Multiple gene dysfunctions lead to high cancer-susceptibility: evidences from a whole-exome sequencing study

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Abstract— A total of \$275 million has been launched to The Cancer Genome Atlas Project for genomic mapping of more than 20 types of cancers. The major challenge is to develop high throughput and cost-effective techniques for human genome sequencing. We developed a targeted exome sequencing technology to routinely determine human exome sequence. As a proof-of-concept, we chose a unique patient, who underwent three high mortalities cancers, i.e., breast, gallbladder and lung cancers, to reveal the genetic cause of high-cancer-susceptibility. Total 24,545 SNPs were detected. 10,868 (44.27%) SNPs were within coding regions, and 1,077 (4.38%) located in the UTRs. 3367 genes were hit by 4480 non-synonymous mutations in CDS with truncation of 30 proteins; and 10 mutations occurred at the splice sites that would generate different protein isoforms. Substitutions or premature terminations occurred in 132 proteins encoded by cancer-associated genes. CARD8 was completely loss; ANAPC1 was pre-translationally terminated from the transcripts of one allele. On the Ras-MAPK pathway, 18 genes were homozygously mutated. 15 growth factors/cytokines and their receptors, 9 transcription factors, 6 proteins on WNT signaling pathway, and 16 cell surface and extracellular proteins may be dysfunctioned. Exome sequencing made it possible for individualized cancer therapy.

Index Terms— ANAPCI, CARD8, Exome, Ras-MAPK pathw ay, SNP, UTRs, WNT Signaling pathw ay.

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1 Introduction

★t has always been bothering physicians to choose correct drugs as the anticancer effects are completely different among patients. This is caused by not only the multiple genetic mutations in human cancers but also a wide variety of single nucleotide polymorphisms (SNPs) of individuals. Mutations in exons, such as mutations on H-RAS, p53 and APC genes, are often found to cause human cancers. Up to date, 73 genes with germline mutations and 412 genes with germline or somatic alterations, including amplification, deletion, rearrangement and point mutations, have been shown to be involved in human cancers in the Cancer Gene Census of Cancer Genome Project database (CGC/CGP). In the Atlas of Genetics and Cytogenetics Oncology and Haematology (AGCOH) database, there are 766 annotated genes that are genetically associated with cancers and other 3,000 other genes are functionally involved in the process of cancer development. Although a great advance has been achieved for early diagnosis of human cancers and anticancer drug development, the mobility of cancer cases is increasing while the average mortality almost remains consistent in the last decades. The random use of anticancer drugs largely neutralized the attempts of anticancer treatment; and cancer is still the second killer of human diseases. Therefore, it is urgently needed to develop genomebased individualized cancer therapy and care.

It is well known that the whole exome constitute only about 1% of the human genome but harbor the major of mutations contribute to cancer development. Therefore, combined with bioinformatics analysis, targeted exome sequencing technology would be a good and practical strategy to largely reduce the cost and labor load. It would also have a great potential to expand our knowledge of rare mutations in cancer development and to accelerate the functional studies of cancer-associated genes. Using high susceptibility of cancers as proof-

of-concept, we observed that 132 genes, which have been shown to be important for cancer development, dysfunctioned or functionally alternated. Of them, only 11 genes were germline-mutated according to CGC/CGP database; while the mutations of other 121 genes were newly identified in germline in cancer patient.

2 MATERIAL AND METHODS

A very unique cancer patient, a chinese women (YH2), was recruited in this study. She underwent breast cancer, gallbladder adenocacinoma and lung cancer at 41, 63 and 66, respectively. She died of recurrence of gallbladder adenocacinoma in liver at 68. The tumors were removed by surgery at the diagnosis and tumor types were determined by histochemisty assays after surgery. There was no family history of cancers. Informed comment was obtained from the patient for this study, and the study was approved by the ethic committee of The Chinese University of Hong Kong.

Exome sequencing

The strategy for exome sequencing was similar as described by Ng et al. In brief, shotgun libraries were generated from 10 ug of blood leukocytes purified genomic DNA (gDNA) using the standard Illumina protocols. The fragments of size 150-200 bp were isolated after electrophoresis on 6% PAGE and hybridized with NimbleGen 2.1M-probe sequence capture array, in which oligos were fixed to cover the human exomes (RefSeq, NCBI 36.3, 33.92 Mb). The captured exomes were applied for direct single-end sequencing on an Illumina Genome Analyzer II. The average read for each probe is 75 bases. Sequences were then aligned to the reference (RefSeq, hg18,19 and YH1) using SOAP aligner, and the mapped bases, depth, coverage and the base distribution were analyzed.

Substitution detection

SNPs were called by SOAPsnp based on the alignments with HapMap database. For each site within the exome targeted region, only copy number <1.5 of the surrounding area was allowed and the depth should range from 10X to 200X. Finally, a Q20 threshold was used to filter unreliable SNPs. After excluding known substitutions from the potential mutations available, the SNPs were annotated and the genes involved in cancer development were revealed by comparison of our data with CGC/CGP and the AGCOH database.

Insertion and deletion detection

For the single reads we produced, the short in-dels <4 bp were also identified by S0APaligner2 in a gap tolerable mode. Local alignments were performed with our custom perl scripts.

3 RESULTS

Exome sequences

Our sequencing strategy was similar to the one published by Ng et al recently but with a larger coverage (33.92 instead of 26.6 megabases). The average sequencing depth was 21.1 (Figure 1). The total reads were about 1.97 Gagabases (GBs) which covered 97.36% of the reference. With SOAPaligner software, 87.92% of bases were aligned to the reference (build 131,10/03/26, hg18 and hg19) and YH genome sequence. The mismatch rate was 0.65%, indicating the data was in high sequencing quality. We detected total 24,545 SNPs. Among them, 10,874 (44.3%) SNPs located in the coding regions and 142 (0.6%) SNPs located in the UTRs. There were 23,604 SNPs were shared among YH1 and dbSNPs, while 941 SNPs were newly identified in the patient after comparative analysis of SNPs in the captured exome. Among them, 8091 SNPs (42.81%) were homozygous. 3058 genes were hit by 4480 nonsynonymous mutations in the coding sequences (CDS). 10 mutations displayed at spice sites, and 8 small in/dels were identified.

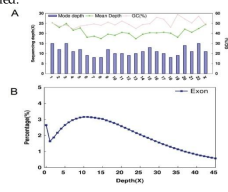


Fig: 1 Targeted capture exome sequencing. A. Chromosome depth and GC distribution in targeted capture exome regions. X axis stands for each chromosome, Y1 axis presents the sequencing depth and YH2 axis is the GC proportion in exon capture region of each chromosome.

Nonsense mutations

We detected 33 nonsense mutations that caused truncation of 30 proteins (Table 1). We found only 3 proteins (PTPN11,

MAGEE2 and IL17RB) have been recorded to have genetic associations with cancer; while 11 other cancer-associated proteins, for the first time, were observed to be mutated in the germline. Particularly, MAGEE2, which has been shown genetic association in melanoma and hepato-cellular carcinoma, was truncated at N-terminal by homozygous mutations. CARD8, a key factor for the recruitment of caspase in apoptosis pathway, was almost completely loss in the patient. ANAPC1, a key components of ana-phase promoting complex that play crucial roles in cell mitosis and protection of the integration of chromosomes from separation, truncated >70% by a heterozygous mutation at Gln465. Some important proteins on the RAS-MAPK signaling pathway, including G protein coupled receptor 1 (GRP1), tyrosine kinase (MAP2K3), and protein tyrosine phosphatase (PTPN11), also prematurely terminated.

TABLE 1: NONSENSE MUTATIONS (ST, STOP; JMML, JUVENILE MYE-LOMONOCYTIC LEUKEMIA; AML, ACUTE MYELOGENOUS LEUKEMIA; MDS, MYELODYSPLASTIC SYNDROME)

Name	T	Muta-	Position	full	Function	Genetic
	Y	tion	(stop)	length		association
	P			(aa)		with dis-
	e					ease(s)
Functional	associ	ated with ca	ancers			
ANAPC1	Н	CAG>TA	AG Q465	1926	anaphase	
	E				promoting	
	T				complex	
GPR1	Н	CGA>TC	GA R236	355	signal trans-	
	E				duction	
	T					
ASCC3	Н	CAG>TA	AG Q87	111	signal trans-	
	E				duction	
	T					
MAP2K3	Н	CAG>TA	AG Q73	318	tyrosine	
	Е				kinase, signal	
	T				transduction	
PTPN11	H	TAT>TA	.G Y197	593	protein tyro-	JMML,
	E				sine phospha-	AML,
MAGEE2	T	G L G . TI	1.C F120	500	tases	MDS
MAGEEZ	H O	GAG>TA	AG E120	523	signal trans- duction	melano ma,
					duction	HCC
CARD8	M H	TGT>TG	A C10	432	caspase	rheumatoid
CARDO	0	101>10	IA CIU	432	recruitment	arthritis
	M				recruitment	artiffus
AB CA10	Н	CGA>TC	GA R132	2 1544	drug transport	
преше	E	convic	J11 K132	2 1544	arag transport	
	Т					
CYP2C18	Н	TAT>TA	A Y68	490	drug metabol-	
	E				ism	
	Т					
IL17RB	Н	CAG>TA	AG Q484	502	cytokine	intestinal
	Е				receptor	inflamma-
	Т					tion

UBE2NL	Н	TTA>TGA	L89	153	ubiquitin
	E				ligation
	T				
FTHL17	Н	GAG>TAG	E148	183	ferritin heavy
	Е				polypeptide-
	T				like protein
TP53RK	Н	CGA>TGA	R152	254	TP53-
1133KK	E	CGA>IGA	K132	234	
	T				regulating kinase
	1				Killase
041					
Others					
SPATA21	Н	CGA>TGA	R467	470	spermatoge-
	E				nesis
	T				
PZP	Н	CAA>TAA	Q598	1483	proteinase
	Е				inhibitor
	T				
UNC5CL	Н	CAG>TAA	Q12	519	NF-kB inhibi-
	E				tor
	T				
TCTE1	Н	CAG>TAG	Q460	502	t-complex-
	E				associated-
	T				testis-
					expressed 1
ASCC3	Н	CAG>TAG	Q87	2203	RNA helicase
	E				
	T				
ZN F75D	Н	CGA>TGA	R331	511	transcriptional
	Е				factor
	T				
DKFZp54	Н	TGG>TGA	W141	150	unknown
7	О				
	M				
LOC1496	Н	CGA>TGA	R37	98	unknown
43	Е				
	T				
MS4A12	Н	CAA>TAA	Q71	267	membrane
	Е				protein
	T				
OR2T5	Н	CGA>TGA	R24	315	olfactory
	Е				receptor
	T				
PZP	Н	CAA>TAA	Q598	1483	pregnancy-
	E				zone protein
	T				Protom
SLC6A18	Н	TAC>TAG	Y319	628	unknown
	Е			220	
	T				
SPATA21	H	CGA>TGA	R467	470	spermatoge-
SIN INZI	н Е	CGA/IGA	N+0/	770	nesis
	E T				110518
ZN F75	Н	CGA>TGA	R331	510	zinc finger
24173		COA>IOA	K331	310	S .
	Е				protein
ZNIEGO	T	TAT TAG	V245	272	ning Carry
ZN F80	Н	TAT>TAG	Y245	273	zinc finger
	E				protein
	T				

Missense mutations

Missensense mutations hit over 3,000 proteins. After aligned with the CGC/CGP and AGCOH databases, we observed important substitutions (most likely causing function alterations) occurred in 132 proteins, which strongly associated with cancer development (Table 3). Among them, 45 have been recorded as somatic mutations and only 11 recorded as germline mutations in cancer patients in the CGC/CGP database. Totally 121 cancer-associated genes were newly found to display mutations in germline; some mutations would cause significant function alterations.

TABLE 2: HOMOZYGOUS MUTATION(S) IN GENES STRONGLY (EITHER GENETICALLY OR FUNCTIONALLY) ASSOCIATED WITH CARCINOGENESIS (*, HETEROZYGOUS MUTATION)

		-			
Name	FL (aa)	mutations	Name	FL (aa)	Mutations
RAS-MAPI	ζ.	_	Wnt signal	ing pathwa	y
signaling pa	thway				
EML4	981	K283E	APC	2843	V1822D
ENPP2	865	S493P	CD97	786	R318Q
EPHA1	976	M900V	DKK2	259	R146Q
FNIP1	1166	G76C, Q648R	DKK3	350	R335G
GPR103	431	L344S		actors/cytol transducer	kines and their recep-
GPR112	3080	T1213N.S1 540P, F1791L, I276M*, P368H*	FGFR4	802	V10I, P136L
GPR116	1346	T604M	IGF2R	2491	R1619G, N2020
GPR142	462	H132N	IL23R	629	Q3H, L310P
GPRC6A	926	P91S	MST1R	1400	Q523R(E), S1195G, R1335G(E)
GRP115	695	K541N	PPARGC 1A	798	G482S
GRP56	693	S281R	TNC	2201	V295M*, Q539R, V605I, E2008Q*
KLK4	251	S22A*, H197Q	TNFRSF 10A	468	H141R, R209T, R441K
KLK5	293	N153D	TNFRSF 17	184	N81S
KLK10	276	S50A, L149P*	TRAF3	568	M129T
KLK11	250	G17E	PLEK2	354	S217C
NIN	2046	Q1125P, G1320E	Cell cycle	control	
RHOD	210	C134R	ATM	3056	N1983S
TEK	1124	I148T*, Q346P	BUB1B	1050	R349Q
Apoptosis/a	nti-apoptos	-			Others
CARD8	432	C10ST	ASXL1	1541	L815P
BCL2L2	194	Q133R	CDH11	796	T255M*,
		-			M275I*,S373A
OPTN	577	M98K*,	BRIP1	1249	S919P

		K322E			
DNA synthesis	repair/RNA		COL1A1	1464	T1075A
ERCC5	1186	G1053R, G1080R, D1104H	GOLGA5	731	A67G*,P350L
FANCA	1454	T266A,A41 2V*,G501S ,P643A*,G 809D, T1328A*	LCP1	627	K533E
DDX43	648	K625E, Q629R	LIFR	1098	D578N
ATM	3056	N1983S	MA- GEE2	523	E120ST(GAG>TA G)
BUB1B	1049	R349Q	MEN1	615	T546A
Transcript	ion factors		NUT	1132	P22L
AFF3	1226	S538N	PDE4DI P	2346	R25L*, A167T*, R681H*, C708R, R1504Q*
CDX2	313	P293S	PMS2	862	P470S*, T485K*,
GATA2	480	A146T	P0U6F2	691	P191L

Homozygous mutations displayed in 58 genes that may contribute to high susceptibility of cancers in this patient. Homozygous missense mutations occurred in 18 genes on RAS-MARK pathway, including G-protein coupled receptors (GPRs), tyrosine kinases and phosphatases (Table 2). On this pathway, heterozygous mutations hit 9 other genes, including AKAP12, CBLB, MAP2K3, MAP3K7IP1, PTPN11, PTPN21, TCL1B and USP6 (Table 3). Although the proteins encoded by these genes play critical roles in cells response to extracellular signalings; however, only EMIA and NIN were recorded somatic mutations in tumors in the CGC/CGP database. The second largest group (10 genes), which were hit by homozygous mutations, were growth factors/cytokines and their receptors. Although only mutation of TNFRSF17 was shown in the intestinal T-cell lymphoma in the database, the products of these genes are important to control cell growth and immune responses to infection and other human diseases including carcinogenesis. On the Wnt signaling pathway, besides APC, homozygous mutations of CD97, DKK2 and DKK3 most likely cause significant alteration of protein functions. The genetic alterations in tumors have not yet recorded. Apart from DDX43, the other homozy-gously mutated genes (ATM, BUB1B, ERCC5 and FANCA) for cell cycle control and DNA/RNA process were shown genetic association with cacinogenesis (Table 2). Besides function association, the germline mutations of transcription factors (AFF3 and POU6F2) have not yet recorded. All 3 apoptotic/anti-apoptotic genes (CARD8, BCL2L2 and OPTN) were newly observed genetic alterations in cancer patients. This would enhance the somatic cells escaping from apoptosis during carcinogenesis.

TABLE 3: MUTATIONS IN THE GENES STRONGLY ASSOCIATED WITH HUMAN CANCERS

Somatic

Muta

FL (aa)1

Gene

	y		tion		line
	p				
	e				
ACSL3	Н	719	L641H	prostic cancer	
	e			•	
	t				
ADAM12	Н	1593	G48R		
	0				
	m				
ADAM8	Н	823	W35R,		
	О		F657L		
	m				
ADAMST5	Н	929	R614H,L		
	e		692P		
	t				
ADAMTS4	Н	1226	S538N		
	0				
	m				
AKAP12	Н	324	K118Q,K	multiple cancers,	
	e		1218I	anti-angiogenesis	
	t				
AKR1C4	Н	324	S145C*,		
	О		Q250R,		
	m		L311V*		
ALOX12	Н	662	N322S		
	0				
	m				
ANAPC1	Н	1926	Q465ST(
	e		GAC-		
	t		>TAC)		
APC	Н	2843	V1288D	colorectal, pan-	the same
	0			creatic, desmoid,	cancers as
	m			hepatoblastoma,	somatic
				glioma, other	mutations
				CNS cancers	
ASNS	Н	561	V210E		
	e				
	t				
ASXL1	Н	1541	L815P	MDS, CMML	
	О				
	m				
ATF6	A	670	A145P,		leukemia,
	T		P157S		lymphoma,
	F				medullob-
	6				lastoma,
					gliom
ATM	Н	3056	N1983S	T-PLL	
	0				
	m				
BCAS1	Н	584	Q24K,		
	О		V163A*		
	m				
BCL2A1	Н	174	C19Y,		
	e		N39K,		
	t		G82D		
BCL9	Н	1426	A218V	B-ALL, Hodgkin	co-
	e			lymphoma	lon/breast/o

Germ

Math												
DMPRIA		t				vary cancer,	DKK3	Н	349	R335G	gas-	
Martial Mart								О				
BMPRIA								m				
Mart						myosarca-	EML4	Н	980	K283E	NSCLC	
BRIP 1						ma		О				
BRIEL	BMPR1A	Н	531	P2T	breast cancer	AML,		m				
BRIPI		e										
BRIPI		t					ENPP2		865	S493E		
BUBIB H 1049 R349Q colorestal cancer gastrointes m colorestal cancer gastrointes m colorestal cancer c	DDID1		1240	CO 1 OD		cancer						
BUBIB H 149 R349Q colorectal cancer, gastriotte- Fact Fact	DKIPI		1249	3919P			ЕРНА 1		976	V160A		
BUBIS							Elimi		710	*10071		
Part	BUB1B		1049	R349O	colorectal cancer.	gastrointes-						
Part						-	ERCC2		759	K751N		skin basal
CARCI		m				lasia,		e				cell, mela-
CARCI						rhabdo-		t				noma,
CARDE						myosarco-						SKC,
Part						ma	ERCC5	Н	1186	G1053R,		skin basal
CARDS	CABC1	Н	647	H85Q				О				
CARDS												melano ma
CARS							FGFR2		820	M186T	_	
CARS	CARD8		432									
CARS							EGER4		802	VIOI	NSCLC	
CBLB	CARS		879		AI CL		TOTKT		002	V101		
CDLB	CARS		017	A//+1	ALCL							
CCND3							FLT3		992	T227M,	AML, ALL	
CCND3	CBLB	Н	Het	N466D	AML			e		D358V		
CCND3		e						t				
CD97		t					FNIP1	Н	1165	G76C,		
CD97	CCND3	Н	292	S259A	MM			О		Q648R		
CD97												
CDH11							FTHL17		183			
The content of the	CD97		785	R318Q								
CDH11							FXVD5		178			
CDX2	CDH11		795	T255M*.	aneurismal bone		THIDS		170			
March Marc	021111		,,,,							1117011		
CENPF					•		GATA2		479	A146T	AML	
CENPF	CDX2	Н	313	P293S	AML			0				
CENPF		0						m				
O R2943G, m t GOLGAS H 730 A67G*,P papillary thyroid COL1A1 H 1465 T1075A o 350L O m m GPR1 H 355 R236st COL1A2 H 1365 P549A dermato fibrosar-coma protube-rans e (CGA-coma protube-rans DDX43 H 647 K625E o m DKK2 H 259 R146Q o P368H*,T O P368H*,T m 1213N,		m					GGH	Н	317	C6R		
Mathematical Reservoir Mathematical Reserv	CENPF	Н	3113									
COL1A1							901.5		5 20			
COL1A2							GOLGA5		730		papillary thyroid	
COL1A2	COLIAI		1465	T1075A						350L		
COL1A2							GDD1		355	P236ct		
o coma protube- m t >TGA) DDX43 H 647 K625E o o m GPR103 H 431 L344S o m GPR112 H 3080 276M*, DKK2 H 259 R146Q o P368H*,T o 1213N, m 1213N,	COL1A2		1365	P549A	dermato fibrosar-		O. Ki		553			
m rans GPR103 H 431 L344S DDX43 H 647 K625E o o m m GPR112 H 3080 276M*, DKK2 H 259 R146Q o P368H*,T o 1213N,	0021112		1000	10.011								
DDX43					-		GPR103		431			
m GPR112 H 3080 276M*, DKK2 H 259 R146Q o P368H*,T o m 1213N,	DDX43	Н	647	K625E				o				
DKK2 H 259 R146Q o P368H*,T o m 1213N,		o						m				
o m 1213N,		m					GPR112	Н	3080	276M*,		
	DKK2	Н	259	R146Q				0				
m S1540P,								m				
		m								S1540P,		

			F1791L			MGC34647	Н	266	Y213st	
GPR116	Н	1345	T604M				e		(TAC-	
	О						t		>TAG)	
	m					MMP10	Н	475	D81Y	
GPR142	Н	462	H132N				e			
	О						t			
	m					MMP11	Н	486	A38V	
GPRC6A	Н	925	P91S				О			
	О						m			
	m					MMP17	Н	602	A182T	
GRP 1 15	Н	694	K541N				o			
	О						m			
	m									
GRP56	Н	692	S281R							
	О					MMP20	Н	482	K18T*,	
	m						О		V275A,T	
HTATIP2	Н	276	S231R				m		281N	
	О					MMP26		260	K43E	
	m						О			
IGF2R	Н	628	Q3H,				m			
	О		L310P			MMP27	Н	512	M30V	
	m						0			
JAG2	Н	1237	E501K				m			
	e					MMP8		467	K87E	
	t						0			
KLK10		275	S50A,				m			
	0		L149P*			MMP9		706	Q279R	
	m						e			
KLK4		250	S22A*,			3.50m4	t	50 4	P4000	
	0		H179Q			MST1	Н	724	R108Q,	breast cancer
	m									
*** ***		202	374 # OD				e		R122Q	
KLK5	Н	292	N153D			MOTER	t	1200		
KLK5	H o	292	N153D			MST1R	t H	1399	Q523R/E,	
	H o m			MIII		MSTIR	t H o	1399	Q523R/E, S1195G,	
KLK5	H o m H	292	N153D K553E	NHL		MSTIR	t H	1399	Q523R/E, S1195G, R1135G/	
	H o m H			NHL			t H o m		Q523R/E, S1195G, R1135G/ E	
LCP1	H o m H o	626	K553E		adana	MSTIR MTHFR	t H o m	1399	Q523R/E, S1195G, R1135G/	
	H o m H o m			salivary	adeno-		t H o m		Q523R/E, S1195G, R1135G/ E	
LCP1	H o m H o m H o	626	K553E		adeno-	MTHFR	t H o m H e t	655	Q523R/E, S1195G, R1135G/ E A222V	
LCP1	H o m H o m H o m	626 1097	K553E D578N	salivary	adeno-		t H o m H e t		Q523R/E, S1195G, R1135G/ E A222V	
LCP1	H o m H o m H o m H o	626	K553E	salivary	adeno-	MTHFR	t H o m H e t	655	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q,	
LCP1	H o m H o m H o m H o	626 1097	K553E D578N	salivary	adeno-	MTHFR	t H o m H e t H	655	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q, G271R	AML
LCP1 LIFR LOX	H o m H o m H o m H o m	626 1097 417	K553E D578N R158Q	salivary	adeno-	MTHFR	t H o m H e t H e	655	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q,	AML
LCP1	H o m H o m H o m H o m H o m H o H	626 1097	K553E D578N	salivary	adeno-	MTHFR	t H o m H e t H e	655	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q, G271R	AML
LCP1 LIFR LOX	H o m H o m H o m H o m H o m H o m	626 1097 417	K553E D578N R158Q	salivary	adeno-	MYEOV MYH11	t H o m H e t H e t	655 312 1937	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q, G271R	AML
LCP1 LIFR LOX LOXL2	H o m H o m H o m H o m H o m H o m m	626 1097 417 773	K553E D578N R158Q M570L	salivary	adeno-	MTHFR	t H o m H e t H e t H H	655	Q523R/E, S1195G, R1135G/ E A222V V159A, R198Q, G271R N1899S	AML
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NUT	Н	1131	P22L	lethal midline		ROS1	Н	2347	T145P	
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OPTN	Н	576	M98K*,			SDC1	Н	310	L136Q	
	0		K322E			~~~	0			
			KJ22L							
DARAM	m	#0.4	****				m			
P2RX7		594	Y155H*,							
	О		R270H*,							
	m		E496A*,			SELE	Н	371	S303R	
			N568I				e			
PBX1	Н	429	G21S	Pre B-ALL			t			
	e					SERPINB5	Н	374	S176P,	
	t						e		I319V	
PDE4D I P		2345	R25L*,	MPD			t			
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						SPRI 4		343		
	m		R681H*,				e		R340K	
			C708R,				t			
			R1504Q*			STEAP2	Н	489	F17C*,	
PDGFRA	Н		S361R,	GIST, idiopathic			O		R456Q*,	
	e		T474M,S	hyperosinophilic			m		M475I	
	t		478P	syndrome		TCF3	Н	653	P479L	pre B-ALL
PLAG1	Н	500	S443R	salivary adeno-			e			
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PMS2	Н	861	P470S*,		colorectal,		e			hood) epithelio-
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	m					TNC	Н	2200	V295M*,	glioma,
PTPN11	Н	592	S189A,	JMML, AML,			О		Q539R,	lung/colon/breas
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PTPN21	п	1173	L385F,			1111 101 10A		707	R209T,	
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	e		V936A				m		R441K	
	t					TNFRSF17	Н	183	N81S	intestinal T-cell
PVRL4	Н	509	F53L				О			lymphoma
	e						m			
						·				

TRAF3	Н	568	M129T		
	О				
	m				
TSC1	Н	365	M322T		
	e				
	t				
USP6	Н	234	Y162H,	aneurysmal bone	
	e		W475R,	cysts	
	t		Y484H		
WISP3	Н	331	Q34H,	colon cancer	hamarto ma,
	e		E100K,		renal cancer
	t		E141K		

ALCL, anaplastic large-cell lymphoma; ALL, acute lymphocytic leukemia; AML, acute myelogenous leukemia; APL, acute promyelocytic leukemia; B-ALI, B-cell acute lymphocytic leukaemia; CMML, chronic myelomonocytic leukemia; CNS, central nervous system; DLBL, diffuse large B-cell lymphoma; DLCL, diffuse large-cell lymphoma; GIST, gastrointestinal stromal tumour, JMML, juvenile myelomonocytic leukemia; MDS, myelodysplastic syndrome, MLCLS, mediastinal large cell lymphoma with sclerosis; MM, multiple myeloma; MPD, Myeloproliferative disorder; NHL, non-Hodgkin lymphoma; NSCLC, non small cell lung cancer, pre-B All, pre-B-cell acute lymphoblastic leukaemia; SKC, skin squamous cell; T-PLL, T cell prolymphocytic leukaemia. *listed as heterozyous mutation.

4 DISCUSSION

The Cancer Genome Atlas project is currently the central task of genome-related research. It remains largely unknown how germline mutations in global contribute to cancersusceptibility, although it is well known some germline mutations in a special gene would cause human cancers (e.g., mutaions in pRB gene leads to retinoblastoma in children). The major challenge is to develop a high throughput and costeffective techniques for genome sequencing. Supported with extensive bioinformatic assays, a US group and us have independently developed cost-effective targeted capture exome sequencing technology to routinely reveal the genetic variations of individuals. However, to our knowledge, the whole exome sequencing on high-cancer-susceptible patient has not yet been studied. In this study, we independently developed a similar technology for the whole exome sequencing. As a pilot study, we showed that homozygous mutations of CARD8 may contribute to the high-cancer-susceptibility in a patient, who underwent three high mortality cancers (breast cancer, gallbladder cancer and lung cancer) in the last three decades. CARD8 was reported to inhibit apoptosis and caspase activation induced by Apaf-1/caspase-9-dependent stimili; however, it was also showed to induce apoptosis in certain cells. It is unclear how the loss of CARD8 contributes the high-cancersusceptibility in this patient. The mutations in other genes, such as genes on RAS-MARK signaling pathway, may also play important roles in high-cancer-susceptibility. However, as some mutations may neutralize or antagonize the other mutations, the exact roles of these mutations are very complicated in the patient. For example, the truncation of MAGEE2 and PTPN11 may neutralize the mutations of tyrosine kinases and GPRs. The roles of these mutations in cancer-susceptibility would be further investigated by identification of more highcancer-susceptibility patients or direct sequencing the tumor samples and paired germline genomes.

In summary, we developed targeted exome capture sequencing technology to characterize the whole-exome of human genome and applied to a high-cancer-susceptible patient. We showed that the truncations of CARD8, MAGEE2, ANAPC1, GPR1, ASCC3, MAP2K3 and PTPN11 be an important reasons for high-cancer-susceptiblity. The non-synonymous mutations in 132 cancer-associated genes, in which most of them have not been reported as germline variations in tumors, may positively or negatively contribute to cancer development. This exome sequencing technology makes it possible for routine dissection of important genes for carcinogenesis and individualized medicine, as the total cost is just less than US\$10,000 per sample. The targeted exome capture sequencing would be a new era of individualized cancer therapy.

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6 DECLARATION

No conflicts of interest.

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