# High-throughput retroviral tagging to identify components of specific signaling pathways in cancer

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Published online: 19 August 2002, doi:10.1038/ng950

Genetic screens carried out in lower organisms such as yeast<sup>1</sup>, Drosophila melanogaster<sup>2</sup> and Caenorhabditis elegans<sup>3</sup> have revealed many signaling pathways. For example, components of the RAS signaling cascade were identified using a mutant eye phenotype in *D. melanogaster* as a readout<sup>2</sup>. Screening is usually based on enhancing or suppressing a phenotype by way of a known mutation in a particular signaling pathway. Such in vivo screens have been difficult to carry out in mammals, however, owing to their relatively long generation times and the limited number of animals that can be screened. Here we describe an in vivo mammalian genetic screen used to identify components of pathways contributing to oncogenic transformation. We applied retroviral insertional mutagenesis in Myc transgenic (EµMyc) mice lacking expression of Pim1 and Pim2 to search for genes that can substitute for Pim1 and Pim2 in lymphomagenesis. We determined the chromosomal positions of 477 retroviral insertion sites (RISs) derived from 38 tumors from  $E\mu Myc Pim1^{-/-} Pim2^{-/-}$  mice and 27 tumors from

EµMyc control mice using the Ensembl and Celera annotated mouse genome databases. There were 52 sites occupied by proviruses in more than one tumor. These common insertion sites (CISs) are likely to contain genes contributing to tumorigenesis. Comparison of the RISs in tumors of *Pim*-null mice with the RISs in tumors of EµMyc control mice indicated that 10 of the 52 CISs belong to the *Pim* complementation group. In addition, we found that *Pim3* is selectively activated in *Pim*-null tumor cells, which supports the validity of our approach.

Retroviral insertions in the genome can transform host cells by activating proto-oncogenes or inactivating tumor-suppressor genes<sup>4</sup>. Multiple rounds of retroviral insertional mutagenesis yield a full-blown tumor in which proviral insertions mark the genes collaborating in stepwise tumor development. Thus, retroviral insertions are instrumental in the clonal outgrowth of the incipient tumor cell. In accordance with this notion, two or three CISs within a single tumor are often occupied by proviruses. We have previously shown co-activation of the Pim



Fig. 1 Retroviral tagging in lymphoma-prone E $\mu$ Myc mice that are sensitized to activation of the Pim pathway. a, MoMuLV infection of E $\mu$ Myc mice yields lymphomas, of which 40% and 15% have retrovirus-activated Pim1 and Pim2 alleles, respectively. b, In 90% of the lymphomas generated in E $\mu$ Myc mice lacking expression of Pim1, the Pim pathway has been activated through proviral insertions near Pim2. c, Retroviral insertional mutagenesis in E $\mu$ Myc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup> mice is expected to yield lymphomas with activated oncogenic Pim signaling, either by mutation of a gene in a parallel pathway (Y), a gene downstream of Pim (X) or a Pim-related gene (Z).

family of serine/threonine kinases and either Mvc or Nmvc1 in retrovirus-induced tumors<sup>5,6</sup>. The cooperation between the Myc and Pim proto-oncogenes was proven using transgenic experiments in which EµMycEµPim1 and EµMycEµPim2 double-transgenic mice succumbed around birth to pre-B cell leukemia<sup>7,8</sup>. Although the frequent retroviral activation of Pim1 established the role of the Pim genes in retrovirus-induced lymphomagenesis, the crucial downstream targets of the Pim kinases are elusive. Candidate Pim substrates such as P100 (ref. 9), CDC25A (ref. HP1γ 10).(ref. 11),TFAF2/SNX6 (ref. 12), SOCS1 (ref. 13) and NFATC (ref. 14) have been described, but it is still unclear to what extent thev contribute to Pimmediated transformation.

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Table 1 • Predicted frequencies of random proviral insertions in the mouse genome												
	Expected number of random CISs <sup>a</sup>		Two insertions		Three insertions		F	Four insertions				
Number of tags	Ef <i>r</i> = 0.001	Ef <i>r</i> = 0.005	Ef <i>r</i> = 0.01	Ef <i>r</i> = 0.001	Ef <i>r</i> = 0.005	Ef <i>r</i> = 0.01	Ef <i>r</i> = 0.001	Ef <i>r</i> = 0.005	Ef <i>r</i> = 0.01	Ef <i>r</i> = 0.001	Ef <i>r</i> = 0.005	Ef <i>r</i> = 0.01
10,000	10	50	100	0.26 kb	1.3 kb	2.6 kb	12 kb	27 kb	39 kb	50 kb	88 kb	113 kb
5,000	5	25	50	0.5 kb	2.6 kb	5.2 kb	24 kb	54 kb	77 kb	99 kb	176 kb	227 kb
2,500	2.5	12.5	25	1.04 kb	5.2 kb	10.4 kb	47 kb	108 kb	155 kb	198 kb	351 kb	454 kb
2,000	2	10	20	1.3 kb	6.5 kb	13 kb	59 kb	135 kb	193 kb	248 kb	439 kb	567 kb
1,000	1	5	10	2.6 kb	13 kb	26 kb	118 kb	269 kb	386 kb	495 kb	878 kb	1,134 kb
500	0.5	2.5	5	5.2 kb	26 kb	52 kb	236 kb	538 kb	772 kb	991 kb	1,757 kb	2,267 kb

Ignoring end-of-chromosome effects, random proviral insertions into the mouse genome follow a Poisson distribution. The expected fraction (Efr) indicates the fraction of the total number of proviral insertion sites expected to be random CIS clusters within the specified distance. For example, 2,500 tags will contain 2.5 CISs consisting of 2 random insertions within 1.04 kb, 2.5 CISs of 3 random insertions within 47 kb, and so on. The calculated distances are based on the available mouse genome sequence at Celera (2.6 × 10<sup>6</sup> kb). The expected number of CISs is the mean number of clusters for  $n = \infty$  experiments.

To gain more insight into the oncogenic signaling network in which the Pim proteins act and to identify crucial downstream Pim targets, we established a mammalian *in vivo* enhancer screen similar to the genetic screens carried out in lower organisms. The synergism between *Myc* and *Pim* in lymphomagenesis probably sensitizes those mice that are deficient for Pim but express high levels of Myc to developing lymphomas in which genes acting either downstream of or parallel to *Pim* have been mutated. In fact, the percentage of retroviral activations of *Pim* observed in EµMyc Pim1<sup>-/-</sup> mice was almost twice that observed in EµMyc mice (Fig. 1*a*,*b*). We observed proviral activations of

*Pim2*, which encodes a protein that is 57% identical to Pim1 (ref. 15), in 90% of the lymphomas in *Pim1*-null mice. These observations underscore the selective advantage of *Pim* activation in the presence of high Myc levels and suggest a strategy for identifying genes that rescue loss of Pim function in lymphomas containing activated *Myc*. In the current study we implement this strategy by infecting  $E\mu Myc$  newborns lacking expression of both *Pim1* and *Pim2* with Moloney murine leukemia virus (MoMuLV) (Fig. 1c), carrying out high-throughput sequence analyses of the proviral insertion sites, mapping the insertions and nearby candidate target genes (using the annotated mouse



Fig. 2 Isolation of the genomic DNA sequences flanking the provirus using a splinkerette-based PCR approach. a, Schematic representation of the amplification procedure; provirus (green), splinkerette (gray). Because a random PCR amplification of the sequences flanking the provirus was preferred, genomic tumor DNA was digested with the restriction enzyme BstYl recognizing Pu-GATC-Py, and not with a methylation-sensitive enzyme that selects for proviral insertions in promoter regions of genes<sup>18,30</sup>. **b**, Southern-blot analysis using a viral LTR probe shows the number of proviral insertions in a tumor. The lym-



phoma analyzed carries a large number of retroviral insertions, indicating that this tumor is of oligoclonal origin. **c**, First radioactive splinkerette-based PCR on the same tumor. The asterisk marks the internal MoMuLV fragment amplified. **d**, Nested radioactive splinkerette-based PCR on the excised fragments. **e**, Final PCR amplification of the provirus flanking sequences yields ready-to-sequence DNA fragments.

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genomic sequence database at Celera Genomics and Ensembl<sup>16</sup>) and assigning CISs to complementation groups.

The concept of insertional mutagenesis has so far been based on the assumption that the existence of CISs is due to a selective advantage associated with insertion in that site. Identifying a large number of insertion sites (as in this study) may, however, increase the likelihood of incorrectly labeling a site as a CIS. It is therefore necessary to attach parameters of significance to the definition of a CIS. We propose to assign to each CIS identified an estimate of the non-randomness of its occurrence. In a set of 500 CISs, ran-

dom insertion events may account for approximately 2.5 clusters, of 2 insertions each, within 26 kb (Table 1). Random occurrence of larger CIS clusters (consisting of three or more retroviral insertions) in a data set of 500 is much less likely. It should be noted that the predicted occurrences of CISs do not objectively equal the likelihood of insertions contributing to tumorigenesis. This is because the measure of non-randomness does not take into account preferential insertion due to chromatin structure or sequence context. 'Cold' and 'hot' spots for transposon insertions have been reported for a variety of genomes.

Table 2 • Common retroviral insertion sites in E $\mu$ Myc tumors								
	Candidate		Candidate			No.	No.	
CIS name	gene	Accession ID	protein family	Mouse chr.	Human chr.	isolated tags	insertions	
Dkmi1	Dst	ENSMUSG0000026131	actin cross-linking protein	1	6p11–p12	2	2	
Dkmi2	Ly108	ENSMUSG0000015314	carcinoembryonic antigen	1	1	2	1	
Cis1	Ptma	ENSMUSG0000026238	nuclear protein	1	2q35–q34	1	ND	
Nki3ª	Zfhx1b	ENSMUSG0000026872	zinc-finger homeobox protein	2	2q22	1	ND	
Dkmi3	Ptpn1	ENSMUSG0000027540/	TYR phosphatase	2	20q13.1–13.2	5	ND	
Dkmi4	<u>Set</u> / ND/	NM 023871	nucleosome assembly protein/	2	9q34	3	ND	
	1190004A01Rik	ENSMUSG00000026785 ENSMUSG00000015335	protein kinase/ ND					
Gfi1b	Gfi1b	ENSMUSG0000026815	transcription factor	2	9q34.13	2	2	
Notch1	Notch1	ENSMUSG0000026923	receptor	2	9q34.3	2	ND	
Bmi1	Bmi1	ENSMUSG0000026739	polycomb protein	2	10p13	17	17	
Evi18	RasGrp1	ENSMUSG0000027347	RAS exchange factor	2	15q15	8	3	
Dkmi5	ND	ND	ND	2	20	2	1	
Dkmi6	Pkig	ENSMUSG0000035268	protein kinase inhibitor	2	20q12–q13.1	2	ND	
Dkmi7	Mcl1	ENSMUSG0000038612	BCL-2–related	3	1q21	3	1	
Dkmi8	Cla3	ENSMUSG0000015749	ND	3	1q21.1	2	ND	
Lef1	Lef1	ENSMUSG0000027985	transcription factor	3	4q23–q25	3	2 <sup>c</sup>	
Evi55ª	Camk2d	ENSMUSG0000027970	SER/THR kinase	3	4	2 <sup>b</sup>	ND	
Cis2	ND	ND	ND	4	9	1	ND	
Nki11ª	Runx3	ENSMUSG0000028814	transcription factor	4	1p36	2 <sup>b</sup>	ND	
Evi62ª	E2f2/ Idb3	ENSMUSG0000007872/ ENSMUSG00000018983	HLH factor/ transcription factor	4	1p36	2 <sup>b</sup>	ND	
Evi143ª	<u>Ak4</u> / Lepr	NM 009647/ ENSMUSG0000028529	adenylate kinase/ leptin receptor	4	9p24–p13	1	ND	
Evi58ª	5830400A04Rik	ENSMUSG0000029204	RAS-related	5	4p13	1	ND	
Gfi1/ Evi5	Gfi1/ Evi5	ENSMUSG0000029275/	transcription repressor/	5	1p22	15	15	
		ENSMUSG0000011831	cell-cycle protein					
Dkmi9	Kdr	ENSMUSG0000029232	TYR kinase receptor	5	4q12	3	3	
Kit	Kit	ENSMUSG0000005672	TYR kinase receptor	5	4q12	1	3	
Dkmi10	ND	ENSMUSG0000035273	heparanase	5	4q21.3	3	2 <sup>c</sup>	
Nki16	ND	ENSMUSG0000029471	SER/THR kinase	5	12q24.31	1	ND	
Evi65ª	Coro1c/ Selp1	ENSMUSG0000004530/ NM 009151	actin binding protein/ selectin	5	12q24.1	1	ND	
Evi78ª	Calm2/ ND	NM 007589/ ENSMUSG0000030349	calcium binding protein/ ribosomal protein	6	2p21	1	ND	
Ccnd2	Ccnd2	ENSMUSG0000000184	cell-cycle regulator	6	12p13	3	5	
Cis3	<u>ND</u> /Wnt5b	mCG49753/ ENSMUSG0000030170	F-box protein/ growth factor	6	12p13.3	1	ND	
Evi167ª	Sema4b	ENSMUSG0000030539	receptor	7	15q26.1	2	ND	
Cis4	PD	LOC243990	ND	7	ND	1	ND	
Dkmi11	PD	mCG60113	ND	7	10q25	2	2	
Dkmi12	Rras2/ Copb1	ENSMUSG0000038142/ ENSMUSG00000030754	RAS-related/ beta coat protein	7	11pter–p15	3	3	
Evi83ª	Swap70	ENSMUSG0000031015	coiled-coil BCR binding protein	7	11p15	1	ND	
Dkmi13	Nttp1	ENSMUSG0000037887	TYR/THR phosphatase	7	11p15.5	3	ND	
Dkmi14	PD	mCG57816	Drosophila protein CG5765	8	13q34	2	2	
Evi86ª	Irs2	ENSMUSG0000038894	docking protein	8	13q34	1	ND	
Evi97ª	ORF23 like/ ND	mCG10088/mCG57228	KIAA1865/ ND	8	14q24.3	1	ND	
Dkmi15	ND	ND	ND	8	16p12	3	4	
Evi92ª	Gab1/ Apm1	ENSMUSG0000003033/	clathrin coat protein/	8	4	1	ND	
	·	ENSMUSG0000031714	growth factor receptor associate	ed				
Cis5	1100001J13Rik/ <i>Mshra</i>	ENSMUSG0000001472/ ENSMUSG00000041188	ND/ receptor	8	16q24.3	1	ND	
Cis6	Lyl1	NM008535	transcription factor	8/ <u>10</u>	19p13.2/ 10q22	2 1	ND	

			Table 2 • (continued)				
Fli1	Fli1	ENSMUSG00000016087	transcription factor	9	11q24.1–24.3	1	ND
Ets1	Ets1	ENSMUSG0000032035	transcription factor	9	11q23.3	2	ND
Dkmi16	Madh3	ENSMUSG0000032402	transcription factor	9	15q14–q15	4	ND
Dkmi17	Tcf12	ENSMUSG0000032228	transcription factor	9	15q21	2	2
Cis7	<i>Kif9</i> / PD	ENSMUSG0000032489/ ENSMUSG0000032483	kinesin-related/ KELCH-like	9	3p21	1	ND
Evi100ª	2700018N07	ENSMUSG0000041012	ND	9	ND	1	ND
Cis8	Mknk2	ENSMUSG0000020190	MAPK interacting protein	10	19p13.3	1	ND
Dkmi18 <sup>d</sup>	PD	ENSMUSG0000020258	ND	9/ <u>11</u>	3p21/ 5q33.1	7	ND
Myb	Myb	ENSMUSG0000019982	transcription factor	10	6q23.3–q24	7	ND
Dkmi19	Hbs1l/Myb	NM019702/	elongation factor/	10	6q23–q24	5	10
Nki28ª	PD/ Galgt1	ENSMUSG00000019982 ENSMUSG00000040462/	transcription factor ND/ transferase	10	12q13	1	ND
Tefo2a	Tefo2a		transcription factor	10	10n12 2	ъ	2
Evi150a	Nfic		transcription factor	10	19012.2	2	
EVI 1 30- Evi 1 063	7910012C11Pik			10	19µ15.5 2n16	1	
EVITUO-	2010013GTTRIK		ND transcription factor	11	2010	1	
EVIS	DCITIA		interloukin recentor	11	5025	4 1b	
1191 Evi150a	li9i Supt4b		transcription suppressor	11	5455 17a21 a22	1-	
EVI159-	Suptan Grb7/Znf12		docking protoin/	11	17q21-q25	1	
CISS	GIDTIZIIITA	ENSIVIUSG0000019512/	transcription factor		17421.2	I	ND
Cis10	ND/ Cdc6	ENSMUSC0000018188	WASP interacting/	11	17021.3	2	ND
CISTU	ND/ COCO	ENSMUSG0000038013/	cell-cycle regulator		17421.5	2	ND
Cis11	Stat5a/ Stat5b/ Stat3	ENSMUSG0000004043/ ENSMUSG00000020919/	transcription factors	11	17q11.2	1	ND
Cic12	PD		transcription factor	17	14~24.2	1	
CIST2 Cic12	FD Irf4		transcription factor	12	14424.5	1	
CISTS Evi112a	II 14 Trim25/ Typrd1	AI 022677/	transcription factor/	11/10	0µ25-µ25	1	
LVIIIZ		ENSMUSG0000020250	thioredoxin reductase	<u>11</u> /10	17911.2/12923.3	1	ND
Dkmi20	Crvabp1	ENSMUSG0000021366	transcription factor	13	6p24–p22.3	2	3
Nki33	PD	ENSMUSG0000021755	ND	13	5p13.2	1	ND
Dkmi21	Ptp4a	ENSMUSG0000022606	TYR phosphatase	15	8q24	2	ND
Pim3	Pim3	AF086624	SER/THRkinase	15	22q13.3	5	9
Evi163ª	PD	ENSMUSG0000022462	amino acid transporter	15	12q13.11	1	ND
Dkmi22	Kcnh3/ PD	ENSMUSG00000037579/ ENSMUSG00000037570	potassium channel/ transcription factor	15	12q13.12	2	ND
Cis14	PD	LOC239926	ND	15	ND	1	ND
Dkmi23	PD/ Runx1	mCG60609/	ND/transcription factor	16	21q22	3	4
		ENSMUSG0000022952					
Evi13	Runx1	ENSMUSG0000022952	transcription factor	16	21q22.12	4	3c
Cis15	1810055P05Rik	ENSMUSG0000023883	transcription factor	17	6q27	1	ND
Pim1e	Pim1	ENSMUSG0000024014	SER/THR kinase	17	6p21	9	-
Evi14	Ccnd3l Tbn pending	ENSMUSG0000034165/ ENSMUSG00000023980	cell-cycle regulator/ chromatin associated protein	17	6p21	6	ND
Dkmi24	PD/PD m0	CG55784/ENSMUSG0000004	1683 ND	17	6p21	2	1
Dkmi25	TsgA2	ENSMUSG0000024034	phosphatidyl inositol kinase	17	21q22.3	2	ND
Dkmi26	Fsrg1	ENSMUSG0000024335	bromodomain containing protein	17	4p16.3	2	ND
Tpl2	Tpl2	ENSMUSG0000024235	SER/THR kinase	18	10p11	7	7
Evi136ª	Egr1	ENSMUSG0000038418	transcription factor	18	5q31.1	1	ND
Evi153ª	<u>Hmg1/</u> 1810041M12Rik	mCG9361/ ENSMUSG0000035765	HMG box protein/ ND	18	13q12	1	ND
Dkmi27	Fbxw4	ENSMUSG0000040913	F-box/WD40-repeat	19	10q24–q25	4	1
Evi17ª	Rasgrp2	ENSMUSG0000032946	RAS exchange factor	19	11q13	1	ND
Dkmi28	Vegfb	ENSMUSG0000024962	growth factor	19	11q13	2	4
Dkmi29	Cd6	ENSMUSG0000024670	scavanger receptor	19	11q13	2	1
Pim2 <sup>e</sup>	Pim2	ENSMUSG0000031155	SER/THR kinase	х	Xp11.23	2	-
Nki37ª	Elf4	ENSMUSG0000031103	transcription factor	х	Xq26	1	ND
Dkmi30	1200013B08Rik	ENSMUSG0000031101	TYR kinase	Х	Xq25-26.3	2	_f

Where gene names are specified, RNA/protein expression altered by proviruses has been demonstrated. Candidate genes are genes adjacent to the provirus. Gene accession number at Ensembl (ENSMUSG/LOC), Celera (mCG) or NCBI. The number of insertions is the number of retroviral insertions observed in 38 E $\mu$ Myc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup> tumors as determined by Southern-blot analysis. 'ND' indicates that the number of insertions was not determined for the CIS. CISs in bold have been described previously or the affected genes have been identified by altered mRNA or protein expression. Underlined gene or chromosome names are those used by Celera. 'PD' indicates a predicted gene according to the Ensembl analysis pipeline, and 'PD' indicates a predicted gene according to the Ensembl analysis pipeline, and 'PD' indicates a predicted gene according to the Celera Discovery System. 'Dkmi' indicates a double knockout Myc insertion. <sup>a</sup>RISs overlapping with CISs identified by Suzuki *et al.* (*Evi*) or Lund *et al.* (*Nki*) based on 3 or more insertions within 100 kb. <sup>b</sup>Two independent retroviral insertions (distance > 26 kb). <sup>c</sup>CIS consists of two clusters, of which only one has been checked by Southern-blot analysis. <sup>d</sup>Gene-rich region of 50 kb harboring, according to Celera, four genes encoding the candidate proteins RAN-related (mCG50456), PP2C-like phosphatase (mCG19525), ACTIN depolymerization factor (*Ptk9I*; mG19506), WD-repeat-containing protein (mCG19514). <sup>e</sup>*Pim1* and *Pim2* insertions were isolated only from EµMyc lymphomas. <sup>f</sup>No rearrangments were observed, indicating the subclonal nature of the retroviral analysis.

CIS name	Gene	Protein family	No. insertions <sup>a</sup>	Insertion type	<i>P</i> value <sup>b</sup>
Pim3	Pim3	SER/THR kinase	9	5' promoter, 5' or 3' enhancer	0.000
Kit	Kit	TYR kinase receptor	3	5' enhancer	0.025
Tpl2	Tpl2	SER/THR kinase	7	activating truncation	0.000
Ccnd2	Ccnd2	cell-cycle regulator	5	5' promoter and 5' enhancer	0.002
Dkmi1	ND	actin cross-linking	2		0.088
Dkmi9	ND	TYR kinase receptor	3		0.025
Dkmi11	ND	ND	2		0.088
Dkmi15 <sup>c</sup>	ND	ND	4		0.007
Dkmi20	ND	transcription factor	3		0.025
Dkmi28	ND	growth factor	4		0.007

<sup>a</sup>Number of proviral insertions in 38 Eµ*Myc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup>* tumors detected by Southern-blot analysis. <sup>b</sup>P value of 38 Eµ*Myc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup>* tumors compared with 89 control tumors using Fisher's exact test. <sup>c</sup>CIS consists of two loci 40 kb or 125 kb apart according to Celera and Ensembl, respectively. Protein descriptions in italics represent the proteins encoded by candidate genes 'ND' means that the genes affected by the retroviral insertions have not been determined.

Confirmation of the contribution of CISs to tumorigenesis relies on the identification of the affected gene and evidence that aberrant expression of that gene reproduces specific aspects of the tumor phenotype.

To identify the genomic sequences flanking most proviruses, we designed an efficient, PCR-based splinkerette amplification procedure (Fig. 2). A splinkerette is an adaptor molecule containing a hairpin loop that prevents nonspecific PCR amplification<sup>17</sup>. The applied splinkerette amplification is preferred over the previously described inverse PCR (IPCR) method<sup>18</sup> for two reasons. First, this technique is not based on a long-range PCR amplification, which may increase the size of the amplified fragments and limit the recovery of provirus flanking sequences. Because the complete annotated mouse genomic sequence is available at Ensembl/Celera, the size of the amplified sequences flanking proviruses no longer has an advantage for identifying CISs. Second, this method does not require cloning in bacterial hosts, which increases the speed of the isolation of proviralflanking sequences. We used this splinkerette procedure to analyze the sequences of 477 RISs from 38 lymphomas from  $E_{\mu}MycPim1^{-/-}Pim2^{-/-}$  mice and 27 lymphomas from control  $E\mu Myc$  mice. This group represents approximately 60% of all retroviral insertions present in the tumors, a substantial fraction of which were of oligoclonal origin. The sequence-analyzed fraction of RISs corresponded to an average of approximately seven insertions per tumor (Table 4). We then compared the 477 RIS sequences against the Celera annotated mouse genomic database and found that 176 of the RISs represented 52 CISs (Table 2; for a complete overview, see Web Table A online). Comparison of the RISs with previously identified CISs, and the RISs and CISs characterized by Suzuki et al. (ref. 19; this issue) and Lund et al. (ref. 25; this issue), yielded a total of 91 independent CISs in this tumor panel (Table 2 and Table 4).

the function of Pim1 or Pim2; genes that encode targets or other downstream components of the *Pim1/Pim2* signaling pathway; or genes whose proteins function in pathways parallel to Pim1 or Pim2 that independently activate a similar crucial oncogene target. One common retroviral insertion that was identified in five independent lymphomas from EuMyc Pim1-/- Pim2-/- mice affected Pim3, the third member of the Pim family and thus a prime candidate for the first category of genes that might substitute for Pim1 or Pim2. At the amino-acid level, Pim3 is 71% and 61% identical to Pim1 and Pim2, respectively. Knockout experiments have shown a high degree of redundancy between Pim1 and Pim3, suggesting a similar function for the encoded proteins (H.M., unpublished data). Southern-blot analysis showed insertions near Pim3 in 9 of 38 lymphomas from EµMyc Pim1-/-*Pim2<sup>-/-</sup>* mice. The discovery that *Pim3* is preferentially activated in tumors lacking expression of Pim1 and Pim2 underscores the pathway-specificity of this screen.

To assign a gene to the Pim complementation group (which consists of genes encoding proteins that can either fully or partially substitute for Pim in lymphomagenesis), retroviral insertions near the corresponding genes should be preferentially absent in tumors of mice that express Pim1 and/or Pim2 or, if present, should be mutually exclusive with insertions near one of the Pim genes. To test the validity of this hypothesis, we carried out Southern-blot analysis for insertions near Pim3 in 89 lymphomas from EµMyc, EµMyc Pim1-/- and EµMyc Pim2-/- control mice, of which 61% showed retroviral activation of either Pim1 or Pim2. Retroviral insertions near Pim3 were observed in only one tumor, and this tumor did not carry insertions near Pim1 or Pim2. We subsequently analyzed the whole tumor panel of 89 lymphomas from EµMyc control and 38 EµMyc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup> mice by Southern blotting, using CISs depicted in Table 2 as probes. Nine CISs, identified as Kit, Ccnd2, Tpl2, Dkmi1, Dkmi9, Dkmi11, Dkmi15, Dkmi20 and Dkmi28, were found to be mutually exclusive with Pim1, Pim2 and Pim3 (Table 3). Within this group of

In  $E\mu Myc$  mice lacking expression of *Pim1*, the pressure to activate the Pim pathway by means of proviral insertions in *Pim2* is new high In Fully.

*Pim2* is very high. In Eµ*Myc* mice nullizygous with respect to both *Pim1* and *Pim2*, the selective advantage conferred by retroviral activation of the Pim pathway is likely to remain unchanged. Therefore, genes that can substitute for *Pim1* and *Pim2* in lymphomagenesis can be expected to fall into one of the following categories (Fig. 1*c*): genes encoding proteins that directly substitute for

Table 4 • Cancer loci are efficiently identified by retroviral tagging										
No. tumors	No. tags	No. tags per tumor	No. tags CISs <sup>a</sup>	% tags CISs	CISs per tumor					
65	477	7.40	230	48	3.53					
	known CISs		86	18	1.32					
	new CISs		90	19	1.38					
	RIS/CISs <sup>b</sup>		39	8	0.6					
n	ew CISs (RIS/RIS	)c	15	3	0.23					
PIN	A-substituting C	llSs	25 <sup>d</sup>		0.68 <sup>e</sup>					

<sup>a</sup>Number of proviral tags identifying a CIS. <sup>b</sup>Single RISs from this study belong to CISs identified by Suzuki *et al.*<sup>19</sup> or Lund *et al.*<sup>20</sup>. <sup>C</sup>Comparison of the RISs from this study with the RISs isolated by Suzuki *et al.*<sup>19</sup> and Lund *et al.*<sup>20</sup> revealed additional CISs. <sup>d</sup>Southern-blot analysis showed 34 RISs substituting for Pim. <sup>e</sup>Calculations are based on 38  $E\mu Myc$ *Pim1<sup>-+</sup>Pim2<sup>-+</sup>* tumors. *Pim-*complementing loci, four of the affected genes (*Pim3, Kit, Ccnd2* and *Tpl2*) were identified by altered expression (data not shown). The observation that these loci belong to the Pim complementation group suggests that the proteins encoded by the affected genes act either downstream of or parallel to Pim.

Despite the unknown position of the proteins encoded by the *Pim*-complementing genes relative to Pim signaling, the varying nature of these proteins argues that Pim proteins, like members of the Myc family, have a central role in a complex signaling network. In a pathway model that fits this hypothesis, Pim acts as a modulator of cross-talk between stem-cell factor–induced Kit signaling and cytokine signaling pathways (see Web Fig. A online). To induce a maximum proliferative effect, cytokines require the synergistic action of stem-cell factor<sup>21–23</sup>. Genes induced by interleukins but not stem-cell factor, such as *Pim*, are prime candidates for involvement in the cross-talk mediating the synergistic proliferative effect<sup>8,24</sup>. The modulating role for Pim is supported by the observations that enforced *Pim1* expression reconstitutes the number of lymphocytes in *Rag*-deficient and common- $\gamma$ -deficient mice<sup>25</sup>, and that Pim1 is recruited to the receptor complexes where it associates with the suppressor of cytokine signaling<sup>13</sup>.

The use of genetically modified mice in combination with high-throughput analyses of retroviral insertions and the availability of the complete mouse genomic sequence have permitted us to focus on specific oncogenic signaling pathways by means of an *in vivo* mammalian genetic screen. The strategy described here can indicate whether the protein encoded by a candidate gene belongs to a particular signaling network, and permits a more focused approach in subsequent biochemical analyses. Thus, it represents a mammalian equivalent of the powerful *D. melanogaster* and *C. elegans* genetic screens. In addition, the methodology we used allows combination of data from independent panels, as illustrated by the additional CISs that were identified upon comparison of the RISs from different panels.

#### Methods

**Mice and MoMuLV infection.** The EµMyc mice<sup>26</sup> were bred with  $Pim1^{-/-}$ Pim 1neo59 mice<sup>24</sup> and Pim2<sup>-/-</sup> Pim2 K180 mice (J.A., unpublished data) to generate EµMyc Pim1<sup>-/-</sup>, EµMyc Pim2<sup>-/-</sup> and EµMyc Pim1<sup>-/-</sup> Pim2<sup>-/-</sup> mice. We infected newborns with 1 × 10<sup>5</sup> infectious units of MoMuLV. We killed moribund mice and isolated lymphomas. All animal experiments were approved by the Dutch Animal Research Committee.

**Statistical analysis.** Because exact calculations and simulations show that end-of-chromosome effects can be ignored for a set of several hundred proviral insertions, random insertions in the genome follow a Poisson distribution in which the distance between two adjacent insertions is exponentially distributed. This means that the probability that the distance is at most *x* equals  $1 - e^{-x/\mu}$ , where  $\mu$  equals the mean distance G/(b + 1), where *G* is the sequenced genome size  $(2.6 \times 10^9)$  and *b* is the total number of insertions. For small values of  $x/\mu$  (<0.05), the probability can be approximated by  $x/\mu$ .

The number of insertions to the right of a selected insertion in a fixed window *W* also follows a Poisson distribution. This means that the probability of at least *m* such extra insertions equals  $1 - \exp(-\lambda)(1 + \lambda + \lambda^2/2 + \dots + \lambda^{m-1}/(m-1)!)$ , where  $\lambda$  equals the mean number of insertions in window *W*:  $W \times b/G$  and  $(m-1)! = 1 \times 2 \times 3 \times \dots \times (m-1)$ . For m = 1 (a cluster of two insertions), this equals  $1 - \exp(-\lambda)$ , or approximately  $\lambda$ . For *m* = 2 (a cluster of three insertions), this equals  $1 - \exp(-\lambda)(1 + \lambda)$ , or approximately  $1 - (1 - \lambda + \lambda^2/2) \times (1 + \lambda) = \lambda^2/2$ . For a cluster of 4 insertions, this equals  $1 - \exp(-\lambda)(1 + \lambda + \lambda^2/2) - \lambda^3/6 + \lambda^4/24) \times (1 + \lambda + \lambda^2/2) = \lambda^3/6 - \lambda^4/8$ . These approximations only hold for a small mean number of insertion clusters in the window (<0.05).

**Southern-blot analysis of CISs.** Genomic tumor DNA ( $10 \mu g$ ) was digested with the appropriate restriction enzyme, separated on a 0.7% agarose gel and transferred to Hybond-N membranes (Amersham). We analyzed the number

of proviral insertions and the number of insertions into the known CISs *Pim1*, *Pim2*, *Bmi1* and *Gfi1* using the probes and restriction enzymes as described previously<sup>5,15,28,29</sup>. Genomic fragments, free of repetitive sequences, that flanked the proviruses and hybridized to a CIS were used as probes to analyze the frequency at which a provirus inserted into these loci.

Isolation of the proviral insertion sites. Tumor DNA ( $3 \mu g$ ) was digested with *Bst*YI (New England Biolabs) and the enzyme was subsequently inactivated. We generated the splinkerette adaptor by annealing the splinkerette oligonucleotides HMSpAA and HMSpBB (primer sequences are available on request). Both oligonucleotides contain modifications of a splinkerette described previously<sup>17</sup>. The oligonucleotides (150 pmol each) were denatured at 95 °C for 3 min and subsequently cooled to room temperature at a rate of 1 °C per 15 s using a thermocycler (PTC100, Perkin Elmer). We ligated 600 ng of genomic tumor DNA digested with *Bst*YI to the splinkerette oligonucleotide (molar ratio 1:10) with 4 U T4 DNA ligase (Roche Diagnostics) in a final volume of 40  $\mu$ l. To avoid amplification of the internal 3' MoMuLV fragment, we digested the ligated fragments with 10 U of *Eco*RV in a total volume of 100  $\mu$ l. Ligation mixtures were desalted in a Microcon YM-30 (Amicon BioSeparations).

We amplified MoMuLV-flanking sequences with a radioactive long terminal repeat (LTR)–specific primer, AB949, and a splinkerette primer, HMSp1 (primer sequences are available upon request). Primer AB949 (10 pmol) was radioactively labeled with [ $\gamma^{32}$ P]ATP (3 µCu) using T4 polynucleotide kinase (PNK) (0.2 U; Roche Diagnostics).

The 50 µl PCR mixture contained 150 ng ligated tumor DNA, 10 pmol each primer, 300 nmol dNTPs, 1 U PfuITurbo and 1 × PfuITurbo buffer (Stratagene). The hot-start PCR conditions were 3 min at 94 °C (2 cycles); 15 s at 94 °C, 30 s at 68 °C, 3.5 min at 72 °C (27 cycles); 15 s at 94 °C, 30 s at 66 °C, 3.5 min at 72 °C, and 5 min at 72 °C. We concentrated radioactive PCR fragments using a Microcon-YM30 (Amicon BioSeparations) and then separated them on a 3.5% denaturing polyacrylamide gel. The gels were dried onto 3-mm Wattman paper and exposed overnight to X-Omat AR films (Kodak). We excised amplified fragments from the gel and boiled them for 30 min in 100 µl TE. We used 1 µl of the DNA solution for a nested amplification with a <sup>32</sup>P-labeled virus-specific primer, HM001, and a non-radioactive splinkerette-specific primer, HMSp2 (primer sequences are available upon request). We carried out nested PCR with 5 pmol of each primer, 200 nM of each dNTP, 1.75 mM Mg, 1 U Taq polymerase (Gibco BRL) and  $1 \times PCR$  buffer (Gibco BRL) in a final volume of 20 µl. The PCR conditions were 15 s at 94 °C, 30 s at 60 °C, 3 min at 72 °C for either 25 (for fragments < 400 bp) or 28 cycles (for fragments > 400 bp). We separated the re-amplified fragments on a 3.5% denaturing polyacrylamide gel and isolated them as described above. We then re-amplified 1 µl of the amplified fragments in a non-radioactive PCR of 25 cycles under the conditions as described for the radioactive nested PCR.

We treated the nested PCR mixture with 0.5 U exonuclease and 0.5 U shrimp alkaline phoshatase, according to the manufacturer's (Amersham) instructions. We used about 25 ng of the PCR product in the sequence reaction containing BigDye terminator mix (Perkin Elmer) and primer HM001. In addition, we used HMSp2 as primer for sequencing of amplified fragments larger than 500 bp. We carried out automated sequence analysis on an ABI 377 (Perkin Elmer). The sequences were processed with Sequencher 3.1.1 and subjected to BLAST analysis against the annotated mouse genome databases at Celera (release 1.2) and Ensembl (version 6.3a.1).

**GenBank accession numbers.** The accession numbers for the 477 flanking sequences of the retroviral insertions in the  $E\mu Myc$  tumors (*Dkm*) are AY127080 through AY127557). Further information is available at http://protagdb.nki.nl.

*Note: Supplementary information is available on the Nature Genetics website.* 

#### Acknowledgments

We wish to thank R. Regnerus for assistance in genotyping the mice; N. Bosnie, L. Rijswijk, A. Zwerver, T. Maidment, C. Spaans and F. van der Ahé for animal care; and J. Jonkers and R. van Amerongen for critical reading of the manuscript. This work was supported by the Dutch Cancer Society (H.M.) and the Leukemia Society of America (J.A.).

#### **Competing interests statement**

The authors declare that they have no competing financial interests.

Received 23 April; accepted 8 July 2002.

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## High-throughput retroviral tagging to identify components of specific signaling pathways in cancer

H Mikkers, J Allen, P Knipscheer, L Romeyn, A Hart, E Vink & A Berns

Nature Genet. 32, 153-159 (2002).

By error, several corrections were not made to proofs while preparing the manuscript for the press.

In the reference list, reference 25 (Losman *et al.*) should be inserted as reference 13. As a consequence, references 13–24 should be renumbered as 14–25.

In the text, the following changes should be made:

On page 154, in the second column, reference 16 should be placed at the end of the sentence "These observations underscore the selective advantage..." Reference 16 should also be removed from the first line on page 155.

On page 155, in the second column, reference 17 should be placed at the end of the sentence "Cold' and 'hot' spots for transposon insertions..."

On page 157, in the first full paragraph, reference 17 should be reference 18, reference 18 should be reference 19, reference 19 should be reference 20, and reference 25 should be reference 21.

On page 158, in the first full paragraph, references 21–23 should be references 22,23.

### Genetics, cytokines and human infectious disease: lessons from weakly pathogenic mycobacteria and salmonellae

T H M Ottenhoff, F A W Verreck, E G R Lichtenauer-Kaligis, M A Hoeve, O Sanal & J T van Dissel

Nature Genet. 32, 97-105 (2002).

On page 97, paragraph 2, line 13, 'IL-29' should be 'IL-27'. On page 98, line 11, 'seemed to be' should be 'was', as this has been demonstrated to be the case. On page 98, Fig. 1, a mistake in the color coding occurred, and the affected genes that appear in purple should be colored red. On page 100, line 1, 'IL12Rb1' should read 'IL12Rβ1'.

### Distal ureter morphogenesis depends on epithelial cell remodeling mediated by vitamin A and Ret

E Batourina, C Choi, N Paragas, N Bello, T Hensle, F D Constantini, A Schuchardt, R L Bacallao & C L Mendelsohn

Nature Genet. 32, 109-115 (2002).

doi:10.1038/ng952

The article was missing a reference to Web Movie A, which should have appeared on the last line of page 110 together with the reference to Web Fig. A.

### Targeted mutation of *CyIn2* in the Williams syndrome critical region links CLIP-115 haploinsufficiency to neurodevelopmental abnormalities in mice

C C Hoogenraad, B Koekkoek, A Akhmanova, H Krugers, B Dortland, M Miedema, A van Alphen, W M Kistler, M Jaegle, M Koutsourakis, N Van Camp, M Verhoye, A van der Linden, I Kaverina, F Grosveld, C I De Zeeuw & N Galjart

Nature Genet. 32, 116-127 (2002).

doi:10.1038/ng954

The article mistakenly contained a note stating that supplementary information was available on the *Nature Genetics* website. Instead, it should have contained a brief paragraph in the Methods section listing a separate website where additional information can be found.

### High-throughput retroviral tagging to identify components of specific signaling pathways in cancer

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#### New genes involved in cancer identified by retroviral tagging

T Suzuki, H Shen, K Akagi, H C Morse III, J D Malley, D Q Naiman, N A Jenkins & N G Copeland

Nature Genet. 32, 166–174 (2002).

doi:10.1038/ng949

By error, the subpanel labels for Fig. 1 on page 172 were offset to the right of the corresponding subpanels.