

# Differences in clinical characteristics and mutational pattern between synchronous and metachronous colorectal liver metastases

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**Purpose:** To investigate differences in clinical characteristics and mutational patterns between synchronous and metachronous colorectal liver metastases (CLMs).

**Patients and methods:** From June 2008 to December 2014, patients with RAS wild-type CLMs treated at Zhongshan Hospital, Fudan University were included. DNA extracted from formalin-fixed paraffin-embedded tissue of primary tumors was sequenced with next-generation sequencing for single-nucleotide polymorphism of 96 genes according to custom panel. Mutations were compared between synchronous and metachronous liver metastases and correlated with clinical characteristics.

**Results:** A total of 161 patients were included: 93 patients with synchronous CLM and 68 patients with metachronous CLM. Patients with metachronous CLM were obviously elder. For pathology of primary tumors, synchronous CLMs were larger in size, poorly differentiated, and more frequently local advanced and lymph node positive. For evaluation of liver metastases, synchronous CLM had more and larger metastatic lesions. The median number of mutations in synchronous CLMs was significantly higher than in metachronous group (22 vs. 18,  $p < 0.001$ ). EGFR rs2227983 is the most prevalent mutation in both groups and only a part of prevalent mutations is shared in both groups. Prevalent mutations were correlated with many clinical characteristics. EGFR rs2227983, RBMXL3 rs12399211, and PTCH1 rs357564 were prognostic for latency of metachronous CLM.

**Conclusion:** Clinically, synchronous CLMs, compared with metachronous CLMs, were younger and showed heavier tumor burden for both primaries and liver metastases. Genetically, we identified different mutational patterns between synchronous and metachronous CLMs and several correlations between mutations and clinical characteristics. Further researches were needed to confirm these potential key mutations of CLMs.

**Keywords:** colorectal liver metastases, next-generation sequencing, metachronous, synchronous

## Introduction

Colorectal cancer (CRC) is the most common malignancy throughout the world,<sup>1</sup> and liver metastases are the major cause of death in CRC patients. Unfortunately, 40%–50% of CRC patients would develop colorectal liver metastases (CLMs). They are either synchronous or metachronous in presentation with approximately equal incidence.<sup>2,3</sup> Compared with synchronous metastases, patients with metachronous metastases show differences in terms of clinicopathologic characteristics and better prognosis after metastases resections.<sup>4–6</sup> However, in previous studies, synchronous and metachronous liver metastases were usually lumped together, neglecting their clinical and biologic

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differences. It is critical to improve our understanding of the biology of liver metastases, which may help to develop more effective therapeutic strategies.

Biologic differences between synchronous and metachronous CLMs had been studied previously but showed conflicting results.<sup>2,7-11</sup> Available evidence indicated that majority of gene alterations in the primary tumor were maintained in the CLMs, whereas a limited number of studies compared the primary tumors of synchronous and metachronous groups and generally demonstrated no differences in biomarker expression. In consideration of limited number of studies and limited number of biomarkers investigated, it is still not reasonable to deny the hypothesis that the biology of synchronous and metachronous CLMs would be different in view of the known clinicopathologic differences that exist between them. Thus, more comprehensive researches are needed. The use of next-generation sequencing (NGS) for high-throughput genomic analysis has accelerated our understanding of the molecular characteristics of CRC<sup>12-14</sup> and may also serve us an insight on this topic.

Therefore, we conducted NGS of primary tumors of synchronous or metachronous CLM to investigate mutational patterns between them. We also expected to find prognostic or predictive biomarkers among mutations sequenced.

## Patients and methods

### Study population

This study retrospectively included patients with CRC treated at General Surgery Department of Zhongshan Hospital, Fudan University (Shanghai, China) during June 2008 to December 2014. The inclusion criteria were as follows: colorectal adenocarcinoma determined by pathologic evidence; wild-type RAS; liver metastases determined by radiologic and/or pathologic evidence; radical resections of primary tumors; enough formalin-fixed paraffin-embedded (FFPE) tissue of primary tumors; no exposure to any treatment (chemotherapy, targeted therapy, radiotherapy, or interventional therapy) before primary tumor resections. Patients with liver metastases accompanied with other distant metastases were permitted. Adjuvant chemotherapy after primary tumor resection was permitted. Target therapy and interventional therapy were permitted only after the occurrence of liver metastases. All patients provided written and oral informed consent. This study was approved by the ethics committee of Zhongshan Hospital, Fudan University.

Two groups were established in this study: the synchronous-metastases group and the metachronous-metastases

group. The synchronous-metastases group consisted of patients with liver metastases diagnosed together with or within a 6-month interval of the diagnosis of the primary colorectal tumor. The metachronous-metastases group consisted of patients with liver metastases diagnosed >6 months after primary tumor resection. This study was approved by the local ethics committees and all patients provided written and oral informed consent, including research on tumor tissue.

In addition, 10 patients were selected for whole-exome sequencing (WES). All 10 patients were diagnosed with CLM and underwent resection of both primary and metastatic tumors. Exclusion criteria included previous exposure to any treatment for metastatic CRC, family history of CRC, and evidence of a mismatch repair deficiency.

### Study procedure

To investigate mutational pattern of different liver metastases, we examined biomarkers (single-nucleotide polymorphism, SNP) through genome-wide exploration using NGS. To preliminarily select genes to construct a custom panel for target capture sequencing, we performed WES for 10 triplets, each comprising primary colorectal tumor and normal colorectal mucosa and matched liver metastases. Genomic DNA from fresh tissue was samples sequenced on an Ion<sup>TM</sup> Proton (Life Technologies, Carlsbad, CA, USA) platform according to the manufacturer's instructions. Normal colorectal mucosa was sequenced to exclude germline variants. The read alignments and variant analyses were performed according to the predefined workflow.

We constructed a custom panel of 96 genes selected based on driver mutations identified using WES and Tumor Mutation Hotspots Panel version 2 (Life Technologies, Carlsbad, CA, USA). Genomic DNA from FFPE tissue samples of patients in both cohorts was subsequently sequenced for SNPs using an Ion<sup>TM</sup> Torrent Personal Genome Machine (PGM) according to the manufacturer's instructions. For a given gene loci, the fraction of mutant alleles was calculated by dividing the number of mutant reads by the number of total reads. A 5% cutoff value was employed. A sample was considered wild-type for a given gene when all sequenced loci harbored <5% mutant alleles.

### Whole-exome sequencing

DNA was extracted from fresh tumor samples using the MELT<sup>TM</sup> Total Nucleic Acid Isolation Kit (Life Technologies). Quantity and quality were assessed using Qubit 2.0 (Life Technologies). Fifty to hundred nanograms of DNA

for each sample was used for exome capture and library preparation with Ion AmpliSeq™ Exome Kit 4xDuo (Life Technologies) following the manufacturer's instructions. Then libraries were bar-coded with Ion Xpress™ Barcode Adapters Kit (Life Technologies). The concentration of each library was determined by PCR with the Ion Library™ Quantization Kit (Life Technologies). According to the manufacturer's instructions, all libraries were diluted to 100 pM working solutions and then pooled as needed to perform the template preparation with Ion PI™ Template OT2 200 kit v2 (Life Technologies) on Ion One Touch™ 2 System. Quality and quantity were determined with Qubit Ion Sphere™ Quality Control Kit (Life Technologies) for the obtained ion sphere particles. WES was performed on Ion Proton™ platform, using the Ion PI™ Sequencing 200 kit v2 and Ion PI™ Chip kit v2. Sequencing data were analyzed with the Torrent Suite™ Software v4.0 (Life Technologies) using default parameters setting.

## Processing of WES data

The primary WES data were analyzed for single-nucleotide variants (SNVs) following the procedures indicated below.

1. Variants with SNV quality (QUAL)  $\leq 20$  were excluded. QUAL was calculated using Torrent Suite™ software.
2. Variants of normal mucosa were considered background variants. Primary or metastatic tumor samples were filtered using background variants, and the variants were rejected as germline variants or sequencing artifacts when present in the corresponding normal samples.
3. Primary or metastatic tumor-specific SNVs were analyzed using the SeattleSeq SNP Annotation.<sup>15</sup> Known germline mutations from the Exome Sequencing Project<sup>16</sup> and dbSNP databases (build 140)<sup>17</sup> were also excluded. We selected nonsilent mutations, including missense mutations and InDels.
4. Nonsilent mutations were predicted to affect gene function when any of the following criteria were fulfilled: 1) functional impact score of SIFT<sup>18</sup>  $\leq 0.05$ ;<sup>19</sup> 2) functional impact score of PolyPhen-2<sup>20</sup>  $> 0.45$ ;<sup>21</sup> 3) functional impact label of Mutation Assessor<sup>22</sup> was “medium” or “high”;<sup>23</sup> 4) Condel<sup>24</sup> label was “deleterious”;<sup>25</sup> and 5) functional impact score of FATHMM<sup>26</sup>  $< 0.27$
5. The transFIC analysis was performed as previously described.<sup>28</sup> Mutations were considered cancer driver mutations when the outcome of the transFIC analysis was of “high impact”.

## Construction of custom panel

To prepare for target capture sequencing, we constructed a custom panel based on WES data and Tumor Mutation Hotspots Panel version 2. When selecting genes from WES data, genes that were essential for cancer progression, particularly liver metastasis, were considered with priority. On one hand, we searched all mutations in the GeneRIF database<sup>29</sup> using key words “cancer/tumor/carcinoma” and “metasta-/invasion/invade/invasive/migrate”. On the other hand, we focused on mutations of universal genes in primary tumors and corresponding liver metastases. In addition, most genes in the Tumor Mutation Hotspots Panel version 2 were included. The mutation information for these genes was acquired from the Catalog of Somatic Mutations in Cancer (COSMIC) database,<sup>30</sup> and we employed the most frequent mutations to build the panel.

## Target capture sequencing

DNA was extracted from FFPE tumor samples using the RecoverAll™ Total Nucleic Acid Isolation Kit (Life Technologies) according to the manufacturer's instructions. Quantity and quality were assessed using Qubit 2.0 (Life Technologies). Ten nanograms of DNA for each sample was used for library construction and template preparation with same procedures described above in the “Whole-exome sequencing” section. Target capture sequencing was carried out with customized panel using the Ion PGM™ platform (Life Technologies) according to the manufacturer's instructions. The panel consisted of two separate PCR primer pools covering recurrent mutations in 96 genes with 1500X sequence coverage on Ion™ 318 chip. Sequencing data were analyzed with Ion Reporter™ software v4.4 (Life Technologies) using default parameters setting.

## Statistical analysis

The statistical analysis plan was established before the genotyping results were available. Differences in categorical parameters were calculated using a chi-square test or Fisher's exact test. Survival curves were generated using the Kaplan–Meier method and compared using a log-rank test. Hazard ratios and 95% CIs were calculated using the Cox proportional hazards model. For univariate and multivariate analyses of latency of metachronous liver metastases, the Cox proportional hazards model was used. All statistical analyses were conducted using the statistical software SPSS version 18.0 (SPSS Inc., Chicago, IL, USA). A *p*-value  $< 0.05$  was considered statistically significant.

## Results

### Patients and clinical characteristics

A total of 161 patients were included and sequenced. According to the metastatic type, 93 patients were in synchronous-metastases group and 68 patients in metachronous-metastases group. Several differences in clinical characteristics were observed between groups (Table 1). Patients with metachronous CLMs were obviously elder. With regard to pathology of primary tumors, synchronous CLMs were larger in size, poorly differentiated, and more frequently local advanced and lymph node positive. As to evaluation of liver metastases, synchronous CLMs result in more and larger metastatic lesions.

### Summary of WES and target capture sequence

Ten triplets were sequenced with WES for preliminarily data to construct custom panel. Overall, we identified 608 potential somatic driver mutations in 511 genes in primary colorectal tumors and 694 potential somatic driver mutations in 638 genes in liver metastases. The number of somatic mutations in 10 primary colorectal tumor ranged from 41 to 87, with a mean of 60 (Figure 1A), which was not significantly different from that of the non-hypermutated CRCs reported in The Cancer Genome Atlas.<sup>31</sup> When comparing mutations between tumors, 230 mutations in 219 genes were universal in the primary tumor and corresponding liver metastases. In addition, mutations observed in 10 patients were predominated by the C/G>T/A transition (Figure 1B) consistent with the results of previous CRC genomics studies.<sup>31,32</sup>

According to the custom panel, 161 patients were sequenced with target capture sequence for recurrent mutations of 96 genes. Mutations with mutational frequencies >10% and <90% were further investigated for stability. Finally, 48 mutations in 27 genes were included in following analysis (Table 2). Among these mutations, 40 were already documented in dbSNP databases (build 140).

### Correlation between clinical characteristics and mutations

Univariate analysis identified that 18 mutations in 12 genes were correlated with clinical variables (Table 3). MDC1 was associated with the most number of clinical variables, including T stage, N stage, tumor location of primary, and size and number of liver metastases. Mutations of ATAD3B, MAGEC1, and MICB were only associated with pathologic

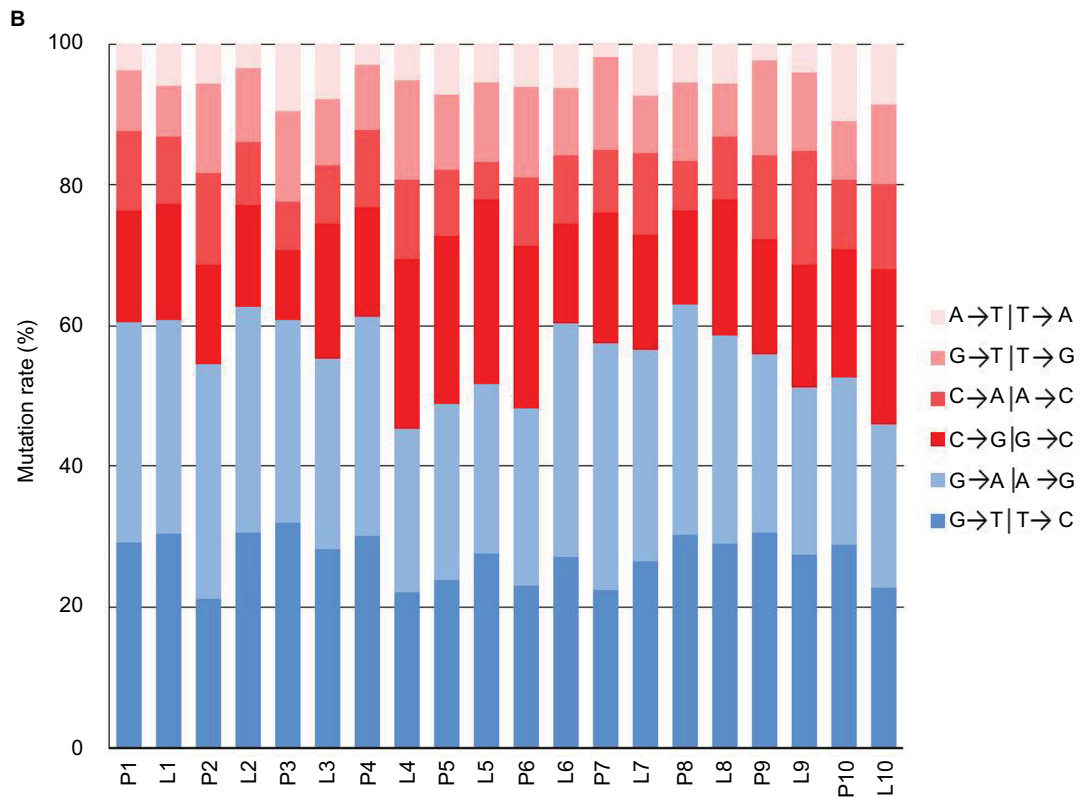
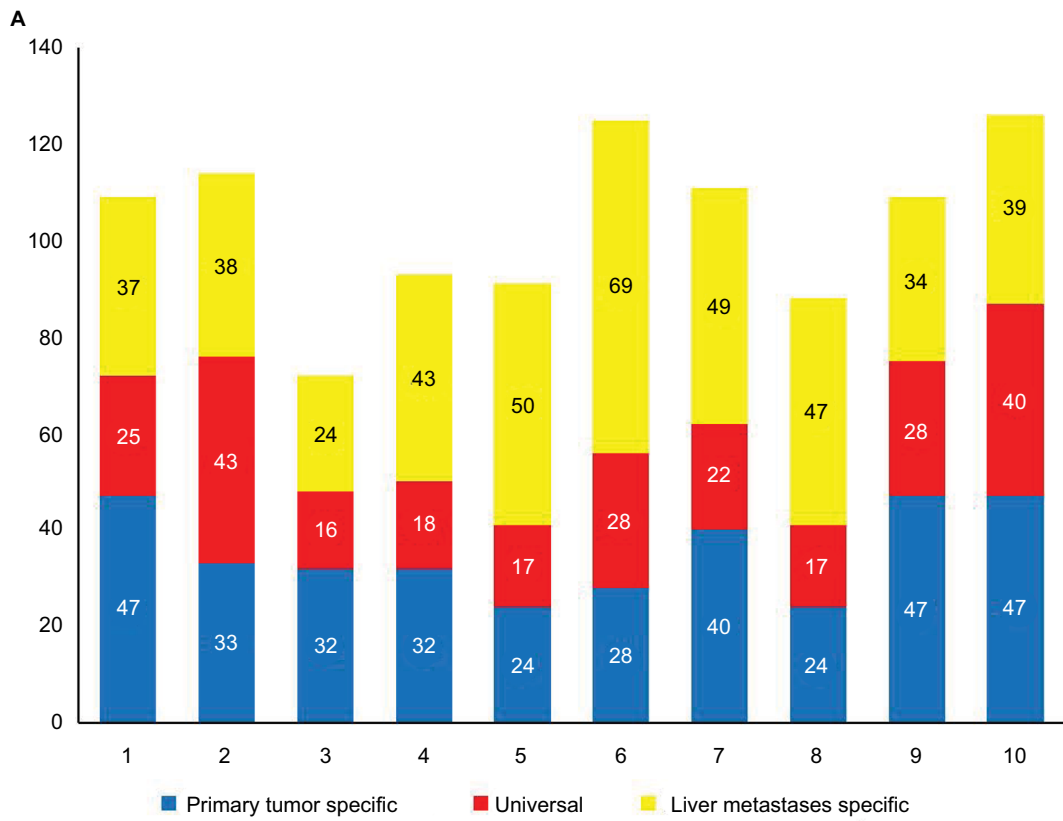
**Table 1** Clinical characteristics of synchronous and metachronous groups

Clinical variables	Synchronous (N=93)	Metachronous (N=68)	p-value
Age (years), mean±SD	55.1±10.3	63.4±11.8	<0.001
Gender, n (%)			0.652
Male	62 (66.7%)	43 (63.2%)	
Female	31 (33.3%)	25 (36.8%)	
CEA level at diagnosis, ng/mL, n (%)			<0.001
≥5	79 (84.9%)	21 (30.9%)	
<5	14 (15.1%)	47 (69.1%)	
Primary tumor location, n (%)			0.428
Right-sided	30 (32.3%)	18 (26.5%)	
Left-sided	63 (67.7%)	50 (73.5%)	
Tumor diameter (cm), mean±SD	5.1±2.1	4.2±1.8	0.011
Histologic grade, n (%)			0.016
Well (Grade 1)	2 (2.2%)	7 (10.3%)	
Moderate (Grade 2)	59 (63.4%)	48 (70.6%)	
Poor (Grades 3 and 4)	32 (34.4%)	13 (19.1%)	
pT stage, n (%)			0.004
T1/T2	3 (3.2%)	13 (19.1%)	
T3/T4	90 (96.8%)	55 (80.9%)	
pN stage, n (%)			<0.001
N0	15 (16.1%)	43 (63.2%)	
N1	43 (46.2%)	18 (26.5%)	
N2	35 (37.6%)	7 (10.1%)	
Tumor deposits, n (%)			0.001
No	47 (50.5%)	53 (77.9%)	
Yes	46 (49.5%)	15 (22.1%)	
Distribution of LM, n (%)			<0.001
Unilobar	34 (36.5%)	47 (69.1%)	
Bilobar	59 (63.5%)	21 (30.9%)	
Numbers of LM			<0.001
Median (IQR)	5 (3–10)	2 (1–4)	
Diameter of the largest LM, cm			0.007
Median (IQR)	38 (27–69)	33 (14–52)	
Accompany with other metastases, n (%)			
Lung	10 (12.9%)	5 (7.3%)	0.764
Retroperitoneal LN	6 (6.4%)	2 (2.9%)	0.598
Others*	3 (3.2%)	2 (2.9%)	0.995

**Note:** \*Includes omentum, ovary, and brain.

**Abbreviations:** CEA, carcinoembryonic antigen; IQR, interquartile range; LM, liver metastases; LN, lymph node.

variables of primary tumor. Most of the mutations associated with variables of liver metastases were also associated with lymph node metastases. Ten correlations with preoperative



**Figure 1** Summary of driver mutations identified by whole-exome sequencing.

**Notes:** (A) Distribution of driver mutations in primary tumor and liver metastases. (B) Distribution of transitions and transversions.

**Abbreviations:** L, liver metastases; P, primary tumor.

**Table 2** List of mutations analyzed

Gene	Single Nucleotide Polymorphism database (dbSNP)	cDNA	Coding sequence (CDS)	Protein	Amino acids	Codons
ACTN4	–	1,276	1,200	400	W/C	tgG/tgC
ATAD3B	rs860213	1,273	1,157	386	R/Q	cGg/cAg
ATAD3B	rs139902189	1,851	1,735	579	R/C	Cgc/Tgc
ATP6V1B1	rs17720303	162	89	30	T/I	aCc/aTc
COL2A1	rs2070739	4,378	4,213	1,405	G/S	Ggc/Agc
CUL9	rs2273709	5,918	5,843	1,948	H/P	cAc/cCc
EGFR	rs2227983	1,586	1,403	468	R/K	aGg/aAg
ERBB2	rs1058808, rs370420724	3,463	3,463	1,155	P/A	Ccc/Gcc
ERBB2	rs1136201	281	283	95	I/V	Atc/Gtc
EZH2	rs2302427	648	526	176	D/H	Gac/Cac
FAM129A	–	2,020	1,826	609	L/P	cTg/cCg
FCGBP	rs11083543	4,026	4,018	1,340	V/L	Gtg/Ctg
FCGBP	rs75388508	52	44	15	T/N	aCc/aAc
HNF1A	rs1169288	222	223	75	I/L	Atc/Ctc
KRT33A	rs12937519	854	809	270	A/V	gCg/gTg
LAMA4	rs1050349	3,754	3,356	1,119	P/R	cCt/cGt
LY6G6D	rs118062293	334	334	112	R/C	Cgt/Tgt
MAGEC1	rs12558365	1,008	722	241	S/F	tCc/tTc
MAGEC1	rs176037	738	452	151	T/I	aCt/aTt
MAP3K19	rs3905317	2,466	2,435	812	E/G	gAa/gGa
MAP3K19	rs1112542	2,057	2,026	676	E/Q	Gag/Cag
MDC1	–	5,320	4,672	1,558	G/S	Ggc/Agc
MDC1	rs144087810	6,265	5,617	1,873	P/A	Ccc/Gcc
MDC1	rs61733213	4,595	3,947	1,316	M/R	aTg/aGg
MDC1	rs9461623	719	721	241	S/P	Tct/Cct
MDC1	–	1,975	1,535	512	R/K	aGa/aAa
MDN1	rs4707569	1,434	1,318	440	F/V	Ttt/Gtt
MEGF6	rs7513275	571	344	115	M/T	aTg/aCg
MEGF6	rs7553399	2,974	2,747	916	R/L	cGg/cTg
MEGF6	rs4648506	3,637	3,410	1,137	G/A	gGc/gCc
MICB	rs1065075	346	238	80	K/E	Aag/Gag
MICB	rs1051788	451	310	104	D/N	Gat/Aat
MICB	rs3134900	408	267	89	I/M	atC/atG
PTCH1	rs357564	4,086	3,941	1,314	P/L	cCc/cTc
PTPN23	rs6780013	2,788	2,452	818	A/T	Gca/Aca
PTPN23	–	3,697	3,361	1,121	S/C	Agc/Tgc
RBMXL3	rs12399211	1,234	1,192	398	D/N	Gac/Aac
RBMXL3	–	3,059	3,017	1,006	G/D	gGc/gAc
RBMXL3	rs6643947	3,187	3,145	1,049	R/G	Agg/Ggg
TAPBP	rs2071888	731	518	173	T/R	aCa/aGa
TAPBP	–	421	208	70	V/M	Gtg/Atg
TAPBP	–	597	251	84	R/P	cGg/cCg
TCF3	rs2074888	1,972	1,475	492	A/V	gCg/gTg
TYK2	rs2304256	1,566	1,084	362	V/F	Gtc/Ttc
WDSUB1	rs7591849	1,107	958	320	R/S	Cgc/Agc
WDSUB1	rs16843852	793	644	215	K/T	aAa/aCa
ZNF462	rs3814538	5,772	5,483	1,828	N/S	aAc/aGc
ZNF462	rs17723637	1,499	1,210	404	M/V	Atg/Gtg

**Table 3** Clinical variables associated with gene mutations

Clinical variable	Mutation	Mutation frequency (vs. no risk variable)	p-value	q-value
Age $\geq 65$ years	PTPN23 c.3361A>T	60.5% (26/43) (vs. 33.9% [40/118])	0.002	0.096
Male gender	KRT33A rs12937519	59.0% (62/105) (vs. 41.1% [23/56])	0.030	0.480
	MAGEC1 rs176037	23.8% (25/105) (vs. 42.9% [24/56])	0.012	0.576
	MICB rs3134900	8.6% (9/105) (vs. 21.4% [12/56])	0.012	0.288
CEA at diagnosis $\geq 5$ ng/mL	ATAD3B rs860213	62% (62/100) (vs. 32.8% [20/61])	<0.001	0.008
	TCF rs2074888	68% (68/100) (vs. 49.2% [30/61])	0.018	0.144
	MAP3K19 rs3905317	61% (61/100) (vs. 42.6% [26/61])	0.023	0.159
	COL2A1 rs2070739	64% (64/100) (vs. 47.5% [29/61])	0.040	0.192
	MEGF6 rs7513275	51% (51/100) (vs. 26.2% [16/61])	0.002	0.032
	MDC1 c.4672G>A	53% (53/100) (vs. 36.1% [55/61])	0.037	0.196
	PTPN23 c.3361A>T	30% (30/100) (vs. 59.0% [36/61])	<0.001	0.012
	MDC1 rs61733213	29% (29/100) (vs. 47.5% [29/61])	0.017	0.163
	MAGEC1 rs176037	22% (22/100) (vs. 44.3% [27/61])	0.003	0.036
Right-sided tumors	MDC1 c.1535G>A	10% (10/100) (vs. 23.0% [14/61])	0.025	0.150
	MDC1 c.4672G>A	62.5% (30/48) (vs. 39.8% [45/113])	0.008	0.384
Poor histologic grade	ATAD3B rs860213	64.4% (29/45) (vs. 45.7% [53/116])	0.033	0.528
	KRT33A rs12937519	68.9% (31/45) (vs. 46.6% [54/116])	0.011	0.528
	ATAD3B rs139902189	44.4% (20/45) (vs. 26.7% [31/116])	0.030	0.720
	MAGEC1 rs12558365	33.3% (15/45) (vs. 18.1% [21/116])	0.037	0.444
T3/T4	MICB rs1065075	33.8% (49/145) (vs. 62.5% [10/16])	0.024	0.576
	MICB rs3134900	11.0% (16/145) (vs. 31.3% [5/16])	0.023	–
Lymph node positive	ATAD3B rs860213	59.2% (61/103) (vs. 36.2% [21/58])	0.005	0.048
	MEGF6 rs7513275	47.6% (49/103) (vs. 31.0% [18/58])	0.041	0.246
	RBMXL3 rs12399211	53.4% (55/103) (vs. 75.9% [44/58])	0.005	0.060
	MDC1 rs144087810	38.8% (40/103) (vs. 13.8% [8/58])	0.001	0.024
	PTPN23 c.3361A>T	29.1% (30/103) (vs. 62.1% [36/58])	<0.001	0.020
	MDC1 rs61733213	27.2% (28/103) (vs. 51.7% [30/58])	0.002	0.032
	MAGEC1 rs176037	23.3% (24/103) (vs. 43.1% [25/58])	0.009	0.617
	MICB rs3134900	7.8% (8/103) (vs. 22.4% [13/58])	0.008	0.064
Numbers of LM $\geq$ median level	MEGF6 rs7513275	47.8% (54/113) (vs. 27.1% [13/48])	0.015	0.240
	RBMXL3 rs12399211	54.0% (61/113) (vs. 79.2% [38/48])	0.003	0.072
	MDC1 rs144087810	34.5% (39/113) (vs. 18.8% [9/48])	0.045	0.540
	PTPN23 c.3361A>T	32.7% (37/113) (vs. 60.4% [29/48])	0.001	0.038
Diameter of the largest LM $\geq$ median level	HNFA1 rs1169288	68.8% (64/93) (vs. 48.5% [33/68])	0.009	0.413
	MDC1 c.4672G>A	53.8% (50/93) (vs. 36.8% [25/68])	0.033	0.531
	MDC1 rs61733213	43.0% (40/93) (vs. 26.5% [18/68])	0.031	0.740

**Note:** q-value for adjusted p-value with false discovery rate (Benjamini–Hochberg procedure).

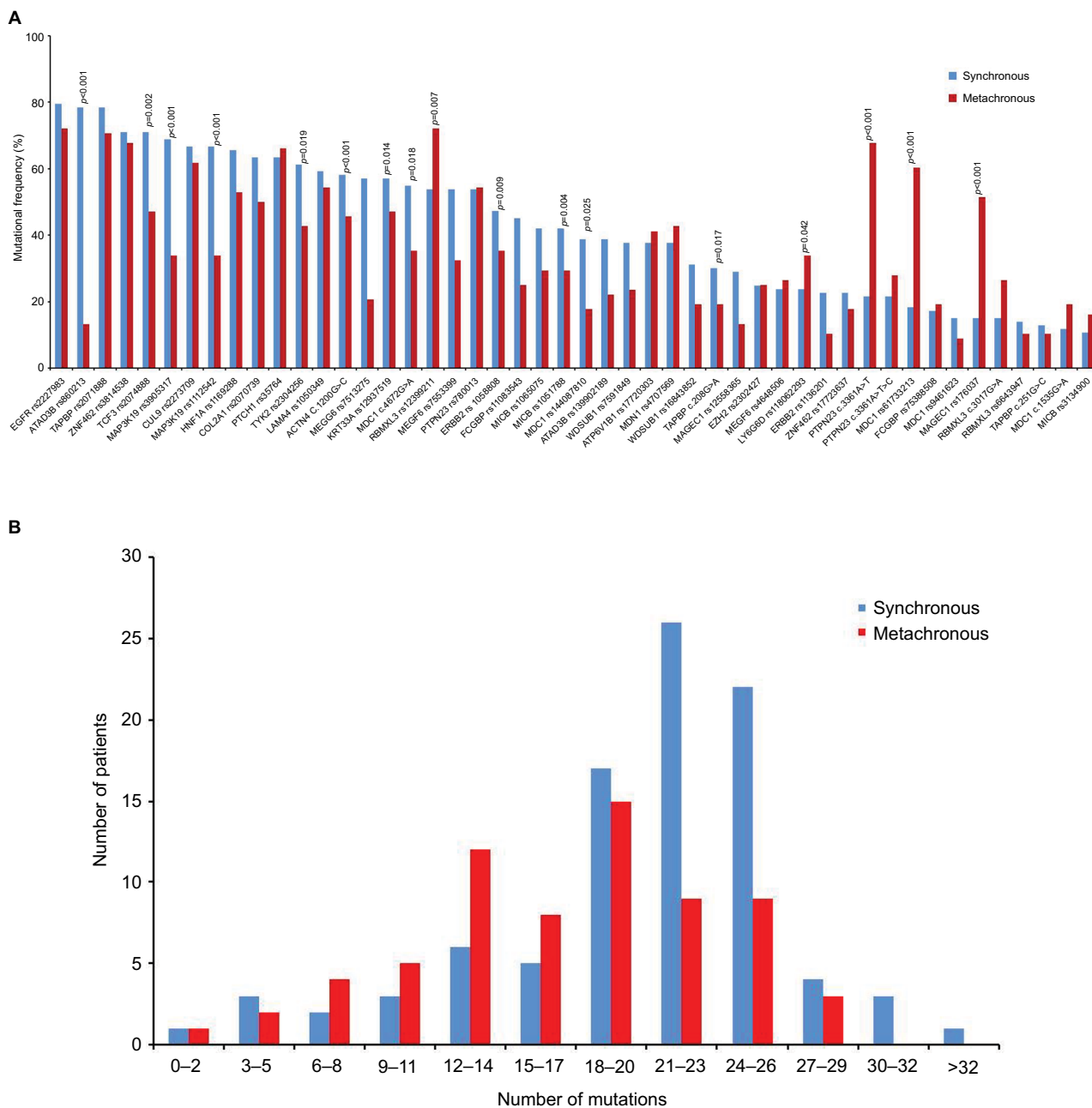
**Abbreviations:** CEA, carcinoembryonic antigen; LM, liver metastases.

CEA level, lymph node metastases, and number of LMs remained significant with false discovery rate <0.05.

### Mutational pattern of synchronous and metachronous liver metastases

The median number of mutations in the synchronous group was 22 (range from 2 to 32), which was significantly higher than 18 (range from 2 to 28) in the metachronous

group ( $p < 0.001$ ). Mutation frequency of the synchronous group, compared with the metachronous group, was significantly higher in 12 mutations and lower in five mutations (Figure 2). For most other mutations, the frequency was numerically higher in the synchronous group. In addition, only half of the most prevalent mutations in the synchronous group were in common with that in the metachronous group (Table 4).



**Figure 2** Mutation frequencies of synchronous and metachronous CLMs. **Notes:** (A) Mutation frequency of most sequenced genes were different between synchronous and metachronous CLMs, and 17 of them were statistically significant. (B) Synchronous CLMs carried significantly more mutations than metachronous CLMs (median number, 22 vs 18,  $p < 0.001$ ). **Abbreviation:** CLMs, colorectal liver metastases.

## Mutations and survival of metachronous liver metastases

The latency of metachronous CLMs is summarized in Figure 3. Half of all metachronous CLMs occurred within 15 months, and 75% occurred within 24 months. For patients with metachronous CLMs, the latency was significantly associated with prevalent mutations in metachronous CLMs, including EGFR rs2227983, RBMXL3 rs12399211, TAPBP rs2071888, PTPN23 c.3361A>T, PTCH1 rs357564, and MDC1 rs61733213. In multivariate analysis, correlation between

latency and EGFR rs2227983, RBMXL3 rs12399211, and PTCH1 rs357564 remained significant (Table 5).

## Discussion

In this study, we conducted NGS in primary tumors of synchronous or metachronous CLM and identified several differences in terms of clinical characteristics and mutational patterns.

For clinicopathologic characteristics, our results were broadly consistent with previous studies.<sup>2,5</sup> Patients from



**Table 4** Prevalent mutations in synchronous and metachronous groups

Prevalent mutations in synchronous group		Prevalent mutations in metachronous group	
Mutation	Frequency (vs. metachronous group) (%)	Mutation	Frequency (vs. synchronous group) (%)
EGFR rs2227983	79.5 (vs. 72.1)	EGFR rs2227983	72.1 (vs. 79.6)
ATAD3B rs860213	78.4** (vs. 13.2)	RBMXL3 rs12399211	72.1** (vs. 53.8)
TAPBP rs2071888	7.84 (vs. 70.6)	TAPBP rs2071888	70.6 (vs. 78.5)
ZNF462 rs3814538	71.0 (vs. 67.6)	ZNF462 rs3814538	67.6 (vs. 71.0)
TCF3 rs2074888	71.0* (vs. 47.1)	PTPN23 c.3361A>T	67.6** (vs. 21.5)
MAP3K19 rs3905317	68.8* (vs. 33.8)	PTCHI rs357564	66.2 (vs. 63.4)
CUL9 rs2273709	66.7 (vs. 61.7)	CUL9 rs2273709	61.7 (vs. 66.7)
MAP3K19 rs1112542	66.7** (vs. 33.8)	MDC1 rs61733213	60.3** (vs. 18.3)

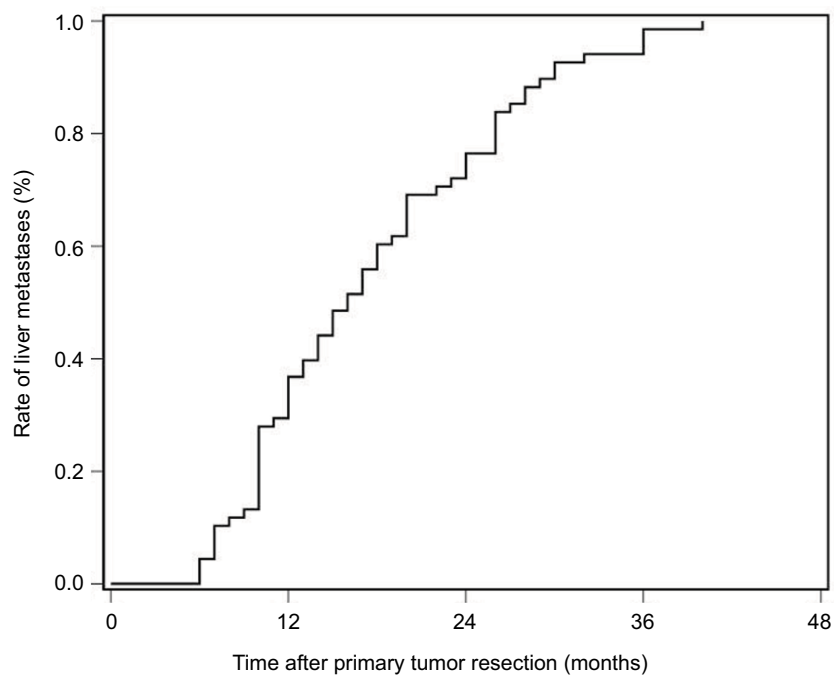
Notes: \* $p < 0.05$ ; \*\* $p < 0.001$ .

the synchronous group were younger than those from the metachronous group. Compared to the metachronous group, patients in the synchronous group showed heavier tumor burden for both primaries and liver metastases. Furthermore,

the synchronous group, more frequently characterized by adverse prognostic factors, had inferior overall survival than the metachronous group (median time 26 vs. 36 months,  $p = 0.001$ ). These results indicated that synchronous CLM may represent a more aggressive tumor subtype than metachronous CLM. All these issues highlight the need for understanding of differences in molecular biology, which results in and keeps the different manifestations between synchronous and metachronous CLMs.

Based on the understanding of the mechanisms of metastases, we extracted our efforts to identify biologic differences between synchronous and metachronous CLMs. In contrast to previous studies focusing on limited gene alterations,<sup>2,8,9,33</sup> we used a multivariable approach, including genome-wide exploration using WES, panel construction based on bioinformatics, and selection of significant biomarkers according to clinical outcomes. The advantage of this multivariable approach is that it is unbiased by biologic assumptions and thereby may find correlations between distinct gene mutations and liver metastases.

Finally, we included and analyzed 48 mutations in 27 genes. The synchronous group harbored significantly more mutations than the metachronous group, representing a more heterogeneous and advanced subgroup of CLM. For the synchronous group, about half of the prevalent mutations were in common with the metachronous group, and the most prevalent was EGFR rs2227983. These mutations

**Figure 3** Timing of metachronous metastases.

Note: Half of all metachronous colorectal liver metastases occurred within 15 months, and 75% occurred within 24 months.

**Table 5** Univariate and multivariate analyses of latency of metachronous colorectal liver metastases

Mutation		Univariate analyses		Multivariate analyses	
		HR (95% CI)	p-value	HR (95% CI)	p-value
EGFR rs2227983	Wild-type/mutant	0.43 (0.23–0.78)	0.006	0.47 (0.25–0.88)	0.019
RBMXL3 rs12399211	Wild-type/mutant	0.36 (0.19–0.67)	0.001	0.41 (0.18–0.92)	0.030
TAPBP rs2071888	Wild-type/mutant	0.49 (0.28–0.85)	0.011	0.71 (0.36–1.40)	0.324
PTPN23 c.3361A>T	Wild-type/mutant	0.49 (0.28–0.87)	0.015	1.71 (0.64–4.56)	0.287
PTCH1 rs357564	Wild-type/mutant	0.39 (0.22–0.669)	0.001	0.50 (0.28–0.91)	0.022
MDC1 rs61733213	Wild-type/mutant	0.56 (0.33–0.95)	0.033	0.88 (0.38–2.04)	0.760

**Abbreviation:** HR, hazard ratio.

were potential key mutations of liver metastases and further research was needed. Another half of the prevalent mutations were private in the synchronous group and were significantly more frequently mutant than the metachronous group. In addition, prevalent mutations private in the metachronous group showed similar trend. Furthermore, these mutations, including RBMXL3 rs12399211, PTPN23 c.3361A>T, PTCH1 rs357564, and MDC1 rs61733213, partly explained the latency of metachronous liver metastases. All these results indicated that the synchronous and the metachronous groups showed different mutational patterns, and these mutations were correlated with several clinical characteristics. Different mutations may be potential key mutations and further researches were needed.

In previous studies on this topic, the majority found differences in molecular marker expression between CLMs and their respective primaries in both the synchronous and metachronous groups. Limited comparison was done between LMs or primaries between the synchronous and the metachronous groups. Pantaleo et al<sup>8</sup> compared the expression signature with reverse transcriptase PCR and enzyme-linked immunosorbent assay and indicated that EGFR and COX-2 are overexpressed in metachronous and synchronous metastases, respectively. van der Wal et al<sup>2</sup> reported that liver parenchyma adjacent to the LMs provides a highly prosperous angiogenic environment in the synchronous group compared to the metachronous groups. However, there were also many studies with negative findings.<sup>34,35</sup> Our results added evidences for the hypothesis that synchronous and metachronous CLMs were biologically different.

According to previous studies, both synchronous and metachronous CLMs would evolve different biologic alterations to their corresponding primaries.<sup>11,32,33,36</sup> Thus, examination of metastatic tissue is crucial as it could help choose optimal treatment according to current biologic status. Simultaneously, studies also demonstrated that the majority

of biologic alterations, especially key alterations, in primaries were maintained in CLMs.<sup>11,32,35</sup> If these key alterations of liver metastases could be detected in primaries, it could have important clinical implications for prediction of occurrence, even timing of liver metastases. Under consideration of the evidence mentioned above, we selected and examined tissue of primary tumors. If only corresponding metastatic tissue were examined, our study may provide more information.

This was a retrospective exploratory study and there were several limitations: first, we sequenced single type of gene alteration in a highly selected cohort. Only SNPs of primary tumors with wild-type RAS were sequenced. Thus, this study provided an incomplete and partial view on differences between different CLMs; second, many correlations were observed between clinical parameters and mutations, but relevant biologic evidences were still limited. More studies were needed to confirm the clinical meanings and explore the underlying mechanism. Third, the influences of tumor heterogeneity and differences between LMs and their respective primaries were not taken into consideration in the study.

## Conclusion

Synchronous CLMs, compared to metachronous CLMs, were younger and showed heavier tumor burden for both primaries and liver metastases. NGS identified different mutational patterns between synchronous and metachronous CLMs. Further researches were needed to confirm the correlation between differences in clinical characteristics and mutational patterns and the mechanism inside.

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## Disclosure

The authors report no conflicts of interest in this work.

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