Identification of DNA methylation biomarkers for risk of liver metastasis in early-stage colorectal cancer

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Abstract

Background: Liver metastases can occur even in CRC patients who underwent curative surgery. While evidence suggested that adjuvant chemotherapy can help to reduce the occurrence of liver metastases for certain patients, it is not a recommended routine as the side effects outweigh the potential benefits, especially in Stage II CRC patients. This study aims to construct a model for predicting liver metastasis risk using differential methylation signals in primary CRC tumors, which can facilitate the decision for adjuvant chemotherapy.

Methods: Fifty-nine stage I/II and IV CRC patients were enrolled. Primary tumor, adjacent normal tissue, and metastatic tumor tissues were subject to targeted bisulfite sequencing for DNA methylation. The Least Absolute Shrinkage and Selection Operator (LASSO) algorithm was used to identify potential DMRs for predicting liver metastasis of CRC.

Results: We identified a total of 241,573 DMRs by comparing the DNA methylation profile of primary tumors of stage II patients who developed metastasis to those who were metastasis-free during the follow up period. 213 DMRs were associated with poor prognosis, among which 182 DMRS were found to be hypermethylated in the primary tumor of patients with metastases. Furthermore, by using the LASSO regression model, we identified 23 DMRs that contributed to a high probability of liver metastasis of CRC. The leave-one-out cross validation (LOOCV) was used to evaluate model predictive performance at an AUC of 0.701. In particular, 7 out of those 23 DMRs were found to be in the promoter region of genes that were previously reported prognostic biomarkers in diverse tumor types, including TNNI2, PAX8, GUF1, KLF4, EVI2B, CEP112, and long non-coding RNA AC011298. In addition, the model was also able to distinguish metastases of different sites (liver or lung) at an AUC of 0.933.

Conclusion: We have identified DNA methylation biomarkers associated with the risk of cancer liver metastasis in early-stage CRC patients. A risk prediction model based on those epigenetic markers was proposed for outcome assessment.
Background

Colorectal cancer (CRC) is one of the most common malignancies worldwide and causes the fifth top cancer mortality in China [1, 2]. The incidence of CRC is associated with age-increasing and a slightly higher risk is found in men than women [3]. The 5-year survival rate for patients with localized CRC is optimistic, while it drops dramatically when distant metastases are developed [4]. Among all possible metastatic sites, liver is the most popular one due to the connection between the intestinal mesenteric drainage and the hepatic portal venous system [5]. Surgical resection is the only treatment needed for stage I CRC [4]. Considering the side effects and the potential benefits, adjuvant chemotherapy is not a recommended routine use in stage II CRC as the overall survival and disease-free survival were not significantly improved, but a decreased trend of tumor relapse was observed [6, 7]. Therefore, it is still debatable about the administration of adjuvant chemotherapies to post-operative stage II CRC patients. According to NCCN Guidelines, the risk assessment for adjuvant-treatment-decision is mainly based on clinical prognosis features such as poorly differentiated histology and lymphatic/vascular invasion. However, this may lead to overtreatment or miss the best timing of treatment as these features are insufficient for early prediction of metastasis. Identification of predictive biomarkers or high-risk molecular features to closely monitor disease progression and metastasis which are critical for determining whether the adjuvant therapy should be administered is urgent for stage II CRC.

Next-generation sequencing is now playing an essential role in early disease detection and precision medicine. Besides genomic alterations, it has been applied to study the epigenetic changes during carcinogenesis and tumor metastasis. DNA methylation (DNAm) is one of the most investigated epigenetic mechanisms which establishes the genomic imprinting marks together with histone modifications and non-coding RNAs [8]. DNAm mainly occurs to the cytosine bases in the ’CpG’ sites, which are enriched in gene promoters [9]. By pre-treating DNA with bisulfite, DNAm can be examined at a single-base resolution using high-throughput sequencing [10]. As aberrant DNAm patterns are considered to be associated with cancer and other diseases, researchers have put a lot of effort into identifying differentially methylated regions (DMRs) to investigate pathology and establish reliable biomarkers for prognosis prediction.

DMRs are widely studied as potential biomarkers for the diagnosis, prognosis, and treatment response in multiple cancer types. For instance, OPCML (opioid-binding cell adhesion molecule) and FLRT2 (fibronectin leucine-rich transmembrane protein 2) were identified as novel DMRs to distinguish prostate tumors and normal tissues [11]. Similar results were found in lung cancer as well, where a list of DMR genes was identified in two patients by comparing tumor and normal tissue specimens [12]. The DNAm levels of a number of genes might contribute to the prediction of tumor progression in gastric cancer [13]. In ovarian cancer, hypermethylation of DMRs were detected in several tumor suppressor genes such as ARHI and PEG3 [14]. While in CRC, a hypomethylated DMR in IGF2 was reported to be associated with poor prognosis [15]. And a comprehensive methylation analysis of DMRs in CRC revealed the higher susceptibility of hypermethylation than hypomethylation in tumor tissues [16]. In this study, we sought to comprehensively study the DMR status in early-stage CRC patients and extracted dominant ones to establish a pilot model to predict liver metastases in stage II CRC.

Methods and materials

Patients recruitment and sample collection

We retrospectively studied CRC patients who received surgery on either primary or metastatic tumor at the Cancer Hospital, Chinese Academy of Medical Sciences between 2012 and 2018. An average 5-year follow-up evaluation was performed for each patient to confirm the status of liver metastasis till December 31, 2019. All patients have provided written informed consent. In total, 81 samples including tumor and tumor adjacent, were collected from 59 patients. All samples were shipped to the central laboratory of a clinical testing center (Nanjing Geneseeq Technology Inc., China) for targeted bisulfite sequencing.

Targeted bisulfite sequencing and identification of differentially methylated region (DMR)

To construct the sequencing libraries, 1 μg of DNA per formalin-fixed paraffin-embedded (FFPE) sample was extracted and then treated with bisulfite. Targeted bisulfite sequencing was then performed on Illumina Hiseq platform (Illumina, San Diego, CA) using pre-designed probes (SeqCap Epi CpGiant, Roche), which targeted a total of ~2.7 × 10^6 CpG sites within ~ 80.5 Mb of genome region. Raw sequencing data were first demultiplexed by bck2fastq and then trimmed by Trimmomatic.
as part of the quality control (QC) protocol [17]. The qualified reads were then mapped onto the human reference genome (GRCh37/UCSC hg19) using the bisulfite sequence aligner Bismark [18] after PCR duplicates removal by Picard toolkit (http://broadinstitute.github.io/picard/).

The methylKit package (version 1.12.0) was used to identify DMRs in R (version 3.6.3) [19]. CpG clusters were tiled into 1000 bp windows with a minimum coverage ≥ 2 to ensure better detection of DMRs based on parameters which were previously reported by Ziller et al. [20]. The methylation level of each DMR was calculated using the total methylated cytosines divided by the total CpGs within each window. Logistic regression was applied to calculate the methylation difference as well as the false discovery rates (FDR) between the test and control groups.

**Identification of prognostic DMRs in stage II CRC patients**

To identify potential methylation markers discriminating between stage II patients with favorable and unfavorable prognoses, we first identified DMRs using 38 stage II tumor samples (21 favorable vs 17 unfavorable). The DMRs, which were significantly different between samples with favorable and unfavorable prognoses (methylation differences ≥ 10%, q value ≤ 0.05), were selected for downstream analysis. We then annotated these DMRs using the latest gene annotation from the GENCODE project (release 19 for GRCh37/UCSC hg19) [21] and chose only DMRs located within the promoter regions, which were defined as 1500 bp upstream and 500 bp downstream of the transcription start sites, using BEDTools (version 2.26.0) [22]. Finally, we performed a Jonckheere trend test on the methylation levels of these DMRs, to identify any DMRs showing a significant trend in different cancer stages, including tumor-adjacent, stage I, II and IV. A total of 70 samples, which included the 38 stage II tumor samples as well as 32 additional samples (11 primary tumor adjacent, 11 stage I and 10 stage IV), were used and the final prognostic DMRs were selected if the trend was statistically significant (p value ≤ 0.05).

**Model predicting liver metastasis using primary CRC tumor samples**

We used a total of 59 primary tumor samples to construct a model for predicting liver metastasis (LIM) in CRC patients. These 59 patient samples, as shown in Additional file 3: Table S1, included 22 patients with LIM and 37 patients without LIM during the follow-up. The machine learning approach was based on the stacked generalized linear model (GLM) of three based models using gradient boosting (GBM), Random Forest and Deep learning algorithms.

To increase the performance of our base model, we have performed tenfold cross-validation based on the training dataset to optimize the base models as well as the GLM stacked model. Each sample was used as a validation set for the LASSO model during the LOOCV, while the rest 58 samples were kept as the training set. The DMRs between primary tumor samples from LIM patients and LIM-free patients were identified using the training dataset only. A selective number of DMRs (methylation differences ≥ 20%, q value ≤ 0.05) were then used as candidates for identifying diagnostic DMRs. XGBoost was used to predict the performance of diagnostic DMRs, and the probability score for the validation set was calculated [23]. In total, the LASSO model was performed 59 times, and the Receiver Operating Characteristic (ROC) curve was constructed using probability scores of all 59 samples.

After the LOOCV was finished, the LASSO model was applied using the entire 59 primary tumor samples. A total of 105 DMRs between patients with/without LIM were identified using the aforementioned 59 primary tumor samples. After applying the LASSO algorithm, we were able to identify 23 DMRs from the total 105 candidate DMRs as the optimal diagnostic markers.

**Model evaluation using liver/lung metastasis tumor samples**

To further evaluate the predicting power of our model, we used 6 liver metastasis tumor samples and 5 lung metastasis tumor samples from CRC patients. Among these 11 metastasis tumor samples, 21 of the 23 previously identified diagnostic markers were available. The probability scores of these 11 samples were calculated using the aforementioned predictive model. A ROC curve was constructed to evaluate the performance of the predictive model by examining if the liver metastasis status were correctly identified for the metastasis tumor samples.

**Results**

**Clinical characteristics of the 59 CRC patients**

In this study, a total of 59 patients who were diagnosed as CRC and treated at the Cancer Hospital, Chinese Academy of Medical Sciences were recruited. The general clinical characteristics of the cohort are summarized in Table 1. There were more male (36/59, 61.0%) than female (23/59, 39.0%) patients and the median age was 61 years old. The majority patients (42/59, 71.2%) were non-smokers. Nearly two-thirds (38/59, 64.4%) were initially diagnosed as stage II and 17 of them developed lung or liver metastasis (LIM:11, LUM: 6) during
the follow-up period. Metastasis was also developed in two of the 11 stage I patients (LIM:1, LUM:1). All of the stage IV patients (N = 10) were initially diagnosed with LIM; meanwhile, LUM was found in two of them. To sum up, LIM were found in 22 of the 29 patients (75.9%) with metastatic tumors, making it more dominant than LUM (31.0%, 2 patients have both LIM and LUM). The median liver-metastasis-free survival time for these 59 patients was 2000 days, while the 1-year liver-metastasis-free rate was 81.4%, as shown in Additional file 1: Figure S1. A total of 81 tissue samples were collected from the entire cohort including 59 primary CRC tumor samples, 11 tumor-adjacent samples, and 11 metastatic tumor samples (6 LIM and 5 LUM) and the detailed information was summarized in Additional file 3: Table S1.

### Identify potential DMRs as prognostic markers

To identify any DMRs markers associated with prognosis, we sought to analyze the methylation status of primary tumor samples, which were grouped based on prognosis. To reduce the disturbance of stages, we only focused on the 38 stage II patients in which the number of favorable and unfavorable prognosis were similar.

These 38 stage II patients were grouped into a test and control group to identify DMRs using the methylkit package (version 1.12.0) in R (version 3.6.3) [19]. The test group was comprised of 17 patients with unfavorable prognosis, who developed either LIM or LUM during the follow-up period. The other 21 patients without metastases were grouped into the control group. The primary tumor samples of these 38 stage II CRC patients were used to identify DMRs as described in Fig. 1. We were able to identify a total of 241,573 DMRs (detail data not shown) between the test group and the control group.

To better understand their potential functional impact, we then annotated our identified DMRs against the latest gene annotation for GRCh37/UCSC hg19 from the GENCODE project (release 19) [21]. In total, 85,668 DMRs were found to be overlapping with a promoter region (Fig. 2B), which was defined as 1500 bp upstream to 500 bp downstream of a transcription start site. We then performed a pathway enrichment test using a subset of 26,105 DMRs, of which the q values met the cutoff of 0.05 (Fig. 2A). These 26,105 DMRs overlapped with the promoter regions of 19,809 genes, which were then used as the input for the enrichment analysis of disease-gene associations in the R package clusterProfiler [24]. The top 20 enriched disease terms are shown in Fig. 2A. These enriched terms included precancerous conditions and disseminated malignant neoplasm, which could be linked to the development of metastatic tumors. It was worth noting that cirrhosis, which was evidently diagnosed within patients with liver tumors [25], was also found among the enriched terms in our results. This was possibly contributed by the fact that the majority (64.7%, 11/17) of the unfavorable prognosis patients were diagnosed with LIM instead of LUM (Additional file 3: Table S1).

To further validate the prognostic power of these DMRs, we then performed the Jonckheere trend test using methylation value in the 59 primary tumor samples of different cancer stages as well as in the 11 tumor-adjacent samples from the 59 CRC patients. For the trend test, these 70 samples were clustered into 4 groups based on their TNM stage information (tumor adjacent, stage I, II and IV, respectively). We used 241,199 DMRs from the total 241,573 DMRs, which had methylation values available in all 70 samples (detail data not shown). For each DMR, a Jonckheere trend test was performed and 15,015 DMRs which were
showing statistical significance were selected for downstream analysis (Fig. 2B). We then selected 8,311 candidate DMRs, which were showing significant differences (q value ≤ 0.05, methylation differences ≥ 10%) between methylation values of the test group and the control group, from the total 241,573 DMRs for further analysis (Fig. 2B).

A final set of 213 DMRs (Additional file 3: Table S2), which met all three of the aforementioned filtering criteria, were selected as the prognostic DMR markers, as shown by the Venn diagram in Fig. 2. We then constructed a heatmap using the methylation values of these 213 DMRs in the 38 stage II primary tumor samples. As shown in Fig. 2C, there was noticeable differences between the group of unfavorable prognosis and favorable prognosis. These 213 DMRs between the 17 unfavorable prognosis and 21 favorable prognosis samples, as shown in Additional file 3: Table S2, included 182 hypermethylated and 31 hypomethylated DMRs. The overwhelming number of hypermethylated DMRs in our results was consistent with the current understanding that DNA hypermethylation associated with cancer are mostly found in gene regions, despite there are more hypomethylation compared to hypermethylation in general [26]. The targeted bisulfite sequencing was performed using SeqCap Epi CpGiant probes, which is biased toward finding hypermethylation as it focused more on genic regions compared to the whole genome bisulfite sequencing approach.

Additionally, the 213 prognostic DMRs have been filtered against gene region and are overlapping with promoter regions, which can enhance such bias. Figure 3 shows the comparison of methylation rate of nine randomly chosen prognostic DMRs between favorable and unfavorable prognosis groups. All of them were susceptible to be hypermethylated when metastasis was developed later on and mostly located in promoter regions of genes which had confirmed associations with certain type of cancer.

We then performed statistical analysis to investigate the potential impact of age and smoking history had on the 213 identified prognostic DMRs. The 38 stage II patients were first split into two different groups comparing their age toward the group median (< 60 years old and ≥ 60 years old, 16 and 22 patients, respectively). For each of the 213 DMRs, raw methylation rates for the 38 stage II patients were used for Wilcoxon test to compare if the two groups are significantly different (fdr adjusted p value or q value <0.1). None of the 213 DMRs showed any statistical significance differences between the two groups, as shown in Additional file 3: Table S3. Similar to the age test, these patients, excluding one patient without smoking history information, were subsequently grouped into smoking [11] and non-smoking [26] groups. The Wilcoxon tests showed that none of these DMRs had significant differences based on patients smoking history (q value <0.1, Additional file 3: Table S3).

**Construct a pilot model for predicting the liver metastasis based on DMRs in primary colorectal tumor**

Since LIM is the more dominant type of metastasis among CRC patients, we set to explore the possibility to predict LIM status using the primary tumor samples. The total 59 primary tumor samples were used to construct a pilot model for predicting the LIM status in these CRC patients. A total of 105 DMRs (methylation differences ≥ 20%, q value ≤ 0.05) were identified between the 22 patients with LIM and 37 patients without LIM during the follow-up. These DMRs were then used as candidates for the predictive model. By utilizing LOOCV and LASSO model as described in Fig. 4A, we were able to generate the probability scores for all 59 samples. A ROC curve was constructed, as shown in Fig. 4B, using these probability scores, yielding an Area Under Curve (AUC) score of 0.7015 (sensitivity = 72.7%, specificity = 70.3%, as shown in Additional file 3: Table S4). The model was then applied to the total 59 primary samples, and 23 out of the 105 DMRs were identified as optimal predictive markers (shown in Table 2). 7 out of 23 (30.4%) were overlapped with a promoter region and 6 (26.1%) were located within the coding region. The rest 7 DMRs were involved in the
59 enrolled patients
  11 stage I
  38 stage II (21 favourable vs 17 unfavourable)
  10 stage IV

81 total samples extracted
  11 tumor adjacent
  59 primary tumor (11 stage I, 38 stage II, 10 stage IV)
  11 metastatic tumor (6 LiM, 5 LUM)

Targeted bisulfite sequencing
  SeqCap Epi CpGiant panel, Roche
  2.7 x 10^6 CpG sites
  80.5Mb genomic region

Aim 1: Identify prognostic DMRs in stage II CRC
  38 stage II tumor samples

Identify DMRs using methylKit
  17 stage II unfavourable prognosis vs
  21 stage II favourable prognosis

241,573 DMRs

Excluded 233,262 DMRs
  q value > 0.05 or
  methylation difference < 10%

Excluded 6,357 DMRs
  Outside promoter region

Excluded 1,741 DMRs
  Jonckheere trend test insignificant (p value > 0.05)

213 prognostic DMRs

Aim 2: Evaluate the feasibility of predicting LiM using DMRs in primary CRC tumor samples
  59 primary tumor samples (All stages included)

Identify DMRs using methylKit
  22 primary tumor samples from LiM patients vs
  37 primary tumor samples from LiM-free patients

105 DMRs
  q value <= 0.05 and
  methylation differences >= 20%

LOOCV

Marker selection
  23 predictive DMRs

Final LiM risk model

Validation set
  6 LiM tumor samples
  5 LUM tumor samples
intergenic regions. Wilcoxon test results shown that none of these 23 DMRs were significantly different between different age/smoking group within these patients (q value < 0.1, Additional file 3: Table S5).

To explore the utility of the established model in discriminating metastatic sites, we generated another ROC curve using the 23 predictive markers on 11 metastatic tumor samples (6 LIM and 5 LUM). The model was showing an excellent performance in discriminating the LIM samples against the LUM samples (AUC = 0.9333, sensitivity = 83.3%, specificity = 100%, Fig. 4C). Furthermore, a Kaplan–Meier curve of LIM-free survival suggested that the patient predicted as LIM positive had a significantly shorter LIM-free survival compared to the LIM predicted negative (Log-rank p = 0.0037, Fig. 4D). We also constructed a Principal Component Analysis (PCA) with the total 81 samples including 11 tumor-adjacent samples, 59 primary tumor samples and 11 metastatic tumor samples. As illustrated by the PCA (Fig. 4E), the LIM positive and LIM negative groups in the primary tumor samples were separated from each other. Similarly, the LIM positive and negative samples from the metastatic tumor can be distinguished. The 11 tumor-adjacent samples were placed closer to the primary without LIM samples yet with trend of separation.

Finally, visible differences can be observed in the heatmap generated using these 23 DMRs in the 59 primary tumor samples (Fig. 4F). Finally, an external validation cohort, which consist of primary tumor samples from 8 CRC patients (4 LIM and 4 LIM-free), was used to further evaluate the model performance. As shown in Additional file 2: Figure S2, our model showed great performance in differentiating the LIM samples from the LIM-free samples (AUC = 0.875, sensitivity = 100%, specificity = 75.0%).

**Discussion**

Despite detail mechanisms not being completely understood, DNAm are believed to have the ability of altering downstream gene expression by affecting the binding of transcription factors and their target sites[27]. Additionally, evidence suggest that DNAm within the gene body, especially the first exon, can be associated with transcriptional silencing[28]. DNAm now is an emerging biomarker for cancer diagnosis and prognosis prediction which plays an important role in establishing epigenetic imprints.

Here we investigated the different DNAm status in CRC patients with and without metastasis within the follow-up period. Using targeted bisulfite sequencing,
we initially identified over 24 thousand of DMRs by comparing the primary tumor samples of 38 stage II patients. With several steps of data filtration, we eventually selected 213 candidate DMRs which might serve as metastasis predictors and the majority were hypermethylated in the unfavorable prognosis group. Evidence suggested that, while there are more cancer-related hypomethylation than hypermethylation in the intergenic region, DNA methylation associated with cancer found in genic regions were mostly hypermethylation [26, 29].

A recent study in 2018 by Hidaka et al. has focused on genes expression regulated by DMRs in 106 CRC patients and found an overall trend toward hypermethylation in CRC tissue samples [16]. This agrees with the fact that we had more hypermethylation in our results, which were all located within promoter regions further enhancing such bias.

Through literature searching, we found that many of them were associated with cancer-related genes. For example, our results suggested that the unfavorable prognosis group were hypermethylated compared to the favorable prognosis group within the region chr1:118147001–118148000. Such hypermethylation in the promoter region would in theory result in downregulation of the \textit{FAM46C} gene, which was acting as an onco-suppressor gene in multiple myeloma [30]. Similarly, our data suggested that the promoter regions of
SORCS3 – AS1 were differentially methylated among the two groups of different prognoses. Interestingly, Schneider et al. reported in 2015 that the methylation level of SORCS3 gene can be associated with tumor progression in gastric cancer [13]. The promoter region of HLA-DQA1 gene, which was part of the human leukocyte antigen (HLA) complex, was differentially methylated and labeled as a prognostic marker in our result. Intriguingly, the expression of HLAB gene, which was also part of the HLA complex, was reportedly associated with tumor progression in CRC [31]. Furthermore, ARHGEF1, CFAP65, PDGFRB and CLECSA were labeled as prognostic marker by the Human Protein Atlas in renal and breast cancer, endometrial cancer, renal and urothelial cancer, and ovarian cancer, respectively [32, 33].

The predictive model established through LASSO and LOOCV was based on 23 DMRs, most of them were located in the gene coding or promoter regions. The seven genes whose promoter contained these DMRs were either previously reported as prognostic markers or showed the predictive potential in multiple cancers. For example, PAX8 and GUF1 were reported as prognostic markers for endometrial cancer, renal cancer and thyroid cancer, respectively [33]. Both KLF4 and EVI2B were identified by the Human Protein Atlas as markers for renal cancer prognoses (favorable and unfavorable, respectively) [33]. Furthermore, TNNI2 was reported to have predictive power for metastatic tumor development in gastric cancer [34]. A recent study suggested that the centrosomal protein 112 (Cep112) was able to act as an oncogene by interacting with genomic instability inducing RNA [35]. Finally, the long non-coding RNA AC011298 was among the six identified prognostic markers identified in a bladder cancer study [36].

In clinical setting, the postoperative treatment-decision for early-stage CRC is challenging. The high odd of metastasis development especially to liver dramatically decreases the five-year survival rate. Therefore, early-prediction of liver metastasis could be powerful to improve prognosis for early-stage CRC patients. Many biomarkers have been investigated for the possibility of predicting metastasis such as microRNAs [37] and specific gene expression level [38]. Nowadays, epigenetic information has drawn a broad attention as predictive biomarkers and DNA methylation status is the most investigated. Previous studies have identified hundreds of DMRs by
comparing different conditions such as stage, prognosis, and histology. However, no solid predictive model has been established. Herein, we sought to explore the possibility of predicting liver metastasis based on the primer tumor DNAm profiles. Taking advantages of the surgery resected tissue biopsy to predict the possibility of metastasis could reduce overtreatment and provide valuable information to guide treatment. We explored the DNAm profiles of the primary tumors to characterize novel DMRs features by comparing favorable and unfavorable stage II CRC patients. By identifying potential DMR markers which could reflect the risk of liver metastasis, we aimed to eventually establish a model to predict the metastasis risk by detecting the primary tumor DMRs. However, due to the restricted cohort size, we could only perform the LOOCV for model selection and validated the predictive model based on the metastatic tumor samples and a small external validation cohort. A larger cohort with methylation values in primary CRC samples would be a great value for DMRs identification and modelling which remained to be completed in the future. Therefore, in the present study, we have identified DNAm biomarkers associated with the risk of cancer liver metastasis in early-stage CRC patients and proposed a pilot risk prediction model based on those epigenetic markers for outcome assessment.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s13148-021-01108-3.

**Additional file 1: Figure S1**. Liver metastasis free survival for the 59 CRC patients. 60% of the 59 CRC patients reached a stable stage of liver metastasis free at 1600 days.

**Additional file 2: Figure S2**. Receiver operating characteristic curve in an external validation cohort. Performance of the 23 predictive DMR markers were tested using 8 primary CRC tumor samples (4 LIM, 4 LIM-free).

**Additional file 3: Table S1**. Clinical characteristics of the 81 samples from 59 CRC patients included in this study.

**Table 2** Annotation of the 23 predictive DMRs identified using 59 primary CRC tumor samples

| Genome position | Gene context | Gene name | Type of DMR |
|-----------------|--------------|-----------|-------------|
| chr1:1861001–1862000 | Promoter | TNN2 | hypo |
| chr2:113994001–113995000 | Promoter | PAX8 | hypo |
| chr2:241626001–241627000 | Promoter | AC011298 | hypo |
| chr4:44679001–44680000 | Promoter | GUF1 | hyper |
| chr9:110248001–110249000 | Promoter | KLF4 | hyper |
| chr17:29641001–29642000 | Promoter | EVI2B | hyper |
| chr17:63739001–63740000 | Promoter | CEP112 | hypo |
| chr12:200977001–200978000 | Gene body | KIF21B | hyper |
| chr12:62765001–62766000 | Gene body | GRIP1 | hyper |
| chr13:11129001–11129000 | Gene body | NAXD | hypo |
| chr19:39314001–39315000 | Gene body | ECH1 | hypo |
| chr2:136279001–136280000 | Gene body | ZRNAB3 | hyper |
| chr7:51148001–51149000 | Gene body | COBL | hypo |
| chr8:72468001–72469000 | Gene body | RP11-1102P16.1 | hyper |
| chr8:99394001–99395000 | Gene body | KB-1458E12.1 | hypo |
| chr16:13440001–13450000 | Gene body | RP11-616M22.7 | hypo |
| chr11:12097001–12098000 | Intergenic | N/A | hyper |
| chr12:9007001–9008000 | Intergenic | N/A | hyper |
| chr2:75504001–75505000 | Intergenic | N/A | hyper |
| chr22:18530001–18531000 | Intergenic | N/A | hyper |
| chr4:153039001–153040000 | Intergenic | N/A | hypo |
| chr7:34344001–34345000 | Intergenic | N/A | hypo |
| chr9:139591001–139592000 | Intergenic | N/A | hypo |

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**Authors’ contributions**

YS and JY conceptualized the study. WL and LG performed the experiments analyzed the data. HZ collected the clinical samples. WT performed the bioinformatics analysis. YM and XW revised the manuscript. All authors approved the final version of manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
All study protocols were approved by the ethics committee of the National Cancer Center/National Clinical Research Center for Cancer/Cancer Hospital, and in accordance with international standards of good clinical practice. Written informed consents were provided by all patients.

Consent for publication
The content of this manuscript has not been previously published and is not under consideration for publication elsewhere.

Competing interests
Wanxiangfu Tang, Yutong Ma, Xiaonan Wang, and Yang Shao are employees of Nanjing Geneseeq Technology Inc., Nanjing, Jiangsu, China. The remaining authors have nothing to declare.

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