHR	Vec	yes	E, U E	Rec Dom	D, Mis, N, F, S Mis
ENSG00000165409 ENSG00000114999	ENSMUSG00000020963 ENSMUSG00000027394	TSHR TTL	yes	yes	L	Dom	T
ENSG000001149999	ENSMUSG00000027394 ENSMUSG0000000804	USP6	yes		L M	Dom	Т
ENSG00000129204 ENSG00000134086	ENSMUSG0000000804 ENSMUSG00000033933	VHL	yes	VAC	™ E, M, O	Rec	
LN300000134000	EN31:10300000033333	VIIL	yes	yes	L, M, U	X-linked	D, Mis, N, F, S
ENSG00000015285	ENSMUSG0000031165	WAS			L	Rec	Mis, N, F, S
ENSG00000109685	ENSMUSG00000057406	WAS WHSC1	yes		L	Dom	т, т
ENSG00000147548	ENSMUSG00000054823	WHSC1L1	yes yes		L	Dom	Т
2.100000017/070	2.1511050000000000002025		,		-	Doni	•

ENSG00000165392	ENSMUSG0000031583	WRN		yes	L, E, M, O	Rec	Mis, N, F, S
ENSG00000184937	ENSMUSG0000016458	WT1	yes	yes	0	Rec	D, Mis, N, F, S
ENSG00000136936	ENSMUSG0000028329	XPA		yes	E	Rec	Mis, N, F, S
ENSG00000154767	ENSMUSG0000030094	XPC		yes	E	Rec	Mis, N, F, S
ENSG00000109906	ENSMUSG0000066687	ZBTB16	yes		L	Dom	т
ENSG00000121741	ENSMUSG0000021945	ZMYM2	yes		L	Dom	т
ENSG00000126746	ENSMUSG0000038346	ZNF384	yes		L	Dom	Т

Appendix B1. *Sleeping Beauty* CISs and predicted CIS genes obtained using the kernel convolution-based framework with a kernel width of 30 kb.

"None" is used where there is no obvious candidate gene associated with the CIS.

"CIS chr" and "CIS position" are the chromosome and base pair coordinates of the CIS, which correspond to the centre of the peak in insertion density identified by the kernel convolution-based method.

"Biotype" indicates whether the gene encodes a protein or miRNA.

"Real" indicates whether the gene is believed to be a real cancer gene candidate.

CIS chr	CIS position	Gene name	Ensembl ID	Biotype	Real?
1	161802810	Tnr	ENSMUSG0000015829	protein_coding	N
1	162750720	None	N/A	N/A	N
1	162810060	Rc3h1	ENSMUSG0000040423	protein_coding	N
1	162918420	Cenpl	ENSMUSG0000026708	protein_coding	N
1	164494053	Q3UWS6_mouse	ENSMUSG0000073516	protein_coding	N
2	26289038	Notch1	ENSMUSG0000026923	protein_coding	Y
2	98464591		ENSMUSG0000075015	protein_coding	Y
3	136852080	Ррр3са	ENSMUSG0000028161	protein_coding	Y
5	28500825	En2	ENSMUSG0000039095	protein_coding	N
7	27307506	Akt2	ENSMUSG0000004056	protein_coding	Y
9	32222013	Fli1	ENSMUSG0000016087	protein_coding	Y
9	45939227	BC033915	ENSMUSG0000034135	protein_coding	Y
10	20825984	Myb	ENSMUSG0000019982	protein_coding	Y
11	11653286	Ikzf1	ENSMUSG0000018654	protein_coding	Y
13	31634753	Foxf2	ENSMUSG0000038402	protein_coding	N
15	3434345	Ghr	ENSMUSG0000055737	protein_coding	N
15	3536237	Ghr	ENSMUSG0000055737	protein_coding	N
15	28104183	ENSMUSESTG00003714390	N/A	EST	N
15	39369050	Rims2	ENSMUSG0000037386	protein_coding	N
16	95500025	Erg	ENSMUSG0000040732	protein_coding	Y
19	32868157	Pten	ENSMUSG0000013663	protein_coding	Y

Appendix B2. MuLV CISs and predicted CIS genes obtained using the kernel convolution-based framework with a kernel width of 30 kb.

"CIS chr" and "CIS" position are the chromosome and base pair coordinates of the CIS, which correspond to the centre of the peak in insertion density identified by the kernel convolution-based method.

"Biotype" indicates whether the gene encodes a protein or miRNA.

CIS chr	CIS position	Gene name	Ensembl ID	Biotype
1	36867188	Tmem131	ENSMUSG0000026116	protein_coding
1	37429920	Mgat4a	ENSMUSG0000026110	protein_coding
1	88266570	1700019017Rik	ENSMUSG0000036574	protein_coding
1	92886988	Lrrfip1	ENSMUSG0000026305	protein_coding
1	93017760	Ramp1	ENSMUSG0000034353	protein_coding
1	127266450	Actr3	ENSMUSG0000026341	protein_coding
1	130440120	Cxcr4	ENSMUSG0000045382	protein_coding
1	135882360	Btg2	ENSMUSG0000020423	protein_coding
1	138068640	5730559C18Rik	ENSMUSG0000041605	protein_coding
1	139624096	A130050007Rik	ENSMUSG0000051480	protein_coding
1	139988370	Ptprc	ENSMUSG0000026395	protein_coding
1	158862420	ENSMUSG0000073531	ENSMUSG0000073531	protein_coding
1	163733040	AI848100	ENSMUSG0000040297	protein_coding
1	165910570	Sell	ENSMUSG0000026581	protein_coding
1	167543910	Rcsd1	ENSMUSG0000040723	protein_coding
1	173513310	Cd48	ENSMUSG0000015355	protein_coding
1	173513310	Slamf7	ENSMUSG0000038179	protein_coding
1	173759850	Slamf6	ENSMUSG0000015314	protein_coding
1	174351491	Ccdc19	ENSMUSG0000026546	protein_coding
1	182195685	Itpkb	ENSMUSG0000038855	protein_coding
1	182545770	Mix11	ENSMUSG0000026497	protein_coding
1	183688021	EG433384	ENSMUSG0000056699	protein_coding
1	193218780	Ppp2r5a	ENSMUSG0000026626	protein_coding
2	6625263	Cugbp2	ENSMUSG0000002107	protein_coding
2	11544848	Il2ra	ENSMUSG0000026770	protein_coding
2	18526199		ENSMUSG0000046809	protein_coding
2	26286578	Notch1	ENSMUSG0000026923	protein_coding
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16	49720413	Cd47
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16	92755807	Runx1 Runx1
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ENSMUSG0000069082 ENSMUSG0000022372 ENSMUSG0000034596 ENSMUSG0000014508 ENSMUSG0000014508 ENSMUSG0000022471 ENSMUSG0000022471 ENSMUSG00000022473 ENSMUSG00000022475 ENSMUSG00000022475 ENSMUSG0000002710 ENSMUSG0000002710 ENSMUSG0000002710 ENSMUSG0000002710 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000075269 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000072286	protein_coding protein_coding
ENSMUSG0000069082 ENSMUSG0000022372 ENSMUSG00000059895 ENSMUSG0000014596 ENSMUSG0000014508 ENSMUSG000002471 ENSMUSG0000002471 ENSMUSG0000002475 ENSMUSG0000002475 ENSMUSG0000002475 ENSMUSG0000002710 ENSMUSG0000002710 ENSMUSG0000002710 ENSMUSG0000005280 ENSMUSG0000005280 ENSMUSG0000005280 ENSMUSG0000005280 ENSMUSG0000005280 ENSMUSG0000005540 ENSMUSG0000005540 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG0000076457 ENSMUSG0000076457 ENSMUSG0000076457 ENSMUSG0000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000072952 ENSMUSG00000076457 ENSMUSG00000022946 ENSMUSG0000002297 ENSMUSG0000002297 ENSMUSG0000003297	protein_coding protein_coding
ENSMUSG0000069082 ENSMUSG0000022372 ENSMUSG00000059895 ENSMUSG0000014596 ENSMUSG00000142351 ENSMUSG000002471 ENSMUSG0000002471 ENSMUSG0000002475 ENSMUSG0000002475 ENSMUSG0000002710 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000055407 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG0000002952 ENSMUSG0000022952 ENSMUSG000000029	protein_coding protein_coding
ENSMUSG0000069082 ENSMUSG0000022372 ENSMUSG0000034596 ENSMUSG0000014508 ENSMUSG0000014508 ENSMUSG000002471 ENSMUSG0000022471 ENSMUSG0000002473 ENSMUSG00000022475 ENSMUSG00000058794 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG0000005540 ENSMUSG0000005540 ENSMUSG00000022952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000076457 ENSMUSG00000022952 ENSMUSG00000076457 ENSMUSG00000022952 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000076457 ENSMUSG00000022952 ENSMUSG000000022952 ENSMUSG000000022952 ENSMUSG00000000000000000000000000000000000	protein_coding protein_coding
ENSMUSG0000069082 ENSMUSG0000022372 ENSMUSG00000059895 ENSMUSG0000014596 ENSMUSG00000142351 ENSMUSG000002471 ENSMUSG0000002471 ENSMUSG0000002475 ENSMUSG0000002475 ENSMUSG0000002710 ENSMUSG0000005718 ENSMUSG0000005718 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000059280 ENSMUSG00000055407 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG0000002952 ENSMUSG0000022952 ENSMUSG000000029	protein_coding protein_coding

17	28241401	E230001N04Rik	ENSMUSG0000066170	protein_coding
17	29125068	Fgd2	ENSMUSG0000024013	protein_coding
17	29221740	Pim1	ENSMUSG0000024014	protein_coding
17	30355057	Dnahc8	ENSMUSG0000033826	protein_coding
17	30796931	Abcg1	ENSMUSG0000024030	protein_coding
17	30927241	Ubash3a	ENSMUSG0000042345	protein_coding
17	30927241	Tmprss3	ENSMUSG0000024034	protein_coding
17	33731492	Brd2	ENSMUSG0000024335	protein_coding
17	33809236	Tap2	ENSMUSG0000024339	protein coding
17	33809236	Psmb8	ENSMUSG0000024338	protein_coding
17	34656136	Msh5	ENSMUSG0000007035	protein coding
17	34814198	Lta	ENSMUSG0000024402	protein_coding
17	34814198	Nfkbil1	ENSMUSG0000042419	protein coding
17	34960681	H2-D1	ENSMUSG0000073411	protein_coding
17	35524109	Ddr1	ENSMUSG0000003534	protein_coding
17	35630144	Ppp1r10	ENSMUSG0000039220	protein coding
17	44245111	Supt3h	ENSMUSG0000038954	protein_coding
17	44956111	Aars2	ENSMUSG0000023938	protein_coding
17	46991279	Ccnd3	ENSMUSG0000034165	protein_coding
17	47060071	Ccnd3	ENSMUSG0000034165	protein_coding
17	47504543	1700122011Rik	ENSMUSG0000042494	protein_coding
17	55928551	Arrdc5	ENSMUSG0000073380	protein_coding
17	71390787	Q8BP09 MOUSE	ENSMUSG0000041913	protein_coding
17	83869302	Наао	ENSMUSG000000673	protein_coding
17	84056736	Zfp36l2	ENSMUSG00000045817	protein_coding
18	4333434	Map3k8	ENSMUSG0000024235	protein_coding
18	5352149	Zfp438	ENSMUSG0000050945	protein_coding
18	15349211	Aqp4	ENSMUSG0000024411	protein coding
18	35081394	Hspa9	ENSMUSG0000024359	protein coding
18	35868041	Tmem173	ENSMUSG0000024349	protein_coding
18	39149231	Arhgap26	ENSMUSG0000036452	protein coding
18	55115767	Zfp608	ENSMUSG0000052713	protein coding
18	60880541	Rps14	ENSMUSG0000024608	protein_coding
18	61000211	Tcof1	ENSMUSG0000024613	protein coding
18	65035691	Nedd4l	ENSMUSG0000024589	protein_coding
18	68349821	D18Ertd653e	ENSMUSG0000024544	protein coding
18	70687614	Mbd2	ENSMUSG0000024513	protein coding
18	75397751	Dym	ENSMUSG00000035765	protein coding
18	85017491	Cyb5	ENSMUSG0000024646	protein coding
19	4144414	Gpr152	ENSMUSG0000044724	protein_coding
19	4299940	Adrbk1	ENSMUSG0000024858	protein coding
19	5810614	Frmd8	ENSMUSG0000043488	protein_coding
19	5810614	Scyl1	ENSMUSG0000024941	protein coding
19	5844306	Frmd8	ENSMUSG0000024941 ENSMUSG0000024816	protein_coding
19	6399309	Rasgrp2	ENSMUSG0000024010	protein coding
19	6399309	Pygm	ENSMUSG0000032648	protein_coding
19	9110974	Ahnak	ENSMUSG0000069833	protein_coding
19	10649744	Cybasc3	ENSMUSG00000034445	
19	25054444	Dock8	ENSMUSG0000052085	protein_coding protein_coding
19	36321364	Pcqf5	ENSMUSG0000052085 ENSMUSG0000024805	protein_coding
19	37556224	Pcgr5 Hhex	ENSMUSG0000024805 ENSMUSG0000024986	
19	37556224	Exoc6	ENSMUSG0000024988 ENSMUSG00000053799	protein_coding
19	41890234	Frat2	ENSMUSG0000003799 ENSMUSG00000047604	protein_coding
19	44437354	Scd1	ENSMUSG0000047804 ENSMUSG00000037071	protein_coding protein coding
19		Ina	ENSMUSG0000037071 ENSMUSG00000034336	
19	47024957 53487874	INA ENSMUSG00000074788		protein_coding
19	5540/0/4	EN31103G000000/4/88	ENSMUSG0000074788	protein_coding

Appendix C1. List showing other cancer-associated datasets in which the MuLV CIS genes appear.

RTCGD (retro) = genes identified by retroviral insertional mutagenesis in the RTCGD database (RETRO = CIS gene in RTCGD dataset)

RTCGD (SB) = genes identified by transposon-mediated insertional mutagenesis in the RTCGD database (SB = CIS gene in RTCGD dataset)

CGC = Cancer Gene Census (Dom = CIS gene is a dominant cancer gene)

COSMIC = genes mutated in COSMIC database (COSMIC = CIS gene mutated in COSMIC)

Sjoblom = candidate cancer genes from Sjöblom et al. (2006) (Sjoblom = CIS gene in Sjoblom dataset)

Nanog BS = genes with Nanog binding site (Nanog = CIS gene in Nanog dataset)

Oct4 BS = genes with Oct4 binding site (OCT4 = CIS gene in Oct4 dataset)

p53 BS = genes with p53 binding site (p53 (dn) = CIS gene downregulated by p53; p53 (up) = CIS

gene upregulated by p53; p53 = effect of p53 binding unknown)

Mullighan Amps/Dels = genes in recurrent amplicons and deletions in acute lymphoblastic leukaemia dataset of Mullighan *et al.* (2007) (AMP = CIS gene in amplicon; DEL = CIS gene in deletion) n/a is used where the gene has no known human orthologue and cannot be compared to human cancerassociated datasets.

Gene name	Ensembl ID	RTCGD (retro)	RTCGD (SB)	CGC	COSMIC	Sjoblom	Nanog BS	Oct4 BS	p53 BS		ighan 5 / Dels
1110036003Rik	ENSMUSG0000006931	-	-	-	-	-	-	-	-	-	-
1190002H23Rik	ENSMUSG0000022018	-	-	-	COSMIC	-	-	-	-	-	-
1300007F04Rik	ENSMUSG0000000686	-	-	-	-	-	-	-	-	-	-
1600014C10Rik	ENSMUSG0000054676	-	-	-	-	-	-	-	-	-	-
1700019017Rik	ENSMUSG0000036574	-	-	-	-	-	Nanog	-	-	-	-
1700081D17Rik	ENSMUSG0000022085	-	-	-	-	-	-	-	-	-	-
1700122011Rik	ENSMUSG0000042494	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
2010106G01Rik	ENSMUSG0000027366	-	-	-	-	-	-	-	-	-	-
2010107G12Rik	ENSMUSG0000029847	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
2310016C08Rik	ENSMUSG0000043421	-	-	-	-	-	-	-	-	-	-
2310066E14Rik	ENSMUSG0000038604	-	-	-	-	-	-	-	-	-	DEL
2410014A08Rik	ENSMUSG0000048578	-	-	-	-	-	Nanog	-	-	-	-
3110082I17Rik	ENSMUSG0000053553	-	-	-	-	-	-	-	-	-	DEL
3930402G23Rik	ENSMUSG0000038917	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
4632428N05Rik	ENSMUSG0000020101	-	-	-	-	-	-	-	-	-	-
4831426I19Rik	ENSMUSG0000054150	RETRO	-	-	-	-	Nanog	-	-	-	-
4932417H02Rik	ENSMUSG0000025583	-	-	-	COSMIC	-	-	OCT4	-	-	-
4932422M17Rik	ENSMUSG0000062946	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
5730559C18Rik	ENSMUSG0000041605	-	-	-	-	-	-	-	-	AMP	-
6330548G22Rik	ENSMUSG0000029402	-	-	-	-	-	-	-	-	-	-
6430598A04Rik	ENSMUSG0000045348	-	-	-	-	-	-	-	-	-	-
A130050007Rik	ENSMUSG0000051480	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
A130092J06Rik	ENSMUSG0000050592	-	-	-	-	-	-	-	-	AMP	-
A2AN91_MOUSE	ENSMUSG0000038578	-	-	-	-	-	-	-	p53	AMP	-
A530013C23Rik	ENSMUSG0000006462	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
A630001012Rik	ENSMUSG0000074025	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
A930002I21Rik	ENSMUSG0000050179	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
AA536749	ENSMUSG0000005417	-	-	-	-	-	-	-	-	-	DEL
Aars2	ENSMUSG0000023938	-	-	-	-	-	-	-	-	-	-
AB041803	ENSMUSG0000044471	RETRO	-	n/a	n/a	n/a	Nanog	-	n/a	n/a	n/a
Abcb9	ENSMUSG0000029408	-	-	-	-	-	-	-	p53	-	-
Abcg1	ENSMUSG0000024030	-	-	-	-	-	-	-	-	-	-
Acot11	ENSMUSG0000034853	-	-	-	-	-	-	-	-	-	-
Actr3	ENSMUSG00000026341	-	-	-	-	-	-	-	-	-	-
Adrbk1	ENSMUSG0000024858	_	-	-	COSMIC	-	-	-	-	-	-
Ahi1	ENSMUSG0000019986	RETRO	-	-	COSMIC	-	Nanog	-	-	AMP	-
Ahnak	ENSMUSG0000069833	-	-	-	-	-	-	-	-	-	-
AI848100	ENSMUSG0000040297	_	-	-	_	-	-	-	-	AMP	-
Ak1	ENSMUSG00000026817	_	-	-	_	-	-	-	-	AMP	-
Akap13	ENSMUSG0000066406	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Akna	ENSMUSG00000039158	-	-	-	-	-	-	-	-	AMP	-
Akt1	ENSMUSG0000001729	RETRO	_	_	_	_	_	-		-	_
Anp32e	ENSMUSG00000015749	-	-	-	_	-	-	-	-	AMP	-
Anxa2	ENSMUSG0000032231	-	_	_	_	_	_	-		-	DEL
Appl2	ENSMUSG0000020263	_	-	-	-	-	-	-	-	-	-
Aqp4	ENSMUSG0000024411	_	_	_	_	_	_	-		-	-
Aqp9	ENSMUSG0000032204	-	_	_	_	_	_	-		-	_
Arhgap26	ENSMUSG0000036452	-	_	Dom	_	_	_	OCT4		-	_
Arhgdib	ENSMUSG0000030220	_	_	-	_	_	_	-	_	_	_
Arhgef10l	ENSMUSG00000040964	_	_	_	_	_		OCT4	_	_	_
Arhgef3	ENSMUSG00000021895	-	-	-	-	-	Napog	0014	-	-	-
Arid1a	ENSMUSG00000021895	- RETRO	-	-	- COSMIC	-	Nanog -	-	-	-	-
	ENSMUSG00000019564	KLIKU	-	-	-	-	-	-	-	-	- DEL
Arid3a Aron21		-	-	-	-	-	-	-	-	-	
Arpp21	ENSMUSG00000032503	-	-	-	-	-	-	-	-	-	-
Arrdc5	ENSMUSG0000073380	-	-	-	-	-	-	-	-	-	-
Art2b	ENSMUSG0000030651	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Asb2	ENSMUSG00000021200	-	-	-		-	-	-	-	-	-
B230120H23Rik	ENSMUSG0000004085	-	-	-	COSMIC	-	-	-	-	-	-
B3gnt2	ENSMUSG0000051650	-	-	-	-	-	-	-	-	-	-
B3gntl1	ENSMUSG00000046605	-	-	-	-	-	-	-	-	-	-
BC008155	ENSMUSG0000057411	-	-	-	-	-	-	-	-	-	-

00007057											
BC027057	ENSMUSG00000049625	-	-	-	-	-	-	-	-	-	-
BC030863 Bcl11a	ENSMUSG00000051586	- RETRO	-	- Dom	-	- Cioblom	-	-	-	AMP	-
Bcl11b	ENSMUSG0000000861 ENSMUSG00000048251	RETRO	-	Dom Dom	COSMIC COSMIC	Sjoblom -	-		-	-	-
Bcl2l1	ENSMUSG0000007659	RETRO	_	-	-	_	Nanog	_	-	-	DEL
Bcl9l	ENSMUSG0000063382	-	-	-	-	-	-	-	-	-	DEL
Bex6	ENSMUSG00000075269	-	-	-	COSMIC	-	-	-	-	-	-
Bid	ENSMUSG0000004446	RETRO	-	-	-	-	-	-	-	AMP	-
Brd2	ENSMUSG0000024335	RETRO	-	-	COSMIC	-	-	OCT4	-	-	-
Btg2	ENSMUSG0000020423	RETRO	-	-	-	-	-	-	-	AMP	-
C130026L21Rik	ENSMUSG0000052848	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
C330016O10Rik	ENSMUSG0000001053	RETRO	-	-	-	-	-	-	-	-	DEL
C330024D12Rik	ENSMUSG0000030553	-	-	-	-	-	-	-	-	-	-
Capsl	ENSMUSG0000039676	-	-	-	-	-	-	-	-	-	-
Cbfa2t3	ENSMUSG0000006362	RETRO	-	Dom	-	-	-	-	-	-	-
Cbl	ENSMUSG0000034342	-	-	Dom	-	-	-	-	-	-	-
Ccdc19	ENSMUSG0000026546	-	-	-	-	-	-	-	-	AMP	-
Ccnd1	ENSMUSG0000070348	RETRO	-	Dom	-	-	-	-	-	-	-
Ccnd2	ENSMUSG0000000184	RETRO	-	Dom	-	-	-	-	-	-	-
Ccnd3	ENSMUSG0000034165	RETRO	-	Dom	-	-	-	-	-	-	-
Ccr7	ENSMUSG0000037944	RETRO	-	-	-	-	-	-	-	-	-
Ccrk	ENSMUSG0000021483	RETRO	-	-	-	-	-	-	-	AMP	-
Cd2	ENSMUSG0000027863	-	-	-	-	-	-	-	-	-	-
Cd27	ENSMUSG0000030336	-	-	-	-	-	-	-	-	-	-
Cd3e Cd47	ENSMUSG0000032093 ENSMUSG00000055447	-	-		-	-	-	-			-
Cd48	ENSMUSG00000015355	_	_	_	_	_	_	_	-	AMP	_
Cd53	ENSMUSG00000013333	_	_	_	_	_	_	_	-	-	_
Cd72	ENSMUSG00000028459	-	_	_	-	-	-	-	-	-	DEL
Cd97	ENSMUSG0000002885	-	-	-	-	-	-	OCT4	-	-	-
Cdkl3	ENSMUSG00000020389	-	-	-	COSMIC	-	-	-	-	-	-
Cecr5	ENSMUSG00000058979	RETRO	-	-	-	-	Nanog	-	-	AMP	-
Chchd7	ENSMUSG0000042198	-	-	-	-	-	-	-	p53	-	-
Chd1	ENSMUSG0000023852	-	-	-	COSMIC	-	Nanog	OCT4	p53	-	-
Chd2	ENSMUSG0000025788	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Chd7	ENSMUSG0000041235	-	-	-	-	-	Nanog	-	-	-	-
Chst3	ENSMUSG0000057337	-	-	-	-	-	Nanog	OCT4	-	-	-
Cldn10a	ENSMUSG0000022132	-	-	-	-	-	-	-	-	-	-
Clec16a	ENSMUSG0000068663	-	-	-	-	-	-	-	-	-	-
Clec2d	ENSMUSG0000030157	-	-	-	-	-	-	-	-	-	-
Colq	ENSMUSG0000057606	-	-	-	-	-	-	-	-	-	-
Coro2a	ENSMUSG0000028337	RETRO	-	-	-	-	-	-	-	AMP	-
Csk	ENSMUSG00000032312	-	-	- ,	COSMIC	- ,	-	-	- ,	- ,	- ,
Cstad	ENSMUSG00000047363	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Cugbp2	ENSMUSG0000002107	-	-	-	-	-	-	OCT4	-	AMP	-
Cxcr4	ENSMUSG00000045382	-	-	-	COSMIC	-	- Nanog	-	-	-	-
Cyb5	ENSMUSG00000024646 ENSMUSG00000034445	-	-	-	-	-	Nanog	-	-	-	-
<i>Cybasc3 D12Ertd553e</i>	ENSMUSG00000020589	RETRO	_	_	_	_	Nanog	_	-	AMP	_
D18Ertd653e	ENSMUSG0000024544	-	_	_	-	-	-	-	-	-	-
Dad1	ENSMUSG0000022174	-	-	-	-	-	-	-	-	-	-
Ddr1	ENSMUSG0000003534	RETRO	-	-	COSMIC	-	-	-	-	-	-
Dhx40	ENSMUSG0000018425	-	-	-	-	-	-	-	-	-	-
Dnahc8	ENSMUSG0000033826	-	-	-	COSMIC	-	Nanog	OCT4	-	-	-
Dock8	ENSMUSG0000052085	-	-	-	-	-	-	-	-	-	DEL
Dopey2	ENSMUSG0000022946	RETRO	-	-	-	-	Nanog	-	-	-	-
Dpp4	ENSMUSG0000035000	-	-	-	-	-	-	-	-	-	-
Dym	ENSMUSG0000035765	RETRO	-	-	-	-	-	-	-	-	-
E230001N04Rik	ENSMUSG0000066170	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
E2f2	ENSMUSG0000018983	RETRO	-	-	-	-	-	-	-	-	-
Edg1	ENSMUSG0000045092	-	-	-	-	-	Nanog	-	-	-	-
EG433384	ENSMUSG0000056699	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Egfl7	ENSMUSG0000026921	-	-	-	-	-	-	-	-	AMP	-
Elf1	ENSMUSG0000036461	-	-	-	-	-	-	-	-	-	DEL
Emp3	ENSMUSG0000040212	-	-	-	-	-	-	-	-	-	-
Eng Epha6	ENSMUSG00000026814 ENSMUSG00000055540	-	-	- n/a	- n/a	- n/a	-	-	- n/2	AMP	- n/a
Erg	ENSMUSG00000055540 ENSMUSG00000040732	- RETRO	- SB	n/a Dom	n/a -	n/a -	-	-	n/a -	n/a -	n/a DEL
Ets1	ENSMUSG00000032035	RETRO	SB	-	COSMIC	_	_	_	-	-	-
Etv6	ENSMUSG0000032033	-	-	- Dom	-	-	-	-	-	-	- DEL
Evi1	ENSMUSG00000027684	RETRO	-	Dom	COSMIC	-	-	-	-	-	-
Evi2b	ENSMUSG0000070354	-	-	-	-	-	-	-	-	-	-
Evi5	ENSMUSG00000011831	-	-	-	-	-	-	-	-	-	-
Exoc2	ENSMUSG0000021357	-	-	-	COSMIC	-	-	-	-	AMP	-
Exoc6	ENSMUSG0000053799	-	-	-	-	-	-	-	-	-	-
Exosc5	ENSMUSG0000061286	-	-	-	-	-	-	-	-	-	-
Fchsd2	ENSMUSG0000030691	-	-	-	-	-	-	-	-	-	-
Fgd2	ENSMUSG0000024013	-	-	-	COSMIC	-	Nanog	-	-	-	-
Fgfr2	ENSMUSG0000030849	-	-	Dom	COSMIC	-	-	-	-	-	-
Fgfr3	ENSMUSG0000054252	RETRO	-	Dom	COSMIC	-	-	-	-	-	-
Fgr	ENSMUSG0000028874	RETRO	-	-	COSMIC	-	Nanog	-	-	-	-
Fli1	ENSMUSG0000016087	RETRO	SB	Dom	COSMIC	-	-	-	-	-	-
Flt3	ENSMUSG00000042817	RETRO	-	Dom	COSMIC	-	-	-	-	-	-
Fmnl1	ENSMUSG0000055805	-	-	-	-	-	Nanog	-	-	-	-
Foxp1	ENSMUSG0000030067	-	-	-	COSMIC	-	Nanog	-	-	-	-
Frat2	ENSMUSG0000047604		-	-		-	-	-	-	-	-
Frmd8 Frmd8	ENSMUSG0000024816	RETRO	-	-	COSMIC	-	-	-	-	- n/a	-
Frmd8 Fut8	ENSMUSG00000043488	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Fut8 Fyb	ENSMUSG00000021065	-	-	-	-	-	- Nanca	- 0CT4	-	-	-
Fyb Gadd45b	ENSMUSG00000022148 ENSMUSG00000015312	-	-	-	-	-	Nanog -	OCT4	-	-	-
Gadd45g	ENSMUSG00000015312 ENSMUSG00000021453	- RETRO		-	-	_	-	- OCT4	_	- AMP	-
Gadd45g Gfi1	ENSMUSG0000021453 ENSMUSG00000029275	RETRO	-	-	-	-	-	-	-	-	2
Gfi1b	ENSMUSG0000025275	RETRO	-	-	-	-	-	-	-	AMP	-
Ggta1	ENSMUSG00000035778	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
- 3				., .	,	, .			., .	., .	., u

Gimap4	ENSMUSG0000054435	-	-	-	-	-	-	-	-	-	-
Gimap6	ENSMUSG00000047867	-	-	-	-	-	-	-	-	-	-
Gm525	ENSMUSG0000072553	-	-	-	-	-	-	-	-	-	-
Gna15	ENSMUSG0000034792	-	-	-	-	-	-	-	-	-	-
Gpr132	ENSMUSG0000021298	-	-	-	-	-	-	-	-	-	-
Gpr152	ENSMUSG00000044724	-	-	-	-	-	-	-	-	-	-
Gpr56 Gran 2	ENSMUSG00000031785 ENSMUSG00000042351	-	-	-	-	-	- Nonog	-	-	-	-
Grap2			-	-	-	-	Nanog	OCT4	-	-	-
Gse1 H2-D1	ENSMUSG0000031822 ENSMUSG00000073411	RETRO	-	- n/a	COSMIC	- n/a	-	-	- n/a	- n/a	- n/a
Haao	ENSMUSG0000000673	_	_	n/a	n/a	n/a	_	_	p53	n/a	n/a
Hdac7a	ENSMUSG00000022475	-	-	-	COSMIC	_	-	-	p55	-	-
Hhex	ENSMUSG0000024986	RETRO	_	_	-		_	OCT4	_	_	_
Hibadh	ENSMUSG0000024980	-	_	_	_	_	Nanog	-	-	-	DEL
Hipk1	ENSMUSG0000008730	-	-	-	COSMIC	-	-	-	-	-	-
Hmga1	ENSMUSG00000046711	RETRO	-	Dom	-	-	-	-	-	-	-
Hnrpf	ENSMUSG0000042079	-	-	-	-	-	-	-	-	-	-
Hoxa7	ENSMUSG0000038236	RETRO	-	-	-	-	-	-	-	-	DEL
Hrbl	ENSMUSG0000029722	-	-	-	-	-	-	-	-	-	-
Hspa9	ENSMUSG0000024359	RETRO	-	-	-	-	-	-	-	-	-
Hvcn1	ENSMUSG0000064267	-	-	-	-	-	-	-	-	-	-
Ier2	ENSMUSG0000053560	RETRO	-	-	-	-	-	-	-	-	-
Ifnar1	ENSMUSG0000022967	RETRO	-	-	-	-	Nanog	-	p53	AMP	-
Ikzf1	ENSMUSG0000018654	RETRO	-	Dom	-	-	-	-	-	-	DEL
Ikzf3	ENSMUSG0000018168	RETRO	-	-	COSMIC	-	-	OCT4	-	-	DEL
II16	ENSMUSG0000001741	RETRO	-	-	COSMIC	-	-	-	-	-	-
Il21r	ENSMUSG0000030745	-	-	Dom	-	-	-	-	-	-	-
Il2ra	ENSMUSG0000026770	RETRO	-	-	-	-	-	-	-	AMP	-
Il6st	ENSMUSG0000021756	RETRO	-	-	-	-	Nanog	OCT4	-	-	-
Ina	ENSMUSG0000034336	-	-	-	-	-	Nanog	-	-	-	-
Iqch Irf2bn2	ENSMUSG00000037801		-	-	-	-	-	-	- n=2	- A M D	-
Irf2bp2 Irf4	ENSMUSG00000051495 ENSMUSG00000021356	RETRO -	-	- Dom	-	-	-	-	p53	AMP AMP	-
Itri4 Itpkb	ENSMUSG0000021356 ENSMUSG00000038855	-	-	Dom -	-	-	-	-	_	AMP AMP	-
ItprD Itpr2	ENSMUSG0000038855 ENSMUSG00000030287	-	-	-	- COSMIC	-	-	-	_	-	-
Jazf1	ENSMUSG00000063568	RETRO	_	Dom	-	_	_	_	_	-	DEL
Jph4	ENSMUSG00000022208	-	-	-	-	-	-	-	-	-	-
Jundm2	ENSMUSG0000034271	RETRO	-	-	-	-	-	-	-	-	-
Jup	ENSMUSG0000001552	-	-	-	-	-	-	-	-	-	-
Katnal1	ENSMUSG0000041298	-	-	-	-	-	-	-	-	-	-
Kcnn4	ENSMUSG0000054342	-	-	-	-	-	-	-	-	-	-
Kctd2	ENSMUSG0000016940	RETRO	-	-	-	-	-	OCT4	-	-	-
Kdr	ENSMUSG0000062960	RETRO	-	-	COSMIC	-	-	-	-	-	-
Kit	ENSMUSG0000005672	-	-	Dom	COSMIC	-	-	-	-	-	-
Klf3 Klhije	ENSMUSG0000029178	RETRO	-	-	COSMIC	-	-	-	-	-	-
Klhl25 Ksr1	ENSMUSG00000055652 ENSMUSG00000018334	-	-	-	- COSMIC	-	-	-	-	-	-
Lat	ENSMUSG00000030742	_	_	_	-	_	_	_	_	-	-
Lck	ENSMUSG0000000409	-	-	Dom	COSMIC	-	-	-	-	-	-
Lcp1	ENSMUSG00000021998	RETRO	-	Dom	-	-	-	-	-	-	-
Ldha	ENSMUSG0000063229	-	-	-	-	-	-	-	-	-	-
Lef1	ENSMUSG0000027985	RETRO	-	-	-	-	Nanog	-	-	-	DEL
Lfng	ENSMUSG0000029570	RETRO	-	-	-	-	-	-	-	-	DEL
Lgals9	ENSMUSG0000001123	-		-	-	-	-	-	-	-	DEL
			-		n/2		-			n/2	n/a
Lims1	ENSMUSG0000019920	-	-	n/a	n/a	n/a		-	n/a	n/a	
Lmo2	ENSMUSG00000019920 ENSMUSG00000032698	- RETRO	-	n/a n/a	n/a	n/a n/a	-	-	n/a n/a	n/a	n/a
Lmo2 Lrrc8c	ENSMUSG00000019920 ENSMUSG00000032698 ENSMUSG00000054720	- RETRO -	-			n/a -	-	-			-
Lmo2 Lrrc8c Lrrfip1	ENSMUSG00000019920 ENSMUSG00000032698 ENSMUSG00000054720 ENSMUSG00000026305	- RETRO - -	-				- - Nanog	- - -			n/a - DEL
Lmo2 Lrrc8c Lrrfip1 Lta	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000054720 ENSMUSG0000026305 ENSMUSG0000024402	- RETRO - - -	-			n/a -	-	-	n/a - - -	n/a - - -	-
Lmo2 Lrrc8c Lrrfip1 Lta Lyst	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000054720 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG00000019726	- - -	-			n/a -	- Nanog - Nanog	- - - - -			- DEL - -
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad1l1	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000054720 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000029554	- - - RETRO	-			n/a -	-	- - - - - OCT4	n/a - - -	n/a - - - AMP -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 Mafk	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000054720 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000029554 ENSMUSG0000018143	- - -	-		n/a - - - - -	n/a -	- Nanog - -	- - - - OCT4 -	n/a - - -	n/a - - - AMP - -	- DEL - -
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 Mafk Map3k1	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000026305 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000018143 ENSMUSG00000018143	- - - RETRO RETRO -	-			n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - - - -	n/a - - -	n/a - - - AMP - - -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad1l1 Mafk Map3k1 Map3k8	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000026305 ENSMUSG0000026305 ENSMUSG00000024402 ENSMUSG00000019726 ENSMUSG00000018143 ENSMUSG00000021754 ENSMUSG0000021754	- - - RETRO	-		n/a - - - - -	n/a -	- Nanog - -	- - - - - - 0CT4 - - - -	n/a - - -	n/a - - - AMP - -	- DEL - DEL
Lmo2 Lrrc8c Lrrf1p1 Lta Lyst Mad1l1 Mafk Map3k1 Map3k8 Mbd2	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG00000054720 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000029554 ENSMUSG00000018143 ENSMUSG000000242513	- - - RETRO RETRO - RETRO -	-		n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - OCT4 - - - - -	n/a - - p53 - - - -	n/a - - - AMP - - - AMP -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad1l1 Mafk Map3k1 Map3k8	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000026305 ENSMUSG0000026305 ENSMUSG00000024402 ENSMUSG00000019726 ENSMUSG00000018143 ENSMUSG00000021754 ENSMUSG0000021754	- - - RETRO RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - OCT4 - - - - - - -	n/a - - p53 - - - -	n/a - - - AMP - - -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad1/1 Mafk Map3k1 Map3k8 Mbd2 Mcl1	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000024305 ENSMUSG0000024402 ENSMUSG0000029554 ENSMUSG0000029554 ENSMUSG0000021754 ENSMUSG0000024513 ENSMUSG0000024513 ENSMUSG0000024513	- - - RETRO RETRO - RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - -	n/a - - p53 - - - - - -	n/a - - - AMP - - - AMP -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 Mafk Map3k1 Map3k8 Mbd2 Mc11 Med13	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000026305 ENSMUSG0000026305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG00000018143 ENSMUSG00000024754 ENSMUSG0000024235 ENSMUSG0000024513 ENSMUSG0000038612 ENSMUSG0000038612	- RETRO RETRO - RETRO - RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - -	n/a - - p53 - - - - - -	n/a - - AMP - - AMP - AMP -	- DEL - DEL
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Maf111 Mafk Map3k1 Map3k8 Mbd2 Mc11 Mc11 Mc13 Mef2d Metrn1 Mgat1	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000026305 ENSMUSG000002402 ENSMUSG0000029534 ENSMUSG0000029554 ENSMUSG0000024513 ENSMUSG0000024513 ENSMUSG0000024513 ENSMUSG0000034512 ENSMUSG0000034297 ENSMUSG00000034297 ENSMUSG0000039208 ENSMUSG0000039208	- RETRO RETRO - RETRO - RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - - - -	n/a - - p53 - - - - - -	n/a - - AMP - - AMP - AMP -	- DEL - DEL
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Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 MafX Map3k1 Map3k1 Mbd2 Mcl1 Mef2d Mef2d Metrn1 Mgat1 Mgat4a Midn	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG00000054720 ENSMUSG0000024305 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000029554 ENSMUSG00000018143 ENSMUSG00000024235 ENSMUSG0000024235 ENSMUSG00000024235 ENSMUSG00000034297 ENSMUSG00000034297 ENSMUSG0000003208 ENSMUSG00000026110 ENSMUSG00000026110	- RETRO RETRO - RETRO - RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - - - -	n/a - p53 - - - - - - - - - - - - -	n/a - - - - - - - - - - - - - - - - - - -	- DEL - DEL DEL - - - - - - - - -
Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 MafK Map3k1 Map3k8 Mbd2 Mc11 Mc11 Mc113 Mef2d Metrn1 Mgat1 Mgat4a Mix11	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000024720 ENSMUSG0000024305 ENSMUSG0000024305 ENSMUSG0000029554 ENSMUSG0000029554 ENSMUSG0000024513 ENSMUSG0000024513 ENSMUSG00000034297 ENSMUSG00000034297 ENSMUSG00000034297 ENSMUSG00000034208 ENSMUSG00000034287 ENSMUSG0000002346 ENSMUSG00000026497	- RETRO RETRO - RETRO - RETRO -			n/a - - - - -	n/a -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - - - -	n/a - - 53 - - - - - - - - - - - - - -	n/a - - - - - - - - - - - - - - - - - - -	- DEL - DEL DEL - - - - - - - - - - - - - - - - -
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Lmo2 Lrrc8c Lrrfip1 Lta Lyst Mad111 Mafk Map3k1 Map3k8 Mbd2 Mc11 Med13 Mef2d Mef2d Mef2d Met7d Mgat4a Midn Mix11 Mknk2 mmu-mir-142	ENSMUSG0000019920 ENSMUSG0000032698 ENSMUSG0000024720 ENSMUSG0000024720 ENSMUSG0000024402 ENSMUSG0000019726 ENSMUSG0000029554 ENSMUSG0000024253 ENSMUSG0000024235 ENSMUSG0000024235 ENSMUSG00000034297 ENSMUSG00000034297 ENSMUSG0000003208 ENSMUSG00000026110 ENSMUSG0000026110 ENSMUSG00000026110 ENSMUSG0000026110 ENSMUSG0000026110 ENSMUSG0000026110 ENSMUSG0000026120 ENSMUSG0000026120	- RETRO RETRO - RETRO - RETRO -		n/a - - - - - - - - - - - - - - - - - - -	n/a - - - - COSMIC - - - - - - - - - - - - - - - - - - -	n/a - Sjoblom - - - - - - - - - - - - - - - - - - -	- Nanog - - Nanog	- - - - - - - - - - - - - - - - - - -	n/a - p53 - - - - - - - - - - - - - - - - - - -	n/a - - - - - - - - - - - - - - - - - - -	- DEL - DEL DEL - - - - DEL - DEL - - n/a
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Nfkb1	ENSMUSG0000028163	RETRO	-	-	COSMIC	-	-	-	-	-	-
Nfkbil1	ENSMUSG0000042419	-	-	-	-	-	-	-	-	-	-
Nid1	ENSMUSG0000005397	-	-	-	-	-	-	-	-	AMP	-
Natah 1	ENEMUECOOOOOOOOOOO	DETRO	CD	Dam	COCMIC				p53		
Notch1	ENSMUSG0000026923	RETRO	SB	Dom Dom	COSMIC	-	-	-	(up)	AMP	-
NP_001074704.1 Nsmce1	ENSMUSG00000070576 ENSMUSG00000030750	-	-	Dom	-	-	-	-	-	-	-
Ntn1	ENSMUSG00000020902	_	_	-	-	-	Nanog	-	2	2	DEL
Nup210	ENSMUSG00000030091	-	-	-	COSMIC	-	-	-	-	-	-
Nup214	ENSMUSG00000001855	-	-	Dom	COSMIC	Sjoblom	-	-	_	AMP	-
Olfr56	ENSMUSG0000040328	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Orai2	ENSMUSG0000039747	RETRO	-	-	-	-	-	-	-	-	-
	ENSMUSG0000058287	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
	ENSMUSG0000052248	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
	ENSMUSG0000075416	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Ovca2	ENSMUSG0000038268	-	-	-	-	-	-	-	-	-	DEL
Padi2	ENSMUSG0000028927	RETRO	-	-	-	-	-	-	-	-	-
Pag1	ENSMUSG0000027508	-	-	-	-	-	-	-	-	-	-
Paics	ENSMUSG0000029247	-	-	-	-	-	-	-	-	-	-
Park7	ENSMUSG0000028964	-	-	-	-	-	-	-	-	-	-
Parvg	ENSMUSG0000022439	-	-	-	-	-	-	-	-	-	-
Pcgf5	ENSMUSG0000024805	-	-	-	COSMIC	-	-	-	p53	-	-
Pctk2	ENSMUSG0000020015	-	-	-	COSMIC	-	-	-	-	-	-
Pecam1	ENSMUSG0000020717	RETRO	-	-	-	-	Nanog	-	-	-	-
Phyhd1	ENSMUSG0000007476	-	-	-	-	-	-	-	-	AMP	-
Pigv	ENSMUSG0000043257	-	-	-	-	-	Nanog	-	-	-	-
Pik3cd	ENSMUSG0000039936		-	-	-	-	-	OCT4	-	-	-
Pik3r5 Pim1	ENSMUSG00000020901	RETRO	-	- Dom	- COEMIC	-	- Nanog	-	-	-	DEL
Pim1 Pitpnm2	ENSMUSG0000024014 ENSMUSG00000029406	RETRO	-	Dom	COSMIC	_	Nanog	-			-
Plac8	ENSMUSG0000029322	_	_	_	_		_	_	_	_	_
Plekha2	ENSMUSG00000031557	- RETRO	-	-	-	-	-	-			-
Plxnd1	ENSMUSG0000031337	-	_	_	_	_	_	_	-	-	-
Pml	ENSMUSG00000036986	_	_	Dom	COSMIC		_	OCT4	_	_	_
Ppp1r10	ENSMUSG0000039220	-	-	Dom	COSMIC	_	-	0014			-
Ppp1r16b	ENSMUSG0000037754	RETRO	_	_	-		_	_	_	_	DEL
Ppp2r5a	ENSMUSG00000026626	KLIKU	-	-	-	_	-	-		AMP	
Prdm16	ENSMUSG00000039410	RETRO	_	Dom	COSMIC		_	_	_	APT	_
Prkcbp1	ENSMUSG0000039410	RETRO	-	Dom	COSMIC	_	-	- OCT4			- DEL
Prkch	ENSMUSG00000021108	-	_	_	COSMIC		_	-	-	-	DLL
Prr6	ENSMUSG00000018509	_	_	_	-		_	_	_	_	DEL
Pscd4	ENSMUSG0000018008	_	_	_	_	_	_	_	-	-	-
Psma1	ENSMUSG00000030751	_	_	_	_	_	_	-			-
Psmb8	ENSMUSG00000024338	_	_	_	_	_	_	_	-	-	-
Ptbp1	ENSMUSG0000006498	RETRO	_	_	_	_	_	-			DEL
Ptp4a2	ENSMUSG00000028788	-	_	_	_	_	_	_	-	-	-
Ptp4a3	ENSMUSG00000059895	RETRO	-	-	-	-	-	-	-	-	-
Ptprc	ENSMUSG0000026395	-	-	-	COSMIC	_	-	-	-	AMP	-
ripic	EN3140300000020393				COSHIC				p53	APIF	
Ptpre	ENSMUSG0000041836	RETRO	_	_	_	_	_	_	(up)	_	_
Pvt1	ENSMUSG00000072566	-	_	n/a	n/a	n/a	_	_	n/a	n/a	n/a
Pxn	ENSMUSG00000029528	RETRO	_	- Ti/ d	-	11/ d	_	_	-	-	-
Pygm	ENSMUSG00000032648	-	_	_	_	_	_	_	-	-	-
Q8BP09_MOUSE	ENSMUSG00000041913	_	_	_	_		_	_	_	_	_
Rab37	ENSMUSG00000020732	_	_	_	_	_	Nanog	_	-	-	-
Ramp1	ENSMUSG0000034353	RETRO	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a
Rara	ENSMUSG0000037992	-	_	11/04		n, a		0074	n, u	-	-
Rasgrp1	ENG110500000007752			Dom	COSMIC	-	-		-		
Rasgrp2	ENSMUSG0000027347	RETRO		Dom -	COSMIC	-	-	OCT4 -	-	-	
Rassf2	ENSMUSG00000027347	RETRO	SB -	Dom - -	COSMIC - -	-	-	- -	-	-	-
	ENSMUSG0000032946	RETRO RETRO -		Dom - - -	COSMIC - - -		-	- - -	-	-	-
Rchth2	ENSMUSG0000032946 ENSMUSG00000027339			Dom - - -	-			- - - -		-	- - DEI
Rcbtb2 Rcsd1	ENSMUSG00000032946 ENSMUSG00000027339 ENSMUSG00000022106			Dom - - - -	COSMIC - - - COSMIC -			- - - - -		- - - - AMP	- - DEL -
Rcbtb2 Rcsd1 Recql5	ENSMUSG0000032946 ENSMUSG00000027339			Dom - - - - -	-	-		- - - - - -		- - - AMP -	- - DEL - -
Rcsd1 Recql5	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG0000040723 ENSMUSG00000020752			Dom - - - - - -	-	-	-	- - - - - - -		- - - - AMP - -	- - DEL - -
Rcsd1	ENSMUSG00000032946 ENSMUSG00000027339 ENSMUSG00000022106 ENSMUSG00000040723			Dom - - - - - - Dom	-	-	-	- - - - - - - -	· · · ·	- - - AMP - - -	- - DEL - - -
Rcsd1 Recql5 Rhbdf2	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG0000040723 ENSMUSG0000020752 ENSMUSG0000020806				-	-	-	- - - - - - - - - - - -	-	- - - - - - - - -	- - DEL - - - - -
Rcsd1 Recql5 Rhbdf2 Rhoh	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG000002752 ENSMUSG0000020752 ENSMUSG00000020806 ENSMUSG00000029204	RETRO - - - - - -			-	-	-	- - - - - - - - - - - -	-	- - - - - - - - - - -	- - DEL - - - - - -
Rcsd1 Recql5 Rhbdf2 Rhoh Rnf157	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG0000040723 ENSMUSG0000020752 ENSMUSG0000020806 ENSMUSG000002804 ENSMUSG0000052949	RETRO - - - - - -			-	-		- - - - - - - - - - - - - - -	-	- - - - - - - - - - - -	- DEL - - - - - - - - - -
Rcsd1 Recql5 Rhbdf2 Rhoh Rnf157 Rnf166	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG0000040723 ENSMUSG0000020752 ENSMUSG00000202806 ENSMUSG00000029204 ENSMUSG00000052949 ENSMUSG00000052949	RETRO - - - - - -			-	-		- - - - - - - - - - - - - - -	-	- - - - - - - - - - - - - - - - - -	- DEL - - - - - - - - - - - -
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Rcsd1 Recq/5 Rhbdf2 Rhbh Rnf157 Rnf166 Rnf43 Rorc Rpl11 Rpl24 Rps14 Rrs2 Rreb1 Rtn4rl1 Runx1 Runx1 Scd1 Scotin Scube1 Scyl1 Scdk1 Sema4b Sema4b Sema4b Sema4b Sema4b Sema4b Sema5 Sch305	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000027339 ENSMUSG00000240723 ENSMUSG0000020752 ENSMUSG0000029204 ENSMUSG00000029204 ENSMUSG00000029204 ENSMUSG00000028150 ENSMUSG00000028150 ENSMUSG00000028150 ENSMUSG00000028150 ENSMUSG00000024088 ENSMUSG00000045287 ENSMUSG00000045287 ENSMUSG0000002952 ENSMUSG0000002952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000022952 ENSMUSG00000025647 ENSMUSG00000026581 ENSMUSG0000002681 ENSMUSG0000002681 ENSMUSG0000002681 ENSMUSG00000024941 ENSMUSG0000002681 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG0000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG0000002451 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG00000024951 ENSMUSG000002451 ENSMUSG0000002456 ENSMUSG0000002456 ENSMUSG000002456 ENSMUSG000002456 ENSMUSG0000002456 ENSMUSG0000002456 ENSMUSG0000002456 ENSMUSG00000000059248 ENSMUSG0000002482 ENSMUSG0000002482 ENSMUSG0000002882	RETRO - - - RETRO - RETRO RETRO RETRO RETRO RETRO - RETRO RETRO - RETRO - RETRO - RETRO - - RETRO - - RETRO - - - RETRO - - - - - - - - - - - - -		- - - Dom - - - - - - - - - - - - - - - - - - -	COSMIC COSMIC - - - - - - - - - - - - - - - - - - -		- - - - Nanog - Nanog - Nanog			- - - - - - - - - - - - - - - - - - -	- - - - - - - - - - - - - - - - - - -
Rcsd1 Recq/5 Rhbdf2 Rhoh Rnf157 Rnf166 Rnf43 Rorc Rpl11 Rpl24 Rps14 Rrs21 Rth4rl1 Runx1 Runx3 Scd1 Scotin Scube1 Scyl1 Sdk1 Sell Sema4b Sema4b Sema4b Sept9 Serinc3 Set Sh3bp5 Sh3d19 Sirt2 Ski	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000027339 ENSMUSG0000022106 ENSMUSG0000022106 ENSMUSG0000029204 ENSMUSG0000029204 ENSMUSG00000029204 ENSMUSG0000029204 ENSMUSG0000029205 ENSMUSG00000029250 ENSMUSG0000024608 ENSMUSG00000024608 ENSMUSG00000024608 ENSMUSG00000024608 ENSMUSG0000002952 ENSMUSG00000045287 ENSMUSG00000045287 ENSMUSG00000025647 ENSMUSG00000026547 ENSMUSG00000026547 ENSMUSG00000026547 ENSMUSG00000026547 ENSMUSG00000026547 ENSMUSG00000026547 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021452 ENSMUSG00000021892 ENSMUSG00000017707	RETRO - - - RETRO - RETRO RETRO RETRO RETRO RETRO - RETRO RETRO - RETRO - RETRO - RETRO - - RETRO - - RETRO - - - RETRO - - - - - - - - - - - - -		- - - Dom - - - - - - - - - - - - - - - - - - -	COSMIC COSMIC - - - - - - - - - - - - - - - - - - -		- - - - Nanog - Nanog - Nanog			- - - - - - - - - - - - - - - - - - -	- - - - - - - - - - - - - - - - - - -
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Rcsd1 Recql5 Rhbdf2 Rhoh Rnf157 Rnf166 Rnf43 Rorc Rpl11 Rpl24 Rrs2 Rreb1 Rtn4rl1 Runx1 Runx3 Scd1 Scotin Scube1 Scyl1 Scotin Scube1 Scyl1 Scema4b Sema4b Sema4b Sema4b Sema4b Sema4b Set Sch3bp5 Sh3d19 Sirt2 Ski Sla	ENSMUSG0000032946 ENSMUSG0000027339 ENSMUSG0000027339 ENSMUSG0000020752 ENSMUSG0000020752 ENSMUSG0000029204 ENSMUSG00000029204 ENSMUSG00000029204 ENSMUSG0000014470 ENSMUSG00000028150 ENSMUSG00000028150 ENSMUSG00000024608 ENSMUSG00000024608 ENSMUSG00000024608 ENSMUSG00000024527 ENSMUSG00000045287 ENSMUSG00000025523 ENSMUSG00000025523 ENSMUSG00000025523 ENSMUSG00000025527 ENSMUSG00000025527 ENSMUSG00000025527 ENSMUSG00000025547 ENSMUSG00000025647 ENSMUSG00000026581 ENSMUSG00000026581 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000024766 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000024766 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG00000021451 ENSMUSG0000022322 ENSMUSG0000021451 ENSMUSG0000022372 ENSMUSG0000022372	RETRO - - - RETRO - RETRO RETRO RETRO RETRO RETRO - RETRO RETRO - RETRO RETRO - - RETRO - RETRO - - RETRO - - - - - - - - - - - - - - - - - - -		- - - Dom - - - - - - - - - - - - - - - - - - -	COSMIC COSMIC - - - - - - - - - - - - - - - - - - -		- - - - Nanog - Nanog - Nanog			- - - - - - - - - - - - - - - - - - -	- - - - - - - - - - - - - - - - - - -
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Slamf7	ENSMUSG0000038179	-	-	-	-	-	-	-	-	AMP	-
Slc1a3	ENSMUSG0000005360	-	-	-	-	-	-	-	-	-	-
Slc36a3	ENSMUSG0000049491	-	-	-	-	-	-	-	-	-	-
Slc38a1	ENSMUSG0000023169	RETRO	-	-	-	-	-	-	-	-	-
Slc39a13	ENSMUSG0000002105	-	_	_	_	_	_	_	_	_	_
		-	-	-	-	-	-	-	-	-	
Slc43a2	ENSMUSG0000038178	-	-	-	-	-	-	-	-	-	DEL
Smg6	ENSMUSG0000038290	RETRO	-	-	COSMIC	-	-	-	-	-	DEL
Sox19	ENSMUSG00000076431	RETRO	-	-	-	-	-	-	-	AMP	-
Spata13	ENSMUSG0000021990	-	-	-	COSMIC	-	-	-	-	-	-
Spn	ENSMUSG00000051457	-	-	-	-	-	-	-	-	-	-
Spsb4	ENSMUSG0000046997	-	-	-	-	-	-	-	-	-	-
Srgn	ENSMUSG00000020077	-	_	-	-	-	-	-	-	-	-
		DETRO									
Ssbp3	ENSMUSG0000061887	RETRO	-	-	-	-	-	-	-	-	-
St6galnac5	ENSMUSG0000039037	-	-	-	-	-	-	-	-	-	-
Stard10	ENSMUSG0000030688	-	-	-	-	-	-	-	-	-	-
Stard3nl	ENSMUSG0000003062	-	-	-	-	-	-	-	-	-	DEL
Stat5a	ENSMUSG0000004043	RETRO	-	-	-	-	-	-	-	-	-
Stat5b	ENSMUSG0000020919	RETRO	-	-	-	-	-	-	-	-	-
Stk4	ENSMUSG0000018209	-	-	-	-	-	-	-	-	-	DEL
Stmn1	ENSMUSG00000028832	-	-		-	-	-	-	-	-	-
Stra8	ENSMUSG00000029848	_	_	_	_	_	_	_	_	_	_
		-	-	-	-	-	-	-	-	-	-
Supt3h	ENSMUSG0000038954	-	-	-	-	-	-	-	-	-	-
Tap2	ENSMUSG0000024339	RETRO	-	-	-	-	-	-	-	-	-
Tbc1d1	ENSMUSG00000029174	-	-	-	-	-	Nanog	-	-	-	-
Tbxa2r	ENSMUSG0000034881	-	-	-	-	-	-	-	-	-	-
Tceb3	ENSMUSG0000028668	-	-	-	-	-	-	-	-	-	-
Tcf25	ENSMUSG0000001472	-	-	-	-	-	-	-	-	-	-
Tcf7	ENSMUSG0000000782	-	-	-	-	-	-	OCT4	-	-	-
	ENSMUSG00000005718	RETRO						0011			
Tcfap4		KLIKU	-	-	-	-	-	-	-	-	-
Tcof1	ENSMUSG0000024613	-	-	-	-	-	-	-	-	-	-
Tcte3	ENSMUSG0000036648	-	-	-	-	-	-	-	-	-	-
Tgfbr3	ENSMUSG0000029287	-	-	-	-	-	-	OCT4	-	-	-
Thra	ENSMUSG0000058756	-	-	-	-	-	-	-	-	-	-
Thy1	ENSMUSG0000032011	-	-	-	-	-	-	-	-	-	-
Tmem131	ENSMUSG00000026116	-	-	-	-	-	-	OCT4	-	-	-
	ENSMUSG0000024349							0014			
Tmem173		-	-	-	-	-	-	-	-	-	-
Tmem49	ENSMUSG0000018171	-	-	-	-	-	-	-	-	-	-
Tmem90a	ENSMUSG00000071234	-	-	-	-	-	-	-	-	-	-
Tmprss3	ENSMUSG0000024034	-	-	-	-	-	-	-	-	-	-
Tpd52	ENSMUSG0000027506	-	-	-	-	-	Nanog	-	-	-	-
Treh	ENSMUSG0000032098	-	-	-	-	-	-	-	-	-	DEL
Trim47	ENSMUSG0000020773	-	-	-	COSMIC	-	-	-	-	-	
Trp53inp1	ENSMUSG0000028211	_	_	_	-	_	_	_	_	_	_
TTP55IIIP1	LN3M03G0000028211	-	-		-	-	-	-	-	-	-
									p53		
Trpm1	ENSMUSG00000030523	-	-	-	-	-	-	-	(dn)	-	-
Tspan14	ENSMUSG0000037824	-	-	-	-	-	-	-	-	-	-
Tspan2	ENSMUSG0000027858	-	-	-	-	-	-	-	-	-	-
Ttll10	ENSMUSG00000029074	-								-	_
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Tuba8		-	-	-	-	-	-	-	2	AMP	_
Tuba8 Ubac2	ENSMUSG0000030137	-	-	-	-	-	-	-	-	AMP	-
Ubac2	ENSMUSG0000030137 ENSMUSG00000041765	-	-	-	-	-	-	-	-	AMP -	-
Ubac2 Ubash3a	ENSMUSG00000030137 ENSMUSG00000041765 ENSMUSG00000042345	- -	-	-	-	- - -	-	-	-	AMP - -	-
Ubac2 Ubash3a Ube1l	ENSMUSG00000030137 ENSMUSG00000041765 ENSMUSG00000042345 ENSMUSG00000032596	- - -	-	- - -	- - - -	-	-	-	-	AMP - - -	-
Ubac2 Ubash3a	ENSMUSG0000030137 ENSMUSG00000041765 ENSMUSG00000042345 ENSMUSG00000032596 ENSMUSG0000012126	-	-	-	-	-		-	-	AMP - - -	
Ubac2 Ubash3a Ube1l	ENSMUSG00000030137 ENSMUSG00000041765 ENSMUSG00000042345 ENSMUSG00000032596				- - - - - COSMIC	-	-	-	-	AMP - - - -	
Ubac2 Ubash3a Ube11 Ubxd5 Usp52	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG0000012126 ENSMUSG0000005682	- - - -	-		- - - - - COSMIC	-	-	- - - - - - OCT4	-	AMP - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG0000012126 ENSMUSG0000005682 ENSMUSG00000022710	- - - - - - RETRO	-	-	- - - - COSMIC -	-	-	- - - - - - OCT4	-	AMP - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7 Vamp8	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG0000012126 ENSMUSG0000005682 ENSMUSG00000022710 ENSMUSG0000050732	- - - - - RETRO		-	- - - - - COSMIC - -	-	-	- - - - - - - - - - - - - - - - - - -	-	AMP - - - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7 Vamp8 Vdac1	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG00000012126 ENSMUSG0000005682 ENSMUSG00000025710 ENSMUSG00000050732 ENSMUSG00000050732	- - - - - - RETRO -	-	-	- - - - - COSMIC - - -	-	-	- - - - - - - - - - - - - - - - - - -	-	AMP - - - - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7 Vamp8 Vdac1 Vil2	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG00000012126 ENSMUSG0000005682 ENSMUSG00000022710 ENSMUSG0000005732 ENSMUSG00000020402 ENSMUSG0000002397	- - - - - - RETRO - -	-	-	- - - - - - COSMIC - - -	-	-	- - - - - - - - - - - - - -	-	- - - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7 Vamp8 Vdac1 Vil2 Vpreb2	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG00000012126 ENSMUSG000000582 ENSMUSG00000022710 ENSMUSG0000005732 ENSMUSG0000023397 ENSMUSG00000052387	- - - - - - RETRO - - -	-	-	- - - - - - COSMIC - - - - - - -	-	-	- - - - - - OCT4 - - - -	-	AMP - - - - - - - - - - - - - - - - - - -	-
Ubac2 Ubash3a Ube11 Ubxd5 Usp52 Usp7 Vamp8 Vdac1 Vil2	ENSMUSG0000030137 ENSMUSG0000041765 ENSMUSG0000042345 ENSMUSG0000032596 ENSMUSG00000012126 ENSMUSG0000005682 ENSMUSG00000022710 ENSMUSG0000005732 ENSMUSG00000020402 ENSMUSG0000002397	- - - - - - RETRO - - - - -	-	-	- - - - - - COSMIC - - - - - - - - - - -	-	-	- - - - - - - - - - - - - -	-	- - - - - - - -	
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ENSMUSG0000074565	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a	
ENSMUSG0000076378	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a	_

Appendix C2. List showing other cancer-associated datasets in which the *Sleeping Beauty* CIS genes appear.

See Appendix C1 for an explanation of each column.

Gene name	Ensembl ID	RTCGD (retro)	RTCGD (SB)	CGC	COSMIC	Sjoblom	Nanog BS	OCT4 BS	p53 BS	Mulli Amps	ghan / Dels
Akt2	ENSMUSG0000004056	-	-	Dom	-	-	-	-	-	-	-
BC033915	ENSMUSG0000034135	-	-	-	COSMIC	Sjoblom	-	-	-	-	-
Erg	ENSMUSG0000040732	RETRO	SB	Dom	-	-	-	-	-	-	-
Fli1	ENSMUSG0000016087	RETRO	SB	Dom	COSMIC	-	-	-	-	-	-
Ikzf1	ENSMUSG0000018654	RETRO	-	Dom	-	-	-	-	-	-	-
Myb	ENSMUSG0000019982	RETRO	-	-	-	-	-	-	-	AMP	DEL
									p53		
Notch1	ENSMUSG0000026923	RETRO	SB	Dom	COSMIC	-	-	-	(up)	AMP	DEL
Ррр3са	ENSMUSG0000028161	-	-	-	COSMIC	-	-	-	-	-	-
Pten	ENSMUSG0000013663	-	-	Rec	COSMIC	-	-	-	-	-	DEL
	ENSMUSG0000075015	-	-	n/a	n/a	n/a	-	-	n/a	n/a	n/a

Appendix D. Nearest and further genes from CISs with a *P*-value of <0.001 or <0.05 in lists supplied by the Netherlands Cancer Institute.

"CIS chr" and "CIS position" are the chromosome and base pair coordinates of the CIS, which correspond to the centre of the peak in insertion density identified by the kernel convolution-based method.

CIS						
P- value	CIS chr	CIS position	Nearest gene name	Nearest gene Ensembl ID	Further gene name	Further gene Ensembl ID
0.001	1	37217235	Hearest gene name	ENSMUSG0000061123	Tmem131	ENSMUSG0000026116
0.001	1	37780290	Mgat4a	ENSMUSG0000026110	2010300C02Rik	ENSMUSG0000026090
0.001	1	86239852	1700019017Rik	ENSMUSG0000036574		ENSMUSG0000067125
0.001	1	86295450		ENSMUSG0000067125	OTTMUSG0000007209	ENSMUSG0000026238
0.001	1	90879243	Lrrfip1	ENSMUSG0000026305		ENSMUSG0000048068
0.001	1	91009545	Ramp1	ENSMUSG0000034353		ENSMUSG0000047217
0.001	1	125266838	Actr3	ENSMUSG0000026341	Slc35f5	ENSMUSG0000026342
0.001	1	128455260	Cxcr4	ENSMUSG0000045382		ENSMUSG0000064737
0.001	1	133917960	Btg2	ENSMUSG0000020423	Fmod	ENSMUSG0000041559
0.001	1	136100550	Camsap1l1	ENSMUSG0000041570	GPR25_MOUSE	ENSMUSG0000052759
0.001	1	137658090	A130050007Rik	ENSMUSG0000051480	—	ENSMUSG0000065205
0.001	1	156936780	1700057K13Rik	ENSMUSG0000026592	Angptl1	ENSMUSG0000033544
0.001	1	163991159	Sell	ENSMUSG0000026581	Selp	ENSMUSG0000026580
0.001	1	165628770	Creg1	ENSMUSG0000040713	Rcsd1	ENSMUSG0000040723
0.001	1	171607260	Cd48	ENSMUSG0000015355	Slamf7	ENSMUSG0000038179
0.001	1	171851676	Slamf6	ENSMUSG0000015314	Vangl2	ENSMUSG0000026556
0.001	1	172443511	Ccdc19	ENSMUSG0000026546		ENSMUSG0000066683
0.001	1	180313439	Itpkb	ENSMUSG0000038855	ENSMUSG0000056615	ENSMUSG0000056615
0.001	1	180660300	Mixl1	ENSMUSG0000026497	Acbd3	ENSMUSG0000026499
0.001	1	180803760	BC031781	ENSMUSG0000038806		ENSMUSG0000066654
0.001	1	191254946	Dtl	ENSMUSG0000037474	Ppp2r5a	ENSMUSG0000026626
0.001	2	6626911	Cugbp2	ENSMUSG0000002107		ENSMUSG0000069177
0.001	2	11543491		ENSMUSG0000069172	Rbm17	ENSMUSG0000037197
0.001	2	26397160	Notch1	ENSMUSG0000026923	Egfl7	ENSMUSG000002692
0.001	2	26493080	Egfl7	ENSMUSG0000026921	Notch1	ENSMUSG0000026923
0.001	2	28534226	Gfi1b	ENSMUSG0000026815	Gtf3c5	ENSMUSG0000026816
0.001	2	29999738	Set	ENSMUSG0000054766	Pkn3	ENSMUSG0000026785
0.001	2	30588260	Cstad	ENSMUSG0000047363	1700001022Rik	ENSMUSG0000044320
0.001	2	30652628	1700001022Rik	ENSMUSG0000044320	Cstad	ENSMUSG0000047363
0.001	2	31994682	A130092J06Rik	ENSMUSG0000050592	Nup214	ENSMUSG0000001855
0.001	2	32573666	Eng	ENSMUSG0000026814	Ak1	ENSMUSG0000026817
0.001	2	45044311	OTTMUSG0000012358	ENSMUSG0000052248		ENSMUSG0000057684
0.001	2	62252693	Dpp4	ENSMUSG0000035000	Gcg	ENSMUSG0000000394
0.001	2	65193998	9330158F14Rik	ENSMUSG0000061171		ENSMUSG0000060270
0.001	2	72102343		ENSMUSG0000068863	B230120H23Rik	ENSMUSG0000004085
0.001	2	90786925	Sfpi1	ENSMUSG0000002111	Slc39a13	ENSMUSG000000210
0.001	2	103644128	Lmo2	ENSMUSG0000032698		ENSMUSG0000056848
0.001	2	116858239	Rasgrp1	ENSMUSG0000027347	Thbs1	ENSMUSG0000040152
0.001	2	116924160	Rasgrp1	ENSMUSG0000027347	Thbs1	ENSMUSG0000040152
0.001	2	126843788	Dusp2	ENSMUSG0000027368	Stard7	ENSMUSG0000027367
0.001	2	131535747	Rassf2	ENSMUSG0000027339	Slc23a2	ENSMUSG0000027340
0.001	2	132024599	Prei4	ENSMUSG0000027346		ENSMUSG0000068247
0.001	2	152242778	Bcl2l1	ENSMUSG0000007659	Tpx2	ENSMUSG0000027469
0.001	2	156342081	Sla2	ENSMUSG0000027636	Ndrg3	ENSMUSG0000027634
0.001	2	158180458	2310007D09Rik	ENSMUSG0000027654	Ppp1r16b	ENSMUSG0000037754
0.001	2	165342117		ENSMUSG0000051789	Ncoa3	ENSMUSG0000027678
0.001	2	165410755	Ncoa3	ENSMUSG0000027678		ENSMUSG00000051789
0.001	2	166824814	SIc9a8	ENSMUSG0000039463	B4galt5	ENSMUSG00000017929
0.001	2	167210199	A530013C23Rik	ENSMUSG0000006462	Ptpn1	ENSMUSG00000027540
0.001	2	167357666	Ptpn1	ENSMUSG0000027540	A530013C23Rik	ENSMUSG0000006462
0.001	2	169676198	Zfp217	ENSMUSG00000052056		ENSMUSG0000055544
0.001	2	180161851	Slco4a1	ENSMUSG0000038963	mmu-mir-133a-2	ENSMUSG0000065460
0.001	3	29414978	Evi1	ENSMUSG0000027684	Mds1	ENSMUSG0000051636
0.001	3	29514295	Mds1	ENSMUSG0000051636	Evi1	ENSMUSG0000027684
0.001	3	51372385		ENSMUSG0000069019	Maml3	ENSMUSG0000061143
0.001	3	87886915	Mef2d	ENSMUSG0000001419		ENSMUSG0000068928
0.001	3	88367220	Arhgef2	ENSMUSG0000028059	Rxfp4	ENSMUSG0000004974
0.001	3	89142895	Zbtb7b	ENSMUSG0000028042	Flad1	ENSMUSG0000042642
0.001	3	93865375	Rorc	ENSMUSG0000028150	Lingo4	ENSMUSG000004450
0.001	3	95157949	Adamtsl4	ENSMUSG0000015850	Mcl1	ENSMUSG0000038612
0.001	3	95452585		ENSMUSG00000057781		ENSMUSG0000058910
0.001	3	100703412	Cd2	ENSMUSG0000027863	N	ENSMUSG000005046
0.001	3	102156745	Tspan2	ENSMUSG0000027858	Ngfb	ENSMUSG0000027859
0.001	3	103210135	Dclre1b	ENSMUSG0000027845		ENSMUSG0000068800
0.001	3	106883096	A930002I21Rik	ENSMUSG00000050179	AI504432	ENSMUSG000005614
0.001	3	114451945	Edg1	ENSMUSG0000045092	Olfm3	ENSMUSG000002796
0.001	3	129871645		ENSMUSG0000058642		ENSMUSG0000004962
0.001	3	129907165	NGL-L 1	ENSMUSG00000058642		ENSMUSG000004962
0.001	3	134486207	Nfkb1	ENSMUSG0000028163	ENSMUSG0000045520	ENSMUSG000004552
0.001	4	3880430	Chchd7	ENSMUSG0000042198		ENSMUSG000006139
0.001	4	11075624	Trp53inp1	ENSMUSG0000028211		ENSMUSG000005816
0.001	4	43376874	Cd72	ENSMUSG0000028459	T (1)	ENSMUSG000006619
0.001	4	46494350	5 //	ENSMUSG0000066194	Tbc1d2	ENSMUSG000003981
0.001	4	59383145	Rod1	ENSMUSG0000028382	A2AN91_MOUSE	ENSMUSG000003857
0.001	4	62489630	Whrn	ENSMUSG0000039137		ENSMUSG000003916
0.001	4	105793040	Ssbp3	ENSMUSG0000061887	Acot11	ENSMUSG000003485
0.001	4	117418370	Tie1	ENSMUSG0000033191	Mpl	ENSMUSG000000638
0.001	4	128586798		ENSMUSG0000066049	BC030183	ENSMUSG0000005049
11 101	4	128794580		ENSMUSG00000049089	Ptp4a2	ENSMUSG000002878
0.001 0.001	4	131936973	Fgr	ENSMUSG0000028874	Ahdc1	ENSMUSG000003769

0.001	4	132051188	ENSMUSG0000066041	ENSMUSG0000066041		ENSMUSG0000066040
0.001	4	132629392	Pigv	ENSMUSG0000043257	Arid1a	ENSMUSG0000007880
0.001 0.001	4 4	132779960 133074230	Ubxd5	ENSMUSG0000064705 ENSMUSG00000012126	Rps6ka1 Sh3bqrl3	ENSMUSG0000003644 ENSMUSG00000028843
0.001	4	134785370	Cnr2	ENSMUSG00000012120	Fuca1	ENSMUSG0000028843
0.001	4	134910709	Tceb3	ENSMUSG00000028668	Id3	ENSMUSG0000007872
0.001	4	134971837	Id3	ENSMUSG0000007872	Tceb3	ENSMUSG0000028668
0.001	4	135058700	E2f2	ENSMUSG0000018983	Ddefl1	ENSMUSG0000036995
0.001	4	139825430	Sdhb	ENSMUSG0000009863		ENSMUSG0000064443
0.001	4	148194053	D4Ertd429e	ENSMUSG00000044700	Pik3cd	ENSMUSG0000039936
0.001 0.001	5 5	32173336 63408106	Tacc3 Klf3	ENSMUSG00000037313 ENSMUSG00000029178	Fgfr3 Tbc1d1	ENSMUSG00000054252 ENSMUSG00000029174
0.001	5	64659750	Rhoh	ENSMUSG0000029204	/beiui	ENSMUSG0000067246
0.001	5	64720625	Chrna9	ENSMUSG0000029205	Rhoh	ENSMUSG0000029204
0.001	5	74573986		ENSMUSG0000067196	Kdr	ENSMUSG0000062960
0.001	5	74669557	Kdr	ENSMUSG0000062960	D/ 0	ENSMUSG0000067196
0.001 0.001	5 5	99594826 104706836	Lrrc8d	ENSMUSG0000067084 ENSMUSG00000046079	Plac8 Lrrc8c	ENSMUSG00000029322 ENSMUSG00000054720
0.001	5	106723992	Rpap2	ENSMUSG00000033773	Gfi1	ENSMUSG00000029275
0.001	5	106805480	Evi5	ENSMUSG00000011831	Gfi1	ENSMUSG0000029275
0.001	5	110555672	C130026L21Rik	ENSMUSG0000052848	Pitpnb	ENSMUSG0000050017
0.001	5	112943262	Coro1c	ENSMUSG0000004530	SelpIg	ENSMUSG0000048163
0.001	5	114275290	Cabp1	ENSMUSG0000029544	2410014A08Rik	ENSMUSG00000048578
0.001 0.001	5 5	114642496 116270566	Pxn Taok3	ENSMUSG00000029528 ENSMUSG00000061288	Pebp1	ENSMUSG0000064401 ENSMUSG00000032959
0.001	5	122293554	4932422M17Rik	ENSMUSG00000062946	геррі	ENSMUSG0000066795
0.001	5	122492536	Bcl7a	ENSMUSG0000029438	Wdr66	ENSMUSG0000029442
0.001	5	123399436		ENSMUSG0000029398	Pitpnm2	ENSMUSG0000029406
0.001	5	135184914	Prkrip1	ENSMUSG0000039737	Orai2	ENSMUSG0000039747
0.001	5	135637883	Mylc2pl	ENSMUSG0000005474	4731417B20Rik	ENSMUSG0000046548
0.001	5 5	136645576 137199166		ENSMUSG0000066690 ENSMUSG0000064676	00100011000	ENSMUSG0000007324 ENSMUSG00000050552
0.001 0.001	5	138379186	Gpr146	ENSMUSG00000044197	0910001L09Rik C130050O18Rik	ENSMUSG00000044092
0.001	5	138777046	Mafk	ENSMUSG0000018143	Ints1	ENSMUSG00000029547
0.001	5	139597070	Lfng	ENSMUSG0000029570	Grifin	ENSMUSG0000036586
0.001	5	146254722	Flt3	ENSMUSG0000042817		ENSMUSG0000057157
0.001	5	147866815		ENSMUSG0000066552	Hmgb1	ENSMUSG0000066551
0.001 0.001	5 6	148368619 29344053	4930588N13Rik BC048651	ENSMUSG0000029660	6330406I15Rik 2310016C08Rik	ENSMUSG00000029659 ENSMUSG00000043421
0.001	6	31202455	AB041803	ENSMUSG00000039742 ENSMUSG00000044471	2310010C08RIK	ENSMUSG00000052894
0.001	6	35041709	2010107G12Rik	ENSMUSG0000029847	Cnot4	ENSMUSG0000038784
0.001	6	48825985	Gimap6	ENSMUSG0000047867	Gimap4	ENSMUSG0000054435
0.001	6	52360435	Hoxa7	ENSMUSG0000038236	Hoxa9	ENSMUSG0000038227
0.001	6	52787635		ENSMUSG0000064792	Tax1bp1	ENSMUSG0000004535
0.001	6	53055274	Jazf1	ENSMUSG0000063568		ENSMUSG0000038157
0.001 0.001	6 6	99854785 121003391	Eif4e3 Cecr5	ENSMUSG00000030068 ENSMUSG00000058979		ENSMUSG0000024343 ENSMUSG00000040893
0.001	6	121362051	BC030863	ENSMUSG00000051586	Bid	ENSMUSG0000004446
0.001	6	125891365	Cd27	ENSMUSG0000030336		ENSMUSG0000064520
0.001	6	127827619	Ccnd2	ENSMUSG0000000184		ENSMUSG0000064191
0.001	6	127922935		ENSMUSG0000064191		ENSMUSG0000057412
0.001	6 6	128009665	Cd69	ENSMUSG00000058306	Clec2d	ENSMUSG0000067666
0.001 0.001	6	129919277 134950855	6009	ENSMUSG00000030156 ENSMUSG00000020939	Bcl2l14	ENSMUSG00000030157 ENSMUSG00000030200
0.001	6	147434665	4933424B01Rik	ENSMUSG00000040250	Deizhi i	ENSMUSG0000067323
0.001	7	19555460	Kcnn4	ENSMUSG0000054342	1500002020Rik	ENSMUSG0000002210
0.001	7	24194070		ENSMUSG0000047345	Sirt2	ENSMUSG0000015149
0.001	7	33353130	1600014C10Rik	ENSMUSG0000054676		ENSMUSG00000051425
0.001 0.001	7 7	40923804 62158275	C330024D12Rik	ENSMUSG00000039955 ENSMUSG00000030553	Ldha EG384639	ENSMUSG0000063229 ENSMUSG00000042668
0.001	7	67496820	C330024D12Kik	ENSMUSG0000065729	20304033	ENSMUSG00000042008
0.001	7	69487127		ENSMUSG00000025750	Akap13	ENSMUSG0000066406
0.001	7	73844970	Zfp710	ENSMUSG0000048897	2610034B18Rik	ENSMUSG0000039043
0.001	7	73943257	Sema4b	ENSMUSG0000030539	Idh2	ENSMUSG0000030541
0.001	7 7	74000850	Sema4b	ENSMUSG00000030539 ENSMUSG00000038797	Cib1	ENSMUSG0000030538
0.001 0.001	7	74632350 95190900	Zscan2 Fchsd2	ENSMUSG00000030691	Iqgap1 P2ry2	ENSMUSG00000030536 ENSMUSG00000032860
0.001	7	95447100	Stard10	ENSMUSG00000030688	Centd2	ENSMUSG0000032812
0.001	7	95667465		ENSMUSG0000066310	mmu-mir-139	ENSMUSG0000065446
0.001	7	108014978	Copb1	ENSMUSG0000030754	Rras2	ENSMUSG0000055723
0.001	7	108155370	Psma1	ENSMUSG0000030751	Pde3b	ENSMUSG0000030671
0.001 0.001	7 7	112137030 119538513	4930583K01Rik Nsmce1	ENSMUSG00000055159 ENSMUSG00000030750	Syt17 Il4ra	ENSMUSG00000058420 ENSMUSG00000030748
0.001	7	119676960	Gtf3c1	ENSMUSG0000032777	Il21r	ENSMUSG0000030745
0.001	7	120402570	Lat	ENSMUSG0000030742		ENSMUSG0000066180
0.001	7	120757800	Coro1a	ENSMUSG0000030707	Mapk3	ENSMUSG0000063065
0.001	7	121141669	AI467606	ENSMUSG0000045165	Qprt	ENSMUSG0000030674
0.001	7	121233233	Cd2bp2	ENSMUSG0000042502	Spn	ENSMUSG00000051457
0.001 0.001	7 7	121331984 123850980	Brwd2	ENSMUSG0000061513 ENSMUSG00000042055	Sephs2	ENSMUSG00000049091 ENSMUSG00000065763
0.001	7	129963457	Ptpre	ENSMUSG0000042033	Mki67	ENSMUSG00000031004
0.001	7	134601900	Olfr524	ENSMUSG00000050366	Cd163/1	ENSMUSG00000025461
0.001	7	135393702	Ifitm3	ENSMUSG0000025492	Ifitm1	ENSMUSG0000025491
0.001	7	139355962		ENSMUSG0000031071	Tpcn2	ENSMUSG00000048677
0.001	7	139459278	Toco 2	ENSMUSG00000031071	Tpcn2	ENSMUSG0000048677
0.001 0.001	7 8	139520040 10298663	Tpcn2 3930402G23Rik	ENSMUSG00000048677 ENSMUSG00000038917	Irs2	ENSMUSG0000031071 ENSMUSG00000038894
0.001	8	83003110		ENSMUSG0000064787	Cd97	ENSMUSG0000002885
0.001	8	83464133	mmu-mir-23a	ENSMUSG0000065611	mmu-mir-181c	ENSMUSG0000065483
0.001	8	83925867	Ier2	ENSMUSG0000053560	Cacna1a	ENSMUSG0000034656
0.001	8	83990843	Nfix	ENSMUSG0000001911	Dand5	ENSMUSG00000053226
0.001 0.001	8 8	104916233 110892610	2310066E14Rik Ldhd	ENSMUSG0000038604 ENSMUSG0000031958	Ctcf Znrf1	ENSMUSG00000005698 ENSMUSG00000033545
0.001	8 8	116868923	Plcg2	ENSMUSG0000034330	4632417N05Rik	ENSMUSG00000033343 ENSMUSG00000034308
0.001	8	119895988	1700016A09Rik	ENSMUSG00000042840	Gins2	ENSMUSG0000031821

0.001	8	121849515	Rnf166	ENSMUSG0000014470	2310061F22Rik	ENSMUSG0000049482
0.001	8	122086021	Cbfa2t3	ENSMUSG0000006362	BC021611	ENSMUSG0000015016
0.001	8	122807753	obideto	ENSMUSG00000057475	Def8	ENSMUSG0000001482
0.001	8	126128303	Tomm20	ENSMUSG00000058779	Delo	ENSMUSG0000069852
					DavialdE	
0.001	9	7246771	Mmp13	ENSMUSG0000050578	Dcun1d5	ENSMUSG0000032002
0.001	9	32432540		ENSMUSG0000064492	Ets1	ENSMUSG0000032035
0.001	9	32533657	Ets1	ENSMUSG0000032035		ENSMUSG0000064492
0.001	9	44013881	Thy1	ENSMUSG0000032011		ENSMUSG00000059979
0.001	9	44227331	Pdzd3	ENSMUSG0000032105	Cbl	ENSMUSG0000034342
0.001	9	44494631	Blr1	ENSMUSG0000047880	Bcl9l	ENSMUSG0000063382
0.001	9	44979071	Cd3d	ENSMUSG0000032094	Cd3e	ENSMUSG0000032093
0.001	9	57755625	Cyp1a1	ENSMUSG0000032315	Csk	ENSMUSG0000032312
	9	69594821			Foxb1	
0.001			Anxa2	ENSMUSG0000032231		ENSMUSG0000059246
0.001	9	71304551	Aqp9	ENSMUSG0000032204	Aldh1a2	ENSMUSG0000013584
0.001	9	72524010	Mns1	ENSMUSG0000032221	Zfp280d	ENSMUSG0000038535
0.001	9	108045941	Ube1/	ENSMUSG0000032596	6230427J02Rik	ENSMUSG0000042106
0.001	9	109100985	Trex1	ENSMUSG00000049734	Scotin	ENSMUSG0000025647
0.001	9	110949851	Als2cl	ENSMUSG0000044037	Tdgf1	ENSMUSG0000032494
0.001	9	112261091	mmu-mir-128b	ENSMUSG0000065441	Pdcd6ip	ENSMUSG0000032504
0.001	9	114407511	Glb1	ENSMUSG0000045594	Ccr4	ENSMUSG0000047898
0.001	9	123784927	Ccr9	ENSMUSG0000029530	Lztfl1	ENSMUSG0000025245
0.001	10	19508682	Ifngr1	ENSMUSG0000020009	Olig3	ENSMUSG0000045591
0.001	10	20924120	Ahi1	ENSMUSG0000019986	Myb	ENSMUSG0000019982
0.001	10	21009938	Myb	ENSMUSG00000019982	Ahi1	ENSMUSG0000019986
					AIIII	
0.001	10	21125838	МуЬ	ENSMUSG0000019982	A4	ENSMUSG0000059894
0.001	10	21171964	CL 22.4C	ENSMUSG0000059894	Myb	ENSMUSG0000019982
0.001	10	40527010	Slc22a16	ENSMUSG0000019834	Cdc2l6	ENSMUSG0000038481
0.001	10	60125292	Chst3	ENSMUSG0000057337	Spock2	ENSMUSG0000058297
0.001	10	62471532	Srgn	ENSMUSG0000020077	2510003E04Rik	ENSMUSG0000036955
0.001	10	80050543	Arid3a	ENSMUSG0000019564	Wdr18	ENSMUSG0000035754
0.001	10	80284284	Cirbp	ENSMUSG0000045193	Midn	ENSMUSG0000035621
0.001	10	80817672	Mobkl2a	ENSMUSG0000003348	NP 082105.3	ENSMUSG0000055862
0.001	10	81464292	Tbxa2r	ENSMUSG0000034881	Gipc3	ENSMUSG0000034872
0.001	10	81637110	Gna15	ENSMUSG0000034792	Edg6	ENSMUSG0000044199
0.001	10	93100182	Pctk2	ENSMUSG00000020015	4930485B16Rik	ENSMUSG0000020014
0.001	10	120266886	4921513I03Rik	ENSMUSG00000044544	1700006J14Rik	ENSMUSG0000034764
0.001	10	128040691	Usp52	ENSMUSG0000005682		ENSMUSG0000064948
0.001	11	5403337	Xbp1	ENSMUSG0000020484	Znrf3	ENSMUSG0000041961
0.001	11	11610839	Ikzf1	ENSMUSG0000018654	Fignl1	ENSMUSG00000035455
0.001	11	18916027	Meis1	ENSMUSG0000020160		ENSMUSG0000069956
0.001	11	22705312		ENSMUSG0000062899	B3gnt2	ENSMUSG0000051650
0.001	11	48835439		ENSMUSG0000040335	Olfr1396	ENSMUSG0000047511
0.001	11	52072054	Vdac1	ENSMUSG0000020402	Tcf7	ENSMUSG0000000782
0.001	11	54874286	10001	ENSMUSG00000044751	Slc36a3	ENSMUSG0000049491
0.001	11	58159185	OTTMUSG0000005737	ENSMUSG00000058287	1810065E05Rik	ENSMUSG0000013653
0.001	11	62272627	Ubb	ENSMUSG0000019505	Prr6	ENSMUSG0000018509
0.001	11	67925874	Ntn1	ENSMUSG0000020902	Stx8	ENSMUSG0000020903
0.001	11	68156194	Pik3r5	ENSMUSG0000020901	Ntn1	ENSMUSG0000020902
0.001	11	72682830	Atp2a3	ENSMUSG0000020788		ENSMUSG0000065298
0.001	11	74740528		ENSMUSG0000061832	Smg6	ENSMUSG0000038290
0.001	11	74900917	Ovca2	ENSMUSG0000038268	mmu-mir-132	ENSMUSG0000065537
0.001	11	75004843	Rtn4rl1	ENSMUSG0000045287	Rpa1	ENSMUSG0000000751
0.001	11	77243999	1300007F04Rik	ENSMUSG0000000686	Taok1	ENSMUSG0000017291
0.001	11	78730665	Ksr1	ENSMUSG0000018334	Lgals9	ENSMUSG0000001123
0.001	11	79250571	1011	ENSMUSG00000046628	Rab11fip4	ENSMUSG0000017639
0.001	11	82817405		ENSMUSG00000035163	Ap2b1	ENSMUSG00000035152
0.001	11	86133847		ENSMUSG0000064526	Med13	ENSMUSG0000034297
			mmu mir 21		Medij	
0.001	11	86318401	mmu-mir-21	ENSMUSG0000065455	D/ 10	ENSMUSG0000069780
0.001	11	86624827	Ypel2	ENSMUSG0000018427	Dhx40	ENSMUSG0000018425
0.001	11	87475267	mmu-mir-142	ENSMUSG0000065420	Supt4h1	ENSMUSG0000020485
0.001	11	88896397	A930013B10Rik	ENSMUSG0000063109	Nog	ENSMUSG0000048616
0.001	11	98296507		ENSMUSG0000065891	Med24	ENSMUSG0000017210
0.001	11	98568907	Thra	ENSMUSG00000058756	Nr1d1	ENSMUSG0000020889
0.001	11	98766517		ENSMUSG0000064491	Rara	ENSMUSG0000037992
0.001	11	98977357	Ccr7	ENSMUSG0000037944	Smarce1	ENSMUSG0000037935
0.001	11	100225567	1110036003Rik	ENSMUSG0000006931	Jup	ENSMUSG0000001552
0.001	11	100668367	Stat5a	ENSMUSG0000004043	Stat5b	ENSMUSG0000020919
0.001	11	100709155	Stat3	ENSMUSG0000004040	Ptrf	ENSMUSG0000004044
0.001	11	102995551	Fmnl1	ENSMUSG00000055805	1700023F06Rik	ENSMUSG0000020940
0.001	11	106494037	Pecam1	ENSMUSG0000020717	Gm885	ENSMUSG0000040528
0.001	11	114981951	Slc9a3r1	ENSMUSG0000020733	Cd300lf	ENSMUSG0000047798
0.001	11	115250587	Kctd2	ENSMUSG00000016940	Armc7	ENSMUSG00000057219
0.001	11				Recal5	
0.001	11	115723675	2210020M01Rik	ENSMUSG00000048442 ENSMUSG00000052791	Кесція	ENSMUSG00000020752 ENSMUSG00000020786
		116181215	B (157		111001 1/000:1	
0.001	11	116238127	Rnf157	ENSMUSG0000052949	1110014K08Rik	ENSMUSG0000050628
0.001	11	116424757	Rhbdf2	ENSMUSG0000020806	Cygb	ENSMUSG0000020810
0.001	11	117148567	Sept9	ENSMUSG0000059248		ENSMUSG0000021618
0.001	11	117605527	Tmc8	ENSMUSG0000050106	6030468B19Rik	ENSMUSG0000025573
0.001	11	119663437	Chmp6	ENSMUSG0000025371		ENSMUSG0000056397
0.001	12	12295330	Mycn	ENSMUSG0000037169		ENSMUSG0000048957
0.001	12	52103161		ENSMUSG0000019260	1110008L16Rik	ENSMUSG0000021023
0.001	12	70425031	Prkch	ENSMUSG0000021108	Tmem30b	ENSMUSG0000034435
0.001	12	82414330	Jundm2	ENSMUSG0000034271	Fos	ENSMUSG0000021250
0.001	12	82502328	Jundm2	ENSMUSG0000034271	Batf	ENSMUSG0000034266
0.001	12	83703031		ENSMUSG00000061115		ENSMUSG0000059114
0.001	12	102594114	Bel11b	ENSMUSG00000057566	Cotd2	ENSMUSG0000064099
0.001	12	103397701	Bcl11b	ENSMUSG00000048251	Setd3	ENSMUSG00000056770
0.001	12	106174472	Hsp90aa1	ENSMUSG0000021270	Wdr20a	ENSMUSG0000037957
0.001	12	108155761	EG382639	ENSMUSG0000037638	Akt1	ENSMUSG0000001729
0.001	12	108336933	Gpr132	ENSMUSG0000021298	Jag2	ENSMUSG0000002799
0.001	13	18833698	Stard3nl	ENSMUSG0000003062		ENSMUSG0000003061
0.001	13	23035361	Hist1h1d	ENSMUSG0000052565	Hist2h3c2	ENSMUSG0000069273
0.001	13	28426378		ENSMUSG0000043626		ENSMUSG0000069256
0.001	13	30230008	Irf4	ENSMUSG0000021356	Exoc2	ENSMUSG0000021357
0.001	13	30308308	Exoc2	ENSMUSG0000021357		ENSMUSG0000069254

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0.001 13 1086.1599 Risk ENAMUSCO00006420 ENAMUSC000006460 ENAMUSC0000006460 0.001 14 2389272 Zink1 ENAMUSC000007317 Pol/I ENAMUSC0000007317 0.001 14 2389272 Zink1 ENAMUSC000007317 Pol/I ENAMUSC0000007318 0.001 14 2389272 Zink1 ENAMUSC000007317 Pol/I ENAMUSC0000007317 0.001 14 6373894 Grad ENAMUSC0000007317 Press/USC000007317 Press/USC0000007317 0.001 14 67472035 Kchh2 ENAMUSC0000007317 Sup1 ENAMUSC00000073147 0.001 14 7387384 Fress/USC000005739 Sup1 ENAMUSC00000072145 0.001 14 11352827 Can100 ENAMUSC0000007215 Sup1 ENAMUSC0000007215 0.001 15 334319 Can10 ENAMUSC0000007215 Sup1 ENAMUSC0000007215 0.001 15 252387 Can10 ENAMUSC0000007215 Sup1 ENAMUSC0000000145	0.001	13	61798581	Ccrk	ENSMUSG0000021483	Ctsl	ENSMUSG0000021477
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0.001 19 40571759 Dntt ENSMUSG0000025014 Blnk ENSMUSG0000061132 0.001 19 41384622 Frat2 ENSMUSG0000047604 ENSMUSG00000067199 0.001 20 6105829 ENSMUSG00000031162 Pim2 ENSMUSG00000031155 0.001 20 6193879 Gata1 ENSMUSG0000031162 Hdac6 ENSMUSG0000031161 0.001 20 42949129 Elf4 ENSMUSG0000005456 ENSMUSG00000046863 0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG00000046863	0.001		36835733	Hhex	ENSMUSG0000024986	Exoc6	ENSMUSG0000053799
0.001 19 41384622 Frat2 ENSMUSG0000047604 ENSMUSG0000067199 0.001 20 6105829 ENSMUSG00000050227 Pim2 ENSMUSG0000031155 0.001 20 6193879 Gata1 ENSMUSG0000031162 Hdac6 ENSMUSG0000031161 0.001 20 42949129 Elf4 ENSMUSG0000065456 ENSMUSG00000046863 0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG0000046863	0.001	19	40571759	Dntt	ENSMUSG0000025014	Blnk	ENSMUSG0000061132
0.001 20 6105829 ENSMUSG0000050227 Pim2 ENSMUSG0000031155 0.001 20 6193879 Gata1 ENSMUSG00000031162 Hdac6 ENSMUSG00000031161 0.001 20 42949129 Elf4 ENSMUSG0000005455 ENSMUSG0000003512 0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG00000046863							
0.001 20 6193879 Gata1 ENSMUSG0000031162 Hdac6 ENSMUSG00000031161 0.001 20 42949129 Elf4 ENSMUSG0000031103 Q8C3U4_MOUSE ENSMUSG00000053512 0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG0000046863						Pim2	
0.001 20 42949129 Elf4 ENSMUSG0000031103 Q8C3U4_MOUSE ENSMUSG00000053512 0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG00000046863				Gata1			
0.001 20 47271754 mmu-mir-106a ENSMUSG0000065456 ENSMUSG00000046863							
						20000 I_NOUSE	
	0.001	20	JUL 1 JJLL	SHOPE	Ensines0000001330		2.13.1333300000000041

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0.05	7	136753950	6330512M04Rik	ENSMUSG00000045777	Tofref26	ENSMUSG0000066099
0.05 0.05	7 8	137972370 21562896	<i>Cars 1700041G16Rik</i>	ENSMUSG00000010755 ENSMUSG00000054822	Tnfrsf26 Myst3	ENSMUSG00000045362 ENSMUSG00000031540
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0.05	9	21207041	Slc44a2	ENSMUSG00000057193	Ap1m2	ENSMUSG0000003309

			- /			
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0.05	10	93279846	Elk3	ENSMUSG0000008398	Lta4h	ENSMUSG0000015889
0.05	10	117780612		ENSMUSG0000069509	Rap1b	ENSMUSG00000052681
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0.05	10	128150682	Rnf41	ENSMUSG0000025373	Smarcc2	ENSMUSG00000025369
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0.05	11	18767407	Meis1	ENSMUSG0000020160		ENSMUSG0000069957
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0.05	17	51109845	Pcaf	ENSMUSG0000000708		ENSMUSG0000063839
0.05	17	63594898	Twsg1	ENSMUSG0000024098	Ralbp1	ENSMUSG0000024096
0.05	17	69257535	Ndc80	ENSMUSG0000024056	Q8BP09_MOUSE	ENSMUSG0000041913
0.05	18	39246982	Arhgap26	ENSMUSG0000036452	Nr3c1	ENSMUSG0000024431
0.05	18	50244433	Tnfaip8	ENSMUSG0000062210		ENSMUSG0000069381
0.05	18	60980215	Rps14	ENSMUSG0000024608	Ndst1	ENSMUSG0000054008
0.05	18	65140045	Nedd4l	ENSMUSG0000024589	A330084C13Rik	ENSMUSG0000055362
0.05	18	80828071	Atp9b	ENSMUSG0000024566	Nfatc1	ENSMUSG0000033016
0.05	19	3938519	Coro1b	ENSMUSG0000024835	Rps6kb2	ENSMUSG0000024830
0.05	19	4011389	Clcf1	ENSMUSG0000040663	Pold4	ENSMUSG0000024854
0.05	19	5629379	Frmd8	ENSMUSG0000043488	Scyl1	ENSMUSG0000024941
0.05	19	7953175	Nxf1	ENSMUSG0000010097	Stx5a	ENSMUSG0000010110
0.05	19	8242954	Scgb1a1	ENSMUSG0000024653		ENSMUSG0000059508
0.05	19	9780269	Cybasc3	ENSMUSG0000034445	Dak	ENSMUSG0000034371
0.05	19	24225959	Dock8	ENSMUSG0000052085		ENSMUSG0000064896
0.05	19	31586856	2700046G09Rik	ENSMUSG0000024893	Sgms1	ENSMUSG0000040451
0.05	19	32424629		ENSMUSG0000052009	Pten	ENSMUSG0000013663
0.05	19	35598899	Pcgf5	ENSMUSG0000024805	Ankrd1	ENSMUSG0000024803
0.05	19	43928459	Scd1	ENSMUSG0000037071	Scd4	ENSMUSG0000050195
0.05	19	45077552	Poll	ENSMUSG0000025218		ENSMUSG0000062336
0.05	19	46522796	Ina	ENSMUSG0000034336	Q99JM3_MOUSE	ENSMUSG0000025039
0.05	19	46999894	Obfc1	ENSMUSG0000042694	Sh3pxd2a	ENSMUSG0000053617
0.05	19	53470079	Pdcd4	ENSMUSG0000024975	Rbm20	ENSMUSG0000043639
0.05	20	11481710		ENSMUSG0000058791	Ddx3x	ENSMUSG0000000787
0.05	20	42661369	1200013B08Rik	ENSMUSG0000031101	Xpnpep2	ENSMUSG0000037005
0.05	20	128113669	Gla	ENSMUSG0000031266	Btk	ENSMUSG0000031264

Appendix E. Human cancer cell lines used in the 10K and SNP 6.0 CGH analyses.

SNP6.0? indicates whether the cell line was used in the higher resolution SNP 6.0 CGH analysis. Abbreviations: auto_ganglia = autonomic ganglia; CNS = central nervous system; haem_and_lymph = haematopoietic and lymphoid tissue.

unknown(...) is used where the cell line was classified as unknown in the 10K analysis, but was later identified for the SNP 6.0 analysis.

Cell line	Tissue	SNP6.0?	Cell line	Tissue	SNP6.0?	Cell line	Tissue	SNP6.0
SW13	adrenal_gland	Y	MOLT-4	haem_and_lymph	Y	SCLC-21H	lung	Y
-			MONO-	····				
GI-CA-N	auto_ganglia		MAC-6	haem_and_lymph	Y	SHP-77	lung	Y
KP-N-YS	auto_ganglia	Y	Mo-T	haem_and_lymph	Y	SK-LU-1	lung	Y
NB-10	auto_ganglia	Ŷ	MUTZ-1	haem_and_lymph	Y	SK-MES-1	lung	Ŷ
NB-13	auto_ganglia	Ý	MV-4-11	haem_and_lymph		SW1573	lung	Ý
		Y			Y			Y
NB-14	auto_ganglia		NALM-1	haem_and_lymph		SW900	lung	
NB-17	auto_ganglia	Y	NALM-6	haem_and_lymph	Y	VMRC-LCP	lung	Y
NB-5	auto_ganglia	Y	NC-37	haem_and_lymph		COLO-680N	oesophagus	Y
NB-6	auto_ganglia	Y	NKM-1	haem_and_lymph	Y	EC-GI-10	oesophagus	Y
NB-7	auto_ganglia	Y	NOMO-1	haem_and_lymph	Y	HCE-4	oesophagus	Y
ACN	auto_ganglia	Y	OCI-AML2	haem_and_lymph	Y	KYSE-140	oesophagus	
BE2-C	auto_ganglia		OPM-2	haem_and_lymph	Y	KYSE-150	oesophagus	
	_5 5		P12-	/ .				
CHP-126	auto_ganglia	Y	ICHIKAWA	haem_and_lymph	Y	KYSE-180	oesophagus	
CHP-212	auto_ganglia	Ý	P30-OHK	haem_and_lymph	Ŷ	KYSE-30	oesophagus	
GI-LI-N	auto_ganglia		P31-FUJ	haem_and_lymph	Ý	KYSE-520	oesophagus	Y
		V			I			1
GOTO-P3	auto_ganglia	Y	PF-382	haem_and_lymph		KYSE-70	oesophagus	
KP-N-RT-								
BM-1	auto_ganglia	Y	QIMR-WIL	haem_and_lymph	Y	OE19	oesophagus	Y
KP-N-S19s	auto_ganglia	Y	Raji	haem_and_lymph	Y	OE33	oesophagus	Y
			Ramos-					
LAN-5	auto_ganglia	Y	2G6-4C10	haem_and_lymph	Y	TE-1	oesophagus	Y
LAN-6	auto_ganglia	Y	REH	haem_and_lymph	Ý	TE-10	oesophagus	Ý
MC-IXC	auto_ganglia	Y	RL	haem_and_lymph	Y	TE-11	oesophagus	Y
MHH-NB-11	auto_ganglia	Y	RPMI-6666	haem_and_lymph	Y	TE-12	oesophagus	Y
VB16-								
RIKEN	auto_ganglia	Y	RPMI-8226	haem_and_lymph	Y	TE-15	oesophagus	Y
NH-12	auto_ganglia	Y	RPMI-8402	haem_and_lymph	Y	TE-5	oesophagus	Y
VH-6	auto_ganglia	Y	RPMI-8866	haem_and_lymph	Y	TE-6	oesophagus	Y
SCCH-26	auto_ganglia	Ŷ	RS4-11	haem and lymph	Ŷ	TE-8	oesophagus	Ý
SIMA	auto_ganglia	Ý	SCC-3	haem_and_lymph	Ŷ	TE-9	oesophagus	Ŷ
		Y		haem and lymph		-		
SK-N-AS	auto_ganglia		SIG-M5		Y	41MCISR	ovary	
SK-N-DZ	auto_ganglia	Y	SKM-1	haem_and_lymph	Y	Caov-3	ovary	Y
rgw	auto_ganglia	Y	SK-MM-2	haem_and_lymph	Y	EFO-21	ovary	Y
GI-1	biliary_tract	Y	SKW	haem_and_lymph		EFO-27	ovary	Y
TK-1	biliary_tract	Y	SKW-3	haem and lymph		IGROV-1	ovary	Y
HuCCT1	biliary_tract	Y	ST486	haem_and_lymph	Y	KGN	ovary	Y
luH-28	biliary_tract	Ý	SU-DHL-1	haem_and_lymph	Y	KURAMOCHI	ovary	Ŷ
IGBC1TKB		Ý	SUP-B8		Ý	OAW28		Ý
	biliary_tract			haem_and_lymph			ovary	
GBC24TKB	biliary_tract	Y	SUP-T1	haem_and_lymph	Y	OAW-42	ovary	Y
CADO-ES1	bone	Y	TALL-1	haem_and_lymph	Y	OC-314	ovary	Y
CAL-72	bone	Y	THP-1	haem_and_lymph		OVCAR3	ovary	Y
ES-1	bone	Y	TUR	haem_and_lymph	Y	OVCAR-4	ovary	Y
S-4	bone	Y	U-266	haem_and_lymph	Y	OVCAR-5	ovary	Y
EW-1	bone	Y	U-698-M	haem_and_lymph	Y	PA-1	ovary	Y
W-12	bone	Ŷ	WSU-NHL	haem_and_lymph	Ŷ	RMG-I	ovary	Ŷ
W-12	bone	Ý	YT		Ý	RTSG		Ý
				haem_and_lymph	I		ovary	
W-16	bone	Y	NB4	haem_and_lymph		SK-OV-3	ovary	Y
EW-18	bone		769-P	kidney	Y	TYK-nu	ovary	Y
W-22	bone	Y	786-0	kidney	Y	AsPC-1	pancreas	Y
W-24	bone	Y	A498	kidney	Y	BxPC-3	pancreas	Y
W-3	bone	Y	ACHN	kidney	Y	Capan-1	, pancreas	
W-7	bone	Ŷ	BB65-RCC	kidney	Ŷ	Capan-2	pancreas	Y
I-EMC-SS	bone	Ý	BFTC-909	kidnev	Ŷ	CFPAC-1	pancreas	Ŷ
10S	bone	Y	CAKI-1	kidney	Y	HPAF-II		Y
			-				pancreas	
luO9	bone	Y	CAL-54	kidney	Y	HuP-T3	pancreas	Y
1G-63	bone	Y	HA7-RCC	kidney	Y	HuP-T4	pancreas	Y
			LB1047-			1		
NOS-1	bone	Y	RCC	kidney	Y	MIA-PaCa-2	pancreas	
			LB2241-				-	
١Y	bone	Y	RCC	kidney	Y	MZ1-PC	pancreas	
Saos-2	bone	Ý	LB996-RCC	kidney	Ý	PANC-03-27	pancreas	Y
SJSA-1		Y			•		•	Y
	bone		OS-RC-2	kidney		PANC-10-05	pancreas	
SK-PN-DW	bone	Y	RCC10RGB	kidney		SW1990	pancreas	Y
J-2-0S	bone	Y	RXF393	kidney	Y	YAPC	pancreas	Y
3T-20	breast		SK-NEP-1	kidney	Y	JAR	placenta	Y
3T-474	breast	Y	SN12C	kidney	Y	JEG-3	placenta	Y
3T-549	breast	Ŷ	SW156	kidney		IST-MES1	pleura	Ŷ
CAL-120	breast	Ý	TK10	kidney	Y	MPP-89	pleura	Ý
		Y			r Y			r Y
CAL-148	breast		U031	kidney		MSTO-211H	pleura	
CAL-51	breast	Y	VMRC-RCZ	kidney	Y	NCI-H2452	pleura	Y
			VMRC-					
CAL-85-1	breast	Y	MELG	large intestine	Y	NCI-H28	pleura	Y
CAMA-1	breast	Ý	293	large_intestine		BPH-1	prostate	Ŷ
COLO-824	breast	Ý	C2BBe1	large_intestine	Y	DU-145	prostate	Ý
DU-4475	breast	Y	CA46	large_intestine	Y	A101D	skin	Y
	breast	Y	COLO-205	large_intestine	Y	A375	skin	Y
EVSA-T HCC1143	breast	Y	COLO-320	large_intestine	Y	A4-Fuk	skin	Y

HCC1187	breast	Y	COLO-678	large_intestine	Y	AKI	skin	
HCC1395	breast	Y	COLO-741	large_intestine	Y	BB132-MEL	skin	
HCC1419	breast		CW-2	large_intestine	Ŷ	BSCC-93	skin	
		Y	GP5d		Y			v
HCC1569	breast			large_intestine		C32	skin	Y
HCC1599	breast	Y	H4	large_intestine	Y	CAL-39	skin	Y
HCC1806	breast	Y	HCC2998	large_intestine	Y	CHL-1	skin	Y
HCC1937	breast	Y	HCT-116	large intestine	Y	COLO-679	skin	Y
HCC1954	breast	Y	HT-115	large_intestine		COLO-792	skin	Y
HCC2157		Ŷ	HT-29		Y		skin	Ý
	breast			large_intestine		COLO-800		T
HCC2218	breast	Y	HT55	large_intestine	Y	COLO-853	skin	
HCC38	breast	Y	LoVo	large_intestine	Y	CP50-MEL-B	skin	Y
HCC70	breast	Y	LS-123	large_intestine	Y	CP66-MEL	skin	Y
								Ý
Hs-578-T	breast	Y	LS-174T	large_intestine	Y	G-361	skin	Ŷ
MCF7	breast	Y	LS-411N	large_intestine	Y	GR-M	skin	
MDA-MB-								
134-VI	breast	Y	LS-513	large_intestine	Y	HMV-II	skin	Y
MDA-MB-								
						100.4		
157	breast	Y	NCI-H508	large_intestine	Y	IGR-1	skin	Y
MDA-MB-								
175-VII	breast	Y	NCI-H630	large_intestine	Y	IPC-298	skin	Y
MDA-MB-				5 =				
231	breast	Y	NCI-H716	large_intestine	Y	IST-MEL1	skin	Y
	Diedst	1	NCI-11/10	large_incestine	1	131-MLLI	36111	
MDA-MB-								
361	breast	Y	NCI-H747	large_intestine	Y	LB2518-MEL	skin	Y
MDA-MB-						LB2531-		
415	breast	Y	RCM-1	large_intestine	Y	MEL-Z	skin	
	brease	•		large_incestine	•		Skill	
MDA-MB-								
453	breast	Y	RKO	large_intestine	Y	LB33-MEL-A	skin	
MDA-MB-						LB373-MEL-		
468	breast		SK-CO-1	large intestine	Y	D	skin	Y
			SNU-C1	large intestine	Y		skin	•
MFM-223	breast					Malme-3M		
MRK-nu-1	breast		SNU-C2B	large_intestine	Y	MEL-HO	skin	
NCI-ADR-								
RES	breast		SW1116	large intestine	Y	MEL-JUSO	skin	
OCUB-M	breast	Y	SW1417	large_intestine	Ŷ	MLMA	skin	Y
		•						
SK-BR-3	breast		SW403	large_intestine	Y	MMAC-SF	skin	Y
SW527	breast		SW48	large_intestine	Y	MZ2-mel	skin	
T47D	breast	Y	SW620	large_intestine	Y	RVH-421	skin	Y
UACC-812	breast	Ý	SW837	large intestine	Ŷ	SK-MEL-2	skin	Ý
						-		
UACC-893	breast	Y	SW948	large_intestine	Y	SK-MEL-24	skin	Y
8-MG-BA	CNS	Y	T84	large_intestine	Y	SK-MEL-28	skin	Y
A172	CNS	Y	C3A	liver	Y	SK-MEL-3	skin	Y
AM-38	CNS	Y	HLE	liver		SK-MEL-30	skin	Y
7411 30	ens	•	HuH-6-	liver		SIX TILL SU	Skill	•
		Y						
Becker			clone5	liver		SR	skin	Y
	CNS							
CAS-1	CNS	Ý	HuH-7	liver		T107	skin	
CAS-1	CNS			liver	Y	T107		Y
CAS-1 CCF-STTG1	CNS CNS	Y Y	SK-HEP-1	liver liver	Y	T107 UACC-257	skin	Y
CAS-1 CCF-STTG1 D-247-MG	CNS CNS CNS	Y Y Y	SK-HEP-1 SNU-387	liver liver liver	Y Y	T107 UACC-257 UACC-62	skin skin	Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED	CNS CNS CNS CNS	Y Y Y Y	SK-HEP-1 SNU-387 SNU-398	liver liver liver liver		T107 UACC-257 UACC-62 WM-115	skin skin skin	Y Y
CAS-1 CCF-STTG1 D-247-MG	CNS CNS CNS	Y Y Y	SK-HEP-1 SNU-387	liver liver liver		T107 UACC-257 UACC-62	skin skin	Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED	CNS CNS CNS CNS	Y Y Y Y	SK-HEP-1 SNU-387 SNU-398	liver liver liver liver		T107 UACC-257 UACC-62 WM-115	skin skin skin	Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG-	CNS CNS CNS CNS CNS	Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423	liver liver liver liver liver	Y	T107 UACC-257 UACC-62 WM-115 HUTU-80	skin skin skin small_intestine	Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG	CNS CNS CNS CNS CNS CNS	Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449	liver liver liver liver liver liver	Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204	skin skin skin small_intestine soft_tissue	Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG	CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-475	liver liver liver liver liver liver liver	Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673	skin skin skin small_intestine soft_tissue soft_tissue	Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG	CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-449 SNU-475 T24	liver liver liver liver liver liver liver liver	Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401	skin skin small_intestine soft_tissue soft_tissue soft_tissue	Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG	CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-475	liver liver liver liver liver liver liver	Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673	skin skin skin small_intestine soft_tissue soft_tissue	Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG	CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-449 SNU-475 T24	liver liver liver liver liver liver liver liver	Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401	skin skin small_intestine soft_tissue soft_tissue soft_tissue	Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG 05MG GAMG GB-1 GI-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-475 T24 A549 ABC-1	liver liver liver liver liver liver liver lung lung	Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN	liver liver liver liver liver liver liver lung lung lung	Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T	liver liver liver liver liver liver liver lung lung lung lung	Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1	liver liver liver liver liver liver liver lung lung lung lung lung	Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T	liver liver liver liver liver liver liver lung lung lung lung	Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1	liver liver liver liver liver liver liver lung lung lung lung lung	Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-429 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6	liver liver liver liver liver liver liver lung lung lung lung lung lung	Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-3 Calu-6 ChaGo-K-1	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-2 Calu-6 ChaGo-K-1 COLO-668	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1	skin skin skin soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-3 Calu-6 ChaGo-K-1	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-42 KNS-41 KS-1 LN-405 MOG-G-CCM MOG-G-CCM	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COLO-668 COR-L105	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-1MS-1 SK-LMS-1 SK-UT-1	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GGI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-499 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH300 SJRH30	skin skin skin soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-42 KNS-41 KS-1 LN-405 MOG-G-CCM MOG-G-CCM	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COLO-668 COR-L105	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-1MS-1 SK-LMS-1 SK-UT-1	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G- UVW NMC-G1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872	skin skin small_intestine soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COLO-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L47 COR-L88	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH300 SJRH300 SJRH	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GGI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 K	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L28 DMS-114	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-81 LN-405 MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COLO-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L47 COR-L88	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GGI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 K	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-398 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L28 DMS-114	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-UCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 Colu-68 COR-L23 COR-L279 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 SOR-L27 COR-L27 SO	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KNS-81 KS-1 UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L23 COR-L27 COR-L47 COR-L47 COR-L47 COR-L47 COR-L47 COR-L47 DMS-153 DMS-53	liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH300 SJRH300 SJRH	skin skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-425 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L279 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-273 DMS-79	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW	skin skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG OBTRG- 05MG DK-MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L779 COR-L7	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 T17 SW684 SW82 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-425 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L279 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-79	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW	skin skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-UVW NMC-G1 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-429 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 Colu-68 COR-L23 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 COR-L27 DMS-114 DMS-153 DMS-273 DMS-79 DV-90 EBC-1	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SJRH30 SK-LMS-1 SJRH300 SJRH300	skin skin small_intestine soft_tissue	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GGI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-273 DMS-53 DMS-79 DV-90 EBC-1 EKVX	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-42 KNS-81 LN-405 MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL-668 COR-L105 COR-L23 COR-L279 COR-L279 COR-L279 COR-L279 COR-L279 COR-L279 COR-L279 COR-L279 DMS-114 DMS-153 DMS-79 DV-90 EBC-1 EKVX EPLC-272H	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM NOS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 Colu-668 COR-L23 COR-L279 COR-L279 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-2727H HOP-62	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH30	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T88G U-118-MG U251	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L68 DMS-114 DMS-153 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM NOS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 Colu-668 COR-L23 COR-L279 COR-L279 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-2727H HOP-62	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH300 SJRH300 SJRH3	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-42 KNS-81 LN-405 MOG-G-CCM SEC SEC SEC SEC SEC SEC SEC SEC SEC SEC	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L28 DMS-114 DMS-153 DMS-73 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 IST-SL1	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1	skin skin small_intestine soft_tissue stomach stomach stomach stomach stomach	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF239 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L270 COR-L2	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-42 KNS-81 LN-405 MOG-G-CCM SEC SEC SEC SEC SEC SEC SEC SEC SEC SEC	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L47 COR-L68 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1	skin skin small_intestine soft_tissue stomach stomach stomach stomach stomach	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GGI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KNS-42 KNS-81 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM NMC-G1 ND-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYY YYYYY YYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L688 DMS-114 DMS-153 DMS-273 DMS-73	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1 MKN28 MKN45	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYY YYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF239 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L47 COR-L68 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG OBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PF5K SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L279 COR-L270 COR-L	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SJR	skin skin small_intestine soft_tissue soft	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-4275 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L27 COR-L27 COR-L47 COR-L47 COR-L48 DMS-114 DMS-153 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH300 SJRH3	skin skin skin small_intestine soft_tissue	YYY YYYYYY YYYYYYYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOS-76 PFSK SF126 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L47 COR-L47 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad LCLC-103H	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1 MKN28 MKN45 MKN7 NCI-N87 NCI-SNU-1	skin skin skin small_intestine soft_tissue	YYY YYYYYY YYYY YYYY YYYYYYYYY YYY YYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG OBTRG- 05MG DBTRG- 05MG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G- UVW NMC-G1 no-10 no-11 NOS-76 PF5K SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L23 COR-L279 COR-L47 COR-L23 COR-L279 COR-L47 COR-L28 DMS-114 DMS-153 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad LCLC-103H LK-2	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30	skin skin skin small_intestine soft_tissue stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach stomach	YYY YYYYYYY YYYY YYYYYYYYYY YYYY YYYY YYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOS-76 PFSK SF126 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L47 COR-L47 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad LCLC-103H	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1 MKN28 MKN45 MKN7 NCI-N87 NCI-SNU-1	skin skin skin small_intestine soft_tissue	YYY YYYYYY YYYYYYYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM MOG-G-CCM SF126 SF258 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG U-51 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski D0TC2-4510	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L270 COR-L	liver liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 RASS ECC10 ECC12 GC17 GT3TKB KATOIII MKN1 MKN2 MKN7 NCI-N87 NCI-SNU-16 NCI-SNU-15	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYYYYYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DoTc2-4510 HT-3	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L68 DMS-114 DMS-153 DMS-54 DMS-	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 RMS7 A3-KAW AGS ECC10 ECC12 GC17 GT3TKB KATOIII MKN1 MKN28 MKN45 MKN7 NCI-SNU-1 NCI-SNU-1 NCI-SNU-1 NCI-SNU-5 NUGC-3	skin skin skin small_intestine soft_tissue	YYY YYYYY YYYYY YYYYYYY YYYYYYY YYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GB-1 GG-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G- UVW NMC-G1 no-10 no-11 ONS-76 PFSK SF126 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DoTc2-4510 HT-3 ME-180	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-668 COR-L105 COR-L23 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-73 DM	liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SK-UT-1 SW684 SW872 SW982 T-174 TE-159-T TE-441-T VA-ES-BJ 23132-87 A3-KAW AGS ECC10 ECC12 GCIY GT3TKB KATOIII MKN1 MKN28 MKN45 MKN7 NCI-SNU-1 NCI-SNU-16 NCI-SNU-5 NUGC-3 SCH	skin skin skin small_intestine soft_tissue stomach	YYY YYYYYY YYYY YYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-UVW NMC-G1 no-10 no-11 NOS-76 PFSK SF126 SF268 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DoTc2-4510 HT-3 ME-180 OMC-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L27 COR-L27 COR-L279 COR-L270 COR-L27	liver liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYY YYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GAMG GAS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-UVW NMC-G1 no-11 ONS-76 PFSK SF268 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DOTC2-4510 HT-3 ME-180 OMC-1 SiHa	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L23 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad LCLC-103H LK-2 LU-134-A LU-135 LU-135 LU-155 LU-65	liver liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 RMS7 NCI-SNU-1 NCI-SNU-16 NCI-SNU-5 NUGC-3 SCH TGBC11TKB ITO-11	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYY YYYYYYYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GB-1 GI-1 GMS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-CCM MOG-G-UVW NMC-G1 no-10 no-11 NOS-76 PFSK SF126 SF268 SF268 SF539 SK-MG-1 SNB75 SW1783 T986 U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DoTc2-4510 HT-3 ME-180 OMC-1	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYY YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-423 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L27 COR-L27 COR-L279 COR-L270 COR-L27	liver liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 GCT HT-1080 MES-5A RH18 RMS S-117 SJRH30	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYY YYYY YYYYYYYYYYYYYYYYY
CAS-1 CCF-STTG1 D-247-MG D-283-MED D-542-MG DBTRG- 05MG GAMG GAMG GAMG GAS-10 GOS-3 KALS-1 KNS-42 KNS-81 KS-1 LN-405 MOG-G-CCM MOG-G-UVW NMC-G1 no-11 ONS-76 PFSK SF268 SF268 SF539 SK-MG-1 SNB75 SW1783 T98G U-118-MG U251 U-87-MG YH-13 YKG-1 BOKU C-33-A C-4-II Ca-Ski DOTC2-4510 HT-3 ME-180 OMC-1 SiHa	CNS CNS CNS CNS CNS CNS CNS CNS CNS CNS	YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	SK-HEP-1 SNU-387 SNU-387 SNU-423 SNU-423 SNU-423 SNU-449 SNU-475 T24 A549 ABC-1 BEN CAL-12T Calu-1 Calu-3 Calu-6 ChaGo-K-1 COL0-668 COR-L105 COR-L23 COR-L27 COR-L23 COR-L279 COR-L47 COR-L88 DMS-114 DMS-153 DMS-53 DMS-53 DMS-53 DMS-79 DV-90 EBC-1 EKVX EPLC-272H HOP-62 HOP-92 IST-SL1 IST-SL2 KNS-62 LB647- SCLC LC-2-ad LCLC-103H LK-2 LU-134-A LU-135 LU-135 LU-155 LU-65	liver liver liver liver liver liver liver liver lung lung lung lung lung lung lung lung	Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y Y	T107 UACC-257 UACC-62 WM-115 HUTU-80 A204 A673 G-401 G-402 GCT HT-1080 MES-SA RH18 RMS S-117 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 SJRH30 SK-LMS-1 RMS7 NCI-SNU-1 NCI-SNU-16 NCI-SNU-5 NUGC-3 SCH TGBC11TKB ITO-11	skin skin small_intestine soft_tissue soft	YYY YYYYYY YYYY YYYYYYYYYYYYYYYYYYYYYY

			1			D1		
AN3-CA	endometrium	Y	LXF-289	lung		8305C	thyroid	Y
COLO-684	endometrium	Y	NCI-H1048	lung		8505C	thyroid	
EFE-184	endometrium	Y	NCI-H1062	lung		BCPAP	thyroid	Y
ESS-1	endometrium	Y	NCI-H1092	lung		BHT-101	thyroid	Y
HEC-1	endometrium	Y	NCI-H1173	lung		CAL-62	thyroid	Y
KLE	endometrium	Y	NCI-H1184	lung		CGTH-W-1	thyroid	Y
MFE-280	endometrium		NCI-H128	lung		FTC133	thyroid	
MFE-296	endometrium	V	NCI-H1284	lung		K5	thyroid	Y
RL95-2 SNG-M	endometrium endometrium	Y Y	NCI-H1299 NCI-H1304	lung	Y	RO82-W-1 TCO-1	thyroid thyroid	Y
WERI-Rb-1	eye	Y	NCI-H1355	lung lung	I	Π	thyroid	Y
Y79	eye	1	NCI-H1417	lung	Y	22-RVI	unknown	
380	haem_and_lymph		NCI-H146	lung	Ý	CAR-1	unknown	
697	haem_and_lymph	Y	NCI-H1563	lung	Ý	COR-L24	unknown	
ALL-PO	haem_and_lymph	Y	NCI-H157	lung	Y	HTCC3	unknown	
ARH-77	haem_and_lymph	Y	NCI-H1573	lung	Y	IA5	unknown	
ATN-1	haem_and_lymph	Y	NCI-H1581	lung	Y	LC-1-SQ-SF	unknown	
						LCLC-		
BALL-1	haem_and_lymph	Y	NCI-H1618	lung	Y	97TM1	unknown	
BC-1	haem_and_lymph	Y	NCI-H1623	lung	Y	MFN-INO	unknown	
BC-3	haem_and_lymph		NCI-H1648	lung	Y	NB-I-IFO	unknown	
BE-13	haem_and_lymph	Y	NCI-H1650	lung	Y	NCI-H417	unknown	
BL-41	haem_and_lymph	Y	NCI-H1651	lung	Y	SW1315	unknown	
BL-70	haem_and_lymph	Y	NCI-H1666	lung	Y	TCYIK	unknown	
BONNA-12	haem_and_lymph	Y	NCI-H1693	lung	Y	TE-206-T	unknown	Y
DV 172	been and lymph			lung	Y	TuHR-14-	unknown	
BV-173	haem_and_lymph		NCI-H1694	lung	f	ткв	unknown unknown	
C8166	haem_and_lymph	Y	NCI-H1703	lung	Y	NB-12	(auto_ganglia)	Y
00100	naem_anu_iymph	1	NCI-111705	lung	I	ND-12	unknown	'
CESS	haem and lymph	Y	NCI-H1734	lung	Y	NB-SuS-SR	(auto_ganglia)	Y
CLUU	nacm_ana_iymph	•		lang	·	10 505 510	unknown	•
СМК	haem and lymph	Y	NCI-H1755	lung	Y	SK-N-FI	(auto_ganglia)	Y
CML-T1	haem_and_lymph	Ŷ	NCI-H1770	lung	Ý	D-245-MG	unknown (CNS)	Ý
CRO-AP5	haem_and_lymph	Y	NCI-H1792	lung	Y	D-263-MG	unknown (CNS)	Y
CTV-1	haem_and_lymph	Y	NCI-H1793	lung	Y	D-336-MG	unknown (CNS)	Y
Daudi	haem and lymph	Y	NCI-H1838	lung	Y	D-384-MED	unknown (CNS)	Y
DB	haem_and_lymph		NCI-H187	lung	Y	D-392-MG	unknown (CNS)	Y
DEL	haem_and_lymph	Y	NCI-H1882	lung	Y	D-397-MG	unknown (CNS)	Y
DG-75	haem_and_lymph	Y	NCI-H1926	lung	Y	D-423-MG	unknown (CNS)	Y
DOHH-2	haem_and_lymph	Y	NCI-H1930	lung	Y	D-458-MED	unknown (CNS)	Y
EB-2	haem_and_lymph	Y	NCI-H1975	lung	Y	D-502-MG	unknown (CNS)	Y
EB-3	haem_and_lymph	Υ	NCI-H2009	lung	Y	D-538-MG	unknown (CNS)	Y
EHEB	haem_and_lymph	Y	NCI-H2029	lung	Y	D-556-MED	unknown (CNS)	Y
EM-2	haem_and_lymph	Υ	NCI-H2030	lung		D-566-MG	unknown (CNS)	Y
							unknown	
EoL-1-cell	haem_and_lymph	Y	NCI-H2052	lung	Y	ECC4	(gastro_tract)	Y
GA-10-								
Clone-20	haem_and_lymph	Y	NCI-H2073	lung		PLCPRF5	unknown (liver)	Y
GA-10- Clone-4	been and lymph	Y		lung		SW626	unknown (ovory)	Y
CIONE-4	haem_and_lymph	1	NCI-H2085	lung		50020	unknown (ovary) unknown	1
GDM-1	haem and lymph	Y	NCI-H2107	lung	Y	PANC-08-13	(pancreas)	Y
GR-ST	haem_and_lymph	Ý	NCI-H2141	lung		A2058	unknown (skin)	Ý
H9	haem_and_lymph		NCI-H2170	lung		A388	unknown (skin)	Ý
HC-1	haem_and_lymph	Y	NCI-H2195	lung		A431	unknown (skin)	Ý
HDLM-2	haem_and_lymph	Ý	NCI-H2227	lung	Y	GAK	unknown (skin)	Ý
HD-MY-Z	haem_and_lymph	Ŷ	NCI-H2228	lung	Ý	RPMI-7951	unknown (skin)	Ý
HEL-92-1-7	haem and lymph		NCI-H226	lung	Ý	SK-MEL-5	unknown (skin)	Ý
HH	haem_and_lymph	Y	NCI-H2291	lung	Ý	A253	upper_aero_tract	
HL-60	haem_and_lymph	Ŷ	NCI-H23	lung	Ý	BB30-HNC	upper_aero_tract	Y
HT	haem_and_lymph	Ŷ	NCI-H2330	lung	Ý	BB49-HNC	upper_aero_tract	Ý
IM-9	haem and lymph	Y	NCI-H2342	lung		BHY	upper_aero_tract	Y
J45-01	haem_and_lymph		NCI-H2347	lung	Y	Ca9-22	upper_aero_tract	Ý
JiyoyeP-	/ .			5				
2003	haem_and_lymph	Y	NCI-H250	lung	Y	CAL-27	upper_aero_tract	Y
JVM-2	haem_and_lymph	Y	NCI-H292	lung		CAL-33	upper_aero_tract	Y
JVM-3	haem_and_lymph	Y	NCI-H322M	lung	Y	Detroit562	upper_aero_tract	Y
K052	haem_and_lymph	Y	NCI-H345	lung	Y	DOK	upper_aero_tract	Y
K-562	haem_and_lymph	Y	NCI-H358	lung	Y	HCE-T	upper_aero_tract	Y
KARPAS-								
299	haem_and_lymph	Y	NCI-H378	lung		HN	upper_aero_tract	Y
KARPAS-								
422 KARDAS 45	haem_and_lymph	Y	NCI-H441	lung	Y	HO-1-N-1	upper_aero_tract	Y
KARPAS-45	haem_and_lymph	Y	NCI-H446	lung	Y	KOSC-2	upper_aero_tract	Y
KASUMI-1	haem_and_lymph	Y	NCI-H460	lung	Y	LB771-HNC	upper_aero_tract	Y
KE-37	haem_and_lymph	Y	NCI-H510A	lung	Y	RPMI-2650	upper_aero_tract	Y
KM-H2	haem_and_lymph	Y	NCI-H522	lung	Y Y	SAS	upper_aero_tract	Y
KMOE-2	haem_and_lymph	Y Y	NCI-H596	lung	Y Y	SCC-15	upper_aero_tract	Y Y
KMS-12-BM KU812	haem_and_lymph haem_and_lymph	Y Y	NCI-H64	lung	Y Y	SCC-25 SCC-9	upper_aero_tract	Y Y
KU812 KY821	haem_and_lymph	Y	NCI-H650 NCI-H661	lung lung	Y	5637	upper_aero_tract	Y
L-363	haem_and_lymph	Y Y	NCI-H661 NCI-H69		Y Y	639-V	urinary_tract	Y Y
L-540	haem_and_lymph	Y	NCI-H719	lung lung	Y	639-V 647-V	urinary_tract urinary_tract	Y
L-540 LAMA-84	haem_and_lymph	r Y	NCI-H719 NCI-H720	lung	Y	BFTC-905	urinary_tract	Y
LAMA-64 LC4-1	haem_and_lymph	r Y	NCI-H720 NCI-H727	lung	Y	DSH1	urinary_tract	Y
LOUCY	haem_and_lymph	Y	NCI-H748	lung	•	HT-1197	urinary_tract	Ý
LP-1	haem_and_lymph	Ý	NCI-H774	lung	Y	HT-1376	urinary_tract	Ý
MC-1010	haem_and_lymph	Y	NCI-H810	lung	Y	KU-19-19	urinary_tract	'
MEG-01	haem_and_lymph	Ý	NCI-H82	lung	Y	RT-112	urinary_tract	Y
MHH-CALL-	,/F.,						/	·
2	haem_and_lymph	Y	NCI-H835	lung	Y	RT4	urinary_tract	Y
MHH-CALL-								
4	haem_and_lymph		NCI-H889	lung	Y	SW1710	urinary_tract	

MJ ML-2	haem_and_lymph haem_and_lymph	Y Y	PC-14 PC-3 RERF-LC-	lung lung	Y	SW780 UM-UC-3	urinary_tract urinary_tract	Y Y
MN-60 MOLT-13 MOLT-16	haem_and_lymph haem_and_lymph haem_and_lymph	Y Y Y	FM SBC-1 SBC-5	lung lung lung	Y Y Y	SW954 SW962	vulva vulva	Y

Publication

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