DNA methylation-based signature of CD8+ tumor-infiltrating lymphocytes enables evaluation of immune response and prognosis in colorectal cancer

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ABSTRACT

Background Tumor-infiltrating lymphocytes (TILs), especially CD8+ TILs, can be used for predicting immunotherapy responsiveness and survival outcome. However, the evaluation of CD8 + TILs currently relies on histopathological methodology with high variability. We therefore aimed to develop a DNA methylation signature for CD8 + TILs (CD8 + MeTIL) that could evaluate immune response and prognosis in colorectal cancer (CRC).

Methods A CD8 + MeTIL signature score was constructed by using CD8 + T cell-specific differentially methylated positions (DMPs) that were identified from Illumina EPIC methylation arrays. Immune cells, colon epithelial cells, and two CRC cohorts (n=282 and 335) were used to develop a PCR-based assay for quantitative analysis of DNA methylation at single-base resolution (QASM) to determine CD8 + MeTIL signature score.

Results Three CD8 + T cell-specific DMPs were identified to construct the CD8 + MeTIL signature score, which showed a dramatic discriminability between CD8 + T cells and other cells. The QASM assay we developed for CD8 + MeTIL markers could measure CD8 + TILs distributions in a fully quantitative, accurate, and simple manner. The CD8 + MeTIL score determined by QASM assay showed a strong association with histopathology-based CD8 + TIL counts and a gene expression-based immune marker. Furthermore, the low CD8 + MeTIL score (enriched CD8 + TILs) was associated with MSI-H tumors and predicted better survival in CRC cohorts.

Conclusions This study developed a quantitative DNA methylation-based signature that was reliable to evaluate CD8 + TILs and prognosis in CRC. This approach has the potential to be a tool for investigations on CD8 + TILs and a biomarker for therapeutic approaches, including immunotherapy.

INTRODUCTION

Colorectal cancer (CRC) is one of the leading causes of cancer-related death worldwide.1,2 Immunotherapy is playing an increasingly important role in treating CRC in both resectable and non-resectable patients.3 Immune checkpoint inhibitors can boost T cells to attack tumor cells and prolong patient survival.4-6 Meanwhile, immunotherapy for CRC remains challenging because of its heterogeneous immune response.7-8 Tumor-infiltrating lymphocytes (TILs) are associated with tumor immune response and can be applied to predict the response to immunotherapy and survival outcome.9-11 Among them, CD8 + TILs kill cancer cells in a cytotoxic manner and are recognized to affect prognosis, immunotherapy response, and survival outcome in CRC.12-14

Multiple morphology-based methods have been established and widely validated to evaluate TILs in a reproducible way, including the recommendations by an international TILs working group15 and an immunoscore assay.16 In addition, some gene expression-based immune markers were developed to be an adjunct method to assess CD8+ TILs17-19 including the CD8B gene expression marker panel. Moreover, some DNA methylation-based markers for TIL (MeTIL) were recently identified from epigenome-wide analyses in breast cancer, showing good performance in the evaluation of the local tumor immune response and potential to improve the prognostic accuracy in patients with multiple cancer.11

DNA methylation is tissue-specific and plays a decisive role in tissue differentiation.20-21 Thus, DNA methylation profiling varies in cells from different tissues and organs, and it has the potential to be a marker to recognize different cells.22 Among them, the methylation of CpG shores within intermediate-CpG density regions is highly conserved and can discriminate tissue types regardless of species of origin.23 In addition, dramatic hypomethylation of partially methylated domains (PMDs) occurs during lymphocyte activation.
and proliferation, which can be predicted by solo-WCGWs (CpGs flanked by A or T in PMD context) that are mostly located in low-CpG density regions. Unfortunately, these areas of DNA methylation that have the potential to distinguish immune cells from other cells are characterized by low CpG density, which technically hinders the quantitative PCR (qPCR)-based assays in determining the methylation in these regions. To achieve a robust single-base specificity using PCR, we developed a novel assay to best use these low CpG-density methylations for developing biomarkers.

In this study, we aimed to analyze the genome-wide DNA methylation profile of immune cells and colon epithelial cells to identify CD8 + T cell-specific differentially methylated positions (DMPs). Then, we constructed a CD8 + McTIL signature score by using these DMPs to assess CD8 + TILs and survival outcomes in CRC and other cancers. Finally, we developed a qPCR-based assay to determine this score by using low amounts of DNA from formalin-fixed, paraffin-embedded (FFPE) tumor tissue. We demonstrated the utility of this methodology in CD8 + TIL evaluation.

MATERIALS AND METHODS

Study design

This study aimed to identify DNA methylation-based biomarkers that can assess CD8 + TILs in CRC. First, we identified CD8 + T cell-specific DMPs from the discovery set composed of CD8 + T cells, other immune cells, and cancerous and normal colorectal epithelial cells with EPIC methylation array datasets. We then selected DMPs and constructed a CD8 + McTIL signature score model with a machine learning algorithm. We further developed a QASM assay to test selected DMPs in a qPCR-based manner at single-base resolution. Finally, we tested the selected DMPs in multiple cohorts with this method, compared them with CD8B gene expression assay and immunohistochemical (IHC) assay for CD8 + TILs counts, and investigated their prognostic value in CRC and other cancers.

Patient cohorts and tumor samples

Two cohorts with CRC tumor samples from Colon Cancer Family Registry (CCFR) study and Sixth Affiliated Hospital of Sun Yat-sen University (SAH-SYSU) tumor bank were applied in this study. A description of the populations in CCFR study and SAH-SYSU samples has been published previously. Briefly, the CCFR study cohort included 335 patients with pathologically confirmed CRC at Seattle, Ontario, and Mayo CCFR sites, and the SAH-SYSU cohort consisted of 282 patients with pathologically confirmed CRC at SAH-SYSU from 2008 to 2012. All cases underwent radical-intent resection of the primary tumor. Patients with any prior anticancer treatment, concurrent cancer other than CRC, chronic inflammatory bowel disease, familial adenomatous polyposis, and whose DNA sample was degraded or not available were excluded. See online supplemental tables for the baseline characteristics of these cohorts. All samples used in this study were reviewed by pathologists to confirm the diagnosis and ensure that the tumor samples had more than 60% tumor content. To externally validate the findings in other cancer types, we used The Cancer Genome Atlas (TCGA) cohorts with Infinium Human Methylation 450K BeadChip (450K methylation array) profile. The workflow of cohort disposition is summarized in figure 1.

Colon epithelial cells and immune cells

All cancerous and normal colon cell lines (RKO, SW48, HCT8, HCT115, and NCM460) were obtained from American Type Culture Collection. Human-derived immune cells, including CD4+, CD8+, monocytes, and natural killer (NK) and B cells (1505, 1508–1510, 1513; LDE Biotechnology Co., Guangzhou) were enriched by magnetic-activated cell sorting.

Screen of CD8+ T cell-specific DMPs in CRC

A discovery set (n=167) composed of CD8 + T cells, non-CD8 + immune cells, cancerous and normal colorectal epithelial cells with Illumina Infinium MethylationEPIC Beadchip (EPIC methylation array) methylation profile data was obtained from Gene Expression Omnibus datasets with identifiers: GSE103541, GSE118144, GSE118972, GSE111681, and GSE122126. The characteristics of these samples are summarized in the online supplemental tables. The ‘minfi’ R package and Combat tool were used to perform raw data processing, batch effect correction, and DMP analysis as previously described. To identify CD8 + T cell-specific DMPs that can distinguish CD8 + T cells from non-CD8 + immune cells and cancerous and normal colorectal epithelial cells, we performed DMP analysis using their methylation data and selected the top-ranked candidate.

Nucleic acid isolation and bisulfite conversion

Genomic DNA was extracted from 20 mg fresh-frozen tissues using QIAamp DNA Mini Kit (Qiagen, 51306) and treated with sodium bisulfite using EZ DNA Methylation Kit (Zymo, D5002). RNA was extracted from 20 mg fresh-frozen tissues using Rneasy Plus Mini Kit (QIAGEN, 74134).

EPIC methylation array and bisulfite pyrosequencing

Evaluation of genome-wide DNA methylation was performed in 45 fresh-frozen tumor samples from the SAH-SYSU cohort using EPIC methylation array following the manufacturer’s protocol. The raw data were processed with the method described in the screen of CD8 + T cell-specific DMPs. We also evaluated the methylation of selected CpGs in this cohort by bisulfite pyrosequencing as described in the online supplemental materials.

qPCR-based quantitative analysis for single-base methylation (QASM)

The methylation percentage of each candidate CpG site was determined in multiple cohorts using a
MethyLight\textsuperscript{37}-based QASM assay that has been developed and validated in our previous work.\textsuperscript{26} In short, the bisulfite-converted DNA was amplified, in which we exploited the locus-specific PCR primers flanking a pair of methylated and unmethylated probes labeled with the fluorescent dyes 6-carboxyfluorescein (6-FAM) and 2-chloro-7phenyl-1,4-dichloro-6-carboxyfluorescein (VIC), respectively. The methylation percentage was calculated by methylation/(methylation + unmethylation)×100%. QASM was performed using the Applied Biosystems QuantStudio 7 Flex Real-Time PCR System (Thermo). The detailed procedure of QASM assay for CD8 + MeTIL markers can be found in the online supplemental materials.

**IHC assessment of CD8+ TILs (CD8+ PaTIL)**

The FFPE tumor blocks were cut into 4 µm sections. The deparaffinized slides were incubated with monoclonal anti-CD8 antibody (clone C8/144B; DAKO, Kyoto, Japan, SK201) at 1:100 dilution for 16 hours at 4°C. Subsequently, the secondary antibody conjugated with horseradish peroxidase (Envision + Dual Link Kit, DAKO, K5007) was applied. IHC assessment of CD8 + TILs (CD8 + PaTIL) was performed using the Envision + Dual Link Kit.
was performed following the method described by Fortis et al. Briefly, visual enumeration in five representative fields (0.1255 mm² per field) was performed by two independent pathologists, and the mean counts of CD8 + TILs were recorded.

**CDBB gene expression assay for CD8+ TILs (CD8+ ExTIL)**

Total RNA extracted from fresh-frozen tumors was reversely transcribed by ReverTra Ace qPCR RT Master Mix (TOYOBO, FSQ-201) according to the manufacturer’s protocol. Then, CD8B mRNA expression was quantified by qPCR assay. The primer sequence can be found in the online supplemental tables.

**Molecular phenotyping**

Tumor characteristics of microsatellite status, somatic mutations in KRAS codons 12 and 13, BRAF p.V600E mutation, and CpG island methylator phenotype (CIMP) in each cohort were determined as previously described.

**Statistical analysis**

Differences between groups were evaluated using Student’s t-test, Mann-Whitney test or χ² test. Consistencies between results were analyzed using Pearson’s correlation analysis. The maximally selected rank statistics were applied to determine the optimal cut-off of variables in the survival analysis. Cox regression analysis and Kaplan-Meier survival curves with log-rank tests were used to compare overall survival (OS). Gene Ontology (GO) enrichment analysis was performed using GOfuncR package. All the statistical analyses were conducted with R V.3.6.1. All p values were calculated with two-tailed tests and considered significant when p<0.05.

**RESULTS**

**DNA methylation signature of CD8+ TILs in colorectal tumors**

The outline of the study is shown in figure 1. To develop the CD8 + MeTIL signature, we first compared the methylation signature of CD8 + T cells with those of cancerous and normal colorectal epithelial cells and other human-derived immune cells, including CD4 + T cells, B cells, granulocytes, and monocytes, using their genomic methylation data profiled by EPIC array (see online supplemental materials for details). We identified 73 CD8 + T cell-specific DMPs that were highly differentially methylated between CD8 + T cells and other cells in this discovery step (figure 2A and online supplemental material 1).

To confirm the results in the discovery step, we analyzed the biological features of these CD8 + T cell-specific DMPs. Most CpGs were located in the intragenic region (80.6%, 59 probes), including CD8A, CD8B, CD96, and other functional genes in CD8 + T cell (online supplemental material 1), and a few ones were intergenic CpGs (19.2%, 14 probes). We further performed GO analysis for intragenic CpGs and found multiple immune-associated GO terms were top-ranked (figure 2C). This finding confirmed the strong association of identified CpGs with CD8 + T cells.

Next, we analyzed the genomic features of these 73 CpGs. The majority were located in low-CpG density (60.3%, 44 probes) and intermediate-CpG density (34.2%, 25 probes) regions, and few CpGs (5.5%, 4 probes) were located in the high-CpG density region (CpG island) (figure 2B). Of note, 6 CpGs were solo-WCGW CpGs, and 10 CpGs were located in common PMD, according to the previous genomic annotations.

To achieve a shrinking panel that can be easily accessible, we applied the least absolute shrinkage and selection operator (LASSO) machine learning approach to select CpGs that can best represent our list of 73 CD8 + T cell-specific CpGs (online supplemental material 1). The final signature, named CD8 + MeTIL score, included three CpGs (cg02430840, cg06113913, and cg12673499) located in the low-CpG density region. We constructed a CD8 + MeTIL score with their β values and coefficients in LASSO regression: cg02430840 ×0.28724+cg06113913 ×0.40171+cg12673499×0.45016. This score can theoretically adapt values between 0.00 and 1.14, while we received CD8 + MeTIL scores between 0.71 and 1.00 in the discovery cohort.

To explore the distinguishability of the CD8 + MeTIL score, we grouped cells by unsupervised hierarchical cluster analysis using the three CpGs. We found that CD8 + T cells could be well distinguished from other cell types by assessing these three CpGs (figure 2D). Of note, the CD8 + MeTIL score was dramatically lower in CD8 + T cells than that in any of cancerous and normal colorectal epithelial cells, CD4 + T cells, B cells, granulocytes and monocytes (all β >0.4, all β fold change >3.0, all p<0.001; figure 2E). This high specificity of CD8 + MeTIL score in CD8 + T cells was the basis of its ability to determine the abundance of CD8 + T cells in a tumor block. Furthermore, we validated the specificity of CD8 + MeTIL score in an independent sample set with DNA methylation profile of immune cells, cancerous and normal colon epithelial cells, and other cell types that were recognized in the tumor microenvironment (online supplemental material 1).

We next sought to investigate the association between the methylation profiles of the three selected CpGs in the CD8 + MeTIL score and clinicopathological features in the SAH-SYSU cohort of 45 CRC cases with EPIC array data (figure 2F). We found that a lower CD8 + MeTIL score with less methylation in the three CpGs was associated with CRCs with low-frequency microsatellite instability (MSI-H), right-sided location, and high differentiation. These results were consistent with the findings from previous IHC-based CD8 + TIL association studies. The methylation levels of these three CpGs determined by EPIC array were significantly correlated with their methylation levels determined by bisulfite pyrosequencing (figure 3D; r=0.7357–0.9025, all p<0.01).
**Figure 2** Generation of CD8+ MeTIL score using CD8+ T cell-specific DMPS. (A) Unsupervised hierarchical clustering analysis of CD8+ T cells versus other human-derived immune cells, and normal and cancerous colon cells based on β values of 73 candidate DMPs. (B) Statistics of genomic location and motif for the 73 candidate DMPs. (C) Gene Ontology analysis of the genes that 73 candidate DMPs were located in. The three categorized terms including molecular function, biological process, and cell composition were analyzed. (D) The finally selected CpGs in the LASSO model were tested for its representative ability based on the unsupervised hierarchical cluster analysis. Each column represents one sample; red or yellow indicates high methylation levels, while blue indicates low levels. (E) The methylation values of three selected CpGs were used to construct a CD8+ MeTIL score, and the score was calculated for CD8+ T cells, CRC cells, normal colon cells, B cells, CD4+ T cells, granulocytes, and monocytes. Differences in CD8+ MeTIL score between CD8+ T cells and other cell types were assessed with Student's t-test (**p<0.0001). (F) Unsupervised hierarchical clustering analysis in 45 samples from SAH-SYSU cohort using the β values of three selected CpGs. The clinicopathological features of each patient were displayed above the heatmap. BMI, body mass index; CIMP, CpG island methylator phenotype; CRC, colorectal cancer; DMP, differentially methylated position; MHC, major histocompatibility complex; PMD, partially methylated domain; SAH-SYSU, Sixth Affiliated Hospital of Sun Yat-sen University.
Figure 3  Quantitative PCR-based QASM assay for determining CD8+ MeTIL score at single-base resolution. (A) The design of QASM assay for methylation detection of each isolated CpG. Each methylation level was determined merely at probe level, in which a FAM-labeled MGB probe specifically binds to methylated allele (CpG) sequence, and the VIC-labeled probe binds to the unmethylated allele (TpG) sequence. Both alleles were amplified with one pair of primers in the same PCR reaction. (B–D) Scatter plots showing the correlations between any two of the QASM assay (n=45), EPIC methylation array (n=45) and bisulfite pyrosequencing (n=12) in determining methylation percentage of each CpG targeted by cg02430840, cg06113913, and cg12673499, respectively. Pearson’s test was applied and correlation coefficient (r) and p value for each procedure are shown. MGB, minor groove binder.
QPCR-based assay for CD8+ MeTIL score at single-base resolution

To easily obtain CD8 + MeTIL score generated from array-based analysis, we developed a qPCR-based assay (QASM) to determine the methylation of three selected CpGs that were located in the low-CpG density region at single-base resolution. As figure 3A displayed, the methylated and unmethylated alleles of each CpG were quantified at the probe level, in which methylation percentages can be determined by the signal ratio of two probes.

To validate this qPCR-based assay for determining CD8 + MeTIL score, we next compared the results of QASM assay with those of the EPIC array and pyrosequencing in the SAH-SYSU cohort. The qPCR-based and EPIC array-based methylation percentages were linearly related well in each CpG (figure 3B; n=45, r=0.5225–0.7274, all p<0.001). In addition, the methylation percentages determined by QASM assay linearly correlated with those determined by pyrosequencing (figure 3C; n=12, r=0.7170–0.9411, all p<0.01). The CD8 + MeTIL scores generated from QASM assay linearly correlated with the scores generated from both the EPIC array and pyrosequencing (figure 3; n=45 and 12, respectively; r=0.7357 and 0.9025, respectively; all p<0.001). Thus, QASM assay is a technically reliable tool to determine CD8 + MeTIL score.

To measure CD8 + TIL distribution with CD8 + MeTIL score, we first used QASM to determine the methylation percentages of three CpGs in different tissues and cell types, including CRC tissue (n=45) and adjacent normal tissue (n=44), normal (NCM460) and cancerous (RKO, SW48, HCT8, and HCT15) colon epithelial cell lines and multiple immune cells (CD4 + T cells, monocytes, and B cells), which represented the major components of CRC tissues. The CD8 + T cells displayed a significantly lower CD8 + MeTIL score compared with other cells using the QASM assay (all score fold change >3.0, all △ score >2.3, all p<0.001; figure 4A) and were consistent with the results determined from the EPIC array dataset, further supporting the qPCR-based CD8 + MeTIL score as an accurate surrogate for the array-based score to determine the abundance of CD8 + T cells in tumor samples.

Evaluation of CD8+ TIL distribution in CRC with CD8+ MeTIL score

We applied this qPCR-based CD8 + MeTIL score to assess CD8 + TILs in tumor tissues and compared it with CD8 + TILs determined by IHC staining (CD8 + PaTILs) in SAH-SYSU cohorts (figure 4B,C). The CD8 + PaTILs displayed a significant negative correlation with the CD8 + MeTIL score (r=−0.6385, p=0.011). Notably, CD8 + MeTIL low group had significantly more CD8 + TIL counts in IHC staining than the CD8 + MeTIL high group did (p=0.005, figure 4C).

We further compared CD8 + MeTIL score with the CD8 + expression-based TIL (CD8 + ExTIL) panel, which is a molecular signature of CD8B gene expression generated to assess CD8+ MeTIL as previously used to train other panels for evaluation of CD8+ TILs.19 We determined the mRNA expression of CD8B gene expression, encoding a part of the CD8 antigen, in SAH-SYSU cohort. Expectedly, a significant negative correlation was observed between CD8+ MeTIL score and CD8B gene expression signature (r=−0.4079, p=0.008; figure 4D). Taken together, lower CD8+ MeTIL scores determined by QASM assay were associated with higher mean density of CD8+ T cells (CD8+ PaTILs) and increased expression of CD8B gene (CD8+ ExTIL) in CRC tissue.

Next, we evaluated the distribution of CD8+ MeTIL score determined by QASM assay by molecular features of CRC cases from the CCFR cohort. The CD8+ MeTIL score of the MSI-H group was significantly lower compared with those of the low-frequency microsatellite instability (MSI-L)/ microsatellite stability (MSS) group (77.8 vs 82.1, p<0.001; figure 4E), which is consistent with previous findings that MSI-H tumors harbored more CD8+ TILs than MSI-L/MSS tumors did.12 Moreover, patients with CIMP-positive (78.5 vs 81.6, p=0.048) and right-side (78.9 vs 82.1, p=0.001) tumors had significantly lower CD8+ MeTIL score compared with CIMP-negative and left-side tumors, respectively (figure 4E).

Prognostic value of CD8+ MeTIL in CRC and multiple cancers

Considering the role of CD8+ T-cell infiltration in tumor immune response, we sought to test the role of CD8+ MeTIL score in prognosis prediction for CRC. We used QASM assay to determine the CD8+ MeTIL scores in CRCs from the SAH-SYSU (n=237) and CCFR (n=335) cohorts, and grouped patients by the cut-off determined by maximally selected rank statistics (online supplemental material 1). The baseline characteristics of these grouped patients in multiple cohorts are shown in the online supplemental materials. The patients with low CD8+ MeTIL scores had significantly better OS compared with those with high CD8+ MeTIL scores in both SAH-SYSU (figure 5A; HR=0.303, 95% CI 0.109 to 0.842, p=0.022) and CCFR (figure 5C; HR=3.17, 95% CI 1.26 to 7.99, p=0.009) cohorts. This result was further confirmed after adjusted by other significant risk factors, including age, TNM stage, KRAS mutation, and lymphovascular invasion, in multivariate Cox analysis (online supplemental material 1).

Next, we explored whether CD8+ MeTIL score could further stratify MSI-H/MSI-L/MSS patients in marker prediction assays. In survival analysis, patients were grouped according to MS status and CD8+ MeTIL score in SAH-SYSU and CCFR cohort. As expected, the death risk could be further stratified by CD8+ MeTIL score in each microsatellite subgroup (figure 5B,D). Of note, patients with MSI-H and abundant CD8+ TILs (low CD8+ MeTIL score) had the best OS in both cohorts, while MSI-L/MSS and low levels of CD8+ TILs (high CD8+ MeTIL score) predicted poor OS in both cohorts. Though this score was developed in CRC, we found the lower CD8+ MeTIL score was significantly associated with anti-PD-1 treatment response and survival benefit in non-small cell lung.
Figure 4  Evaluation of CD8 + TILs in cell lines and tumor samples using the CD8 + MeTIL score. (A) The CD8 + MeTIL score and methylation level of each CpG determined by QASM assay showed significant differences between CD8 + T cells and other immune cells, normal and cancerous colon cells. (B) Scatter plots showing the correlation between CD8 + MeTIL score determined by QASM assay and CD8 + TIL counts determined by IHC staining with FFPE slides using 23 samples from SAH-SYSU cohort. Pearson’s correlation coefficient and p value are shown. (C) Representative images of IHC stained slides with CD8 + TIL counts from low to high (from left to right) at ×400 and ×800 magnification. Both the CD8 + TIL counts and CD8 + MeTIL score are shown, and a darker color means a higher value. (D) Scatter plots showing the correlation between CD8 + MeTIL determined by QASM assay and CD8 + ExTILs determined by CD8B gene mRNA expression assay using 43 samples from the SAH-SYSU cohort. Pearson’s correlation coefficient and p value are shown. (E) Distribution of CD8 + MeTIL by different molecular phenotypes in CCFR study cohort. The Mann-Whitney test was applied (**p<0.001, *p<0.05). Horizontal bars, median ±IQR. CCFR, Colon Cancer Family Registry; FFPE, formalin-fixed, paraffin-embedded; SAH-SYSU, Sixth Affiliated Hospital of Sun Yat-sen University; TIL, tumor-infiltrating lymphocyte.
Figure 5  The prognostic value of CD8+ MeTIL in CRC and multiple cancers. Patients were grouped by CD8+ MeTIL score in SAH-SYSU cohort (A, n=237) and CCFR study cohort (C, n=335) based on the cut-off determined by maximally selected rank statistics. Kaplan-Meier curves of OS in each cohort with p values from log-rank test are shown. Patients with MSI-H and MSI-L/MSS tumors were further stratified by CD8+ MeTIL score, and four groups were generated and compared in the Kaplan-Meier curves of OS in the SAH-SYSU cohort (B) and the CCFR study cohort (D) with log-rank test. (E) Workflow of DMPS filtering to generate 450K-specific CD8+ MeTIL signature that could be applied in TCGA cohorts with 450K methylation array data. (F) The forest plot showed the HRs and 95% CIs of the 450K-specific CD8+ MeTIL signature in univariate Cox analysis for survival prediction in 24 TCGA cancer types. ACC, adrenocortical carcinoma; BLCA, bladder urothelial carcinoma; BRCA, breast invasive carcinoma; CESC, cervical squamous cell carcinoma and endocervical adenocarcinoma; COREAD, colon and rectum adenocarcinoma; CRC, colorectal cancer; DMP, differentially methylated position; ESCA, esophageal carcinoma; GBM, glioblastoma multiforme; HNSC, head and neck squamous cell carcinoma; KICH, kidney chromophobe; KIRC, kidney renal clear cell carcinoma; KIRP, kidney renal papillary cell carcinoma; LIHC, liver hepatocellular carcinoma; LUAD, lung adenocarcinoma; LUSC, lung squamous cell carcinoma; meso, mesothelioma; OS, overall survival; PAAD, pancreatic adenocarcinoma; PCPG, pheochromocytoma and paraganglioma; sarc, sarcoma; STAD, stomach adenocarcinoma; SAH-SYSU, Sixth Affiliated Hospital of Sun Yat-sen University; THCA, thyroid carcinoma; UCEC, uterine corpus endometrial carcinoma; UCU, uterine carcinosarcoma; UVM, uveal melanoma.
cancer using the EPIC methylation array data (online supplemental material 1).

Acknowledging that the CD8+ MeTIL signature was developed in CRC using EPIC methylation array data, we performed survival analysis in other cancer types using 450K methylation array data available in TCGA. In 6 of 24 tested cancer types (lung adenocarcinoma (LUAD, p=0.033), lung squamous cell carcinoma (LUSC, p=0.038), CRC (colon and rectum adenocarcinoma, p=0.048), cervical squamous cell carcinoma (CESC, p=0.044), breast invasive carcinoma (BRCA, p=0.011), and adrenocortical carcinoma (ACC, p=0.033)), low CD8+ MeTIL scores were significantly associated with a better survival outcome (figure 5F). Taken together, these data suggest that the CD8+ MeTIL score may predict survival outcomes in multiple cancers.

DISCUSSION

In this study, we analyzed the genome-wide DNA methylation profile to identify CD8+ T cell-specific DMPs and constructed a CD8+ MeTIL score to assess CD8+ TILs in CRC. We demonstrated the reliability of CD8+ MeTIL signature in evaluating CD8+ TIL-based immune response by showing the tight association of this signature with IHC-based CD8+ PaTILs and CD8+ ExTIL signature. In addition, we developed a qPCR-based QASM assay that could effectively determine the CD8+ MeTIL score at single-base resolution, enabling the evaluation of CD8+ TILs in a quantitative approach. Finally, we validated the ability of CD8+ MeTIL to stratify microsatellite instability (MSI)/MSS status and prognostic outcomes in two CRC cohorts and identified associations with improved survival for several cancers from a pan-cancer analysis of the TCGA tumors.

The robustness of this study came from our presumption that cancer cells are mixed with normal epithelial cells and multiple subtypes of immune cells in each tumor block. We therefore compared the methylation profiles of CD8+ T cells with those of other potentially mixed cells and obtained 73 CD8+ T cell-specific DMPs. The genomic location of these 73 CD8+ T cell-specific DMPs showed that few CpGs were located in the high-CpG density region (CpG island), while most CpGs were located in low-CpG density regions. These regions are covered more effectively by the additional probes present in EPIC methylation array compared with the 450 k array. Our strict selection criteria that only EPIC array datasets could be included for pooled methylation data of cell lines from published resources were an advantage of this study. Moreover, the CD8+ T cell-specific DMPs are available in DNA methylation array, which facilitates the possibility of analyzing CD8+ TILs using DNA methylation array datasets and uncovering the association with the other omics profiles.

The reliability of applying CD8+ MeTIL signature in CD8+ TILs assessment came from our cross-validation in multiple analyses. First, the GO analysis on the genes that 73 CD8+ T cell-specific DMPs were located in showed that the methylation of these CD8+ T cell-specific CpGs may participate in antitumor immunity through T-cell activation, immune effector process, immune response to tumor cell and cell killing. Therefore, we speculate that these DMPs are not merely markers for CD8+ T cell, but they may also play a role in regulating T-cell function in tumor immune response that needs to be validated in future studies. Second, we found the CD8+ MeTIL was negatively associated with T-cell exhaustion and senescence markers (online supplemental material 1), which supports the previous findings that TILs are functionally impaired with expression of exhaustion and senescence markers in tumor microenvironment. Third, the analysis of clinical and molecular features showed that low CD8+ MeTIL score was related to tumor development in the right-side colon, high differentiation, MSI-H status and wild-TP53. These clinicopathological features have been previously shown to be associated with non-methylation-based measures of CD8+ TILs. Finally, we compared the IHC-based CD8+ PaTILs and CD8+ ExTIL results with the CD8+ MeTIL score in CRC cohorts, where both showed a significant negative correlation. This demonstrated that the signature score could be an alternative and reliable method to assess CD8+ TILs in CRC.

The current qPCR-based methylation assays have been mostly applied to determine the mean value of multiple CpGs in high-CpG density/intermediate-CpG density regions. Unfortunately, the DMPs included in the CD8+ MeTIL score were isolated CpGs within the low-CpG density region. Although the EPIC methylation array and bisulfite pyrosequencing can assay these isolated CpGs at single-base resolution, both of them are more expensive and time-consuming than measurement by a qPCR-based methylation assay. This may explain why the isolated CpGs within the low-CpG density region identified as methylation biomarkers in previous studies were limited to an array-analysis level and could not be extended to further study and clinical use. Thus, we developed a novel qPCR-based QASM assay that has been shown validity in determining single-CpG methylation by an independent group. In the current study, we also showed its validity in measuring the methylation level of each CpG included in the CD8+ MeTIL score. Compared with EPIC methylation array and pyrosequencing, the QASM assay is reliable to determine the CD8+ MeTIL score. This qPCR-based method measures CD8+ TILs in an unbiased, cheap and simple manner and thus is less susceptible to variability across different cohorts. The discrepancy in CD8+ TIL evaluation between the CD8+ PaTIL and CD8+ MeTIL in the scatter plots may be partly explained by the differences in the characteristics of two methods. Considering the methylation biomarkers of cancers are widely used.
in past decades,\textsuperscript{55, 56} we anticipated that our qPCR-based assay for CD8+ MeTIL signature, as measured by the QASM assay, could have been assessed in these tumors. Of note, we demonstrated the lower CD8+ MeTIL signature could predict a response to immunotherapy in non-small cell lung cancer. These results suggested that the CD8+ MeTIL signature may also assess immune responses in other cancers and thus stratify patients for prognosis and potentially for response to immunotherapy. Considering that the CD8+ MeTIL score was generated from CRC, whether the result of this study could be extended to other tumor types might need further study to confirm.

**CONCLUSIONS**

In this study, we developed a DNA methylation-based CD8+ MeTIL signature that was reliable to assess CD8+ TIL-based immune response in CRC. We developed a qPCR-based QASM assay to determine the CD8+ MeTIL signature at single-base resolution and that could evaluate CD8+ TILs in a quantitative and more high-throughput way. We further validated the ability of this signature to stratify MSI/MSS status and prognostic outcomes in two CRC cohorts and TCGA cohorts with other cancers. The CD8+ MeTIL signature, if further validated, has the potential to be a useful biomarker that could be readily deployed in clinical diagnosis to inform the choice of multiple therapeutic approaches, including immunotherapy.

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**Contributors**

Conception, design and supervision of this study; methylation array data processing and bioinformatic analysis; full access to all data for this study and take responsibility for the integrity and accuracy of the data: HY and QZ; manuscript revision: HY, DDB, RKP, and YL; final approval of manuscript: all authors.

**Conflicts of interest**

HY and QZ have a pending patent application for DNA methylation-based TILs.

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**Competing interests** HY and coauthors are the inventors of a provisional patent filed by Sun Yat-sen University on the subject of this work (202011176816.1).

**Patient consent for publication** Not required.

**Ethics approval** This study was approved by the institutional review board of the Sixth Affiliated Hospital of Sun Yat-sen University (2017ZLYEC-006), and written informed consent was obtained from all patients.

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