Prognostic and therapeutic prediction by screening signature combinations from transcriptome–methylome interactions in oral squamous cell carcinoma

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DNA methylation pattern in oral squamous cell carcinoma (OSCC) remains poorly described. This study aimed to perform a genome-wide integrated analysis of the transcriptome and methylome and assess the efficacy of their prognostic signature model in patients with OSCC. We analyzed transcriptome and methylome data from 391 OSCC samples and 41 adjacent normal samples. A total of 8074 differentially expressed genes (DEGs) and 10,084 differentially expressed CpGs (DMCpGs) were identified. Then 241 DEGs with DMCpGs were identified. According to the prognostic analysis, the prognostic signature of methylation-related differentially expressed genes (mrDEGPS) was established. mrDEGPS consisted of seven prognostic methylation-related genes, including ESRRG, CCNA1, SLC20A1, COL6A6, FCGBP, CDKN2A, and ZNF43. mrDEGPS was a significant stratification factor of survival ($P < 0.00001$) irrespective of the clinical stage. The immune effector components, including B cells, $CD4^+$ T cells, and $CD8^+$ T cells, were decreased in the tumor environment of patients with high mrDEGPS. Immune checkpoint expressions, including CTLA-4, PD-1, LAG3, LGALS9, HAVCR2, and TIGHT, were comprehensively elevated ($P < 0.001$). The estimated half-maximal inhibitory concentration difference between low- and high-risk patients was inconsistent among chemotherapeutic drugs. In conclusion, the transcriptome–methylome interaction pattern in OSCC is complex. mrDEGPS can predict patient survival and responses to immunotherapy and chemotherapy and facilitate clinical decision-making in patients with OSCC.

Oral squamous cell carcinoma (OSCC) is one of the most common malignancies in the head and neck region, which impairs the quality of life. Over the past 30 years, the age-standardized incidence rate was 6.2 and 3.6 per 100,000 for males and females, respectively, and the age-standardized death rate was 3.3 and 1.6 per 100,000 for males and females, respectively. However, the 5-year survival rate after surgery or chemoradiotherapy was only 64.4%, according to the 8th edition of the American Joint Committee on Cancer (AJCC), with age of the patient and stage of OSCC as independent prognostic factors. For early stage OSCC, the survival rate did not significantly increase (69.7%), which indicated the difficulty of survival modification in patients with OSCC. With the inclusion of the depth of invasion and extranodal extension, the 8th edition AJCC staging exhibited superior performance in stratifying the survival of patients with OSCC than that by the former edition. However, stratifying the patient survival remained significantly challenging.

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Recently, it has become evident that epigenetic mis-programming constitutes a core component of cancer initiation and progression. Currently, DNA methylation remains the main epigenetic marker that can be measured reliably using genome-wide studies in large numbers of samples. DNA methylation occurs almost exclusively in CpG dinucleotides. The CpG dinucleotides tend to cluster in regions called CpG islands (CGI), while most tissue-specific differentially methylated regions appear outside of CGIs. A comprehensive study of the profiles of different healthy individuals and tissue types enables the estimation of variance of each CpG site in the methylome. The integrated analysis of methylome and clinical data made prognostic classification feasible based on methylome analysis in colorectal cancer, hepatocellular carcinoma, and leukemia. Moreover, there may be a link between DNA methylation and the tumor microenvironment. Infiltrating immune cells are important participants in the tumor microenvironment (TME); they are involved in proliferation, signal maintenance, cell death resistance, invasion, metastasis, and angiogenesis. Tumor-associated macrophages support disease progression and resistance to therapy by providing malignant cells with trophic and nutritional support. Altered signals from tumor cells produce a suppressive tumor microenvironment for enrichment of inhibitory cells. DNA methylation may participate in the changes of infiltrating immune cells, reflecting a specific immune response to the cancer cell. Immune TME (TIME) is associated with therapy responsiveness of the immune-checkpoint blockade.

Previous studies have preliminarily depicted the roles of DNA methylation in gene expression, patient prognosis, and immune markers in head and neck cancers. However, a comprehensive analysis of transcriptome–methylome interactions and clinical characteristics may be required to mine the potential of methylation orchestration in the management of OSCC. Therefore, this study aimed to perform a genome-wide integrated analysis of the transcriptome and methylome, depict the complex pattern between methylation and gene expression, identify methylation-related differentially expressed genes (mDEGs), and assess the efficacy of their prognostic signature model in predicting patient survival, TIME alterations, and responses to immunotherapy and chemotherapy in patients with OSCC.

Results
Patterns of DMCpGs and differential gene expression (DEG). A flowchart of the study is shown in Fig. 1. A total of 8074 DEGs and 10,084 DMCpGs were identified. Clustering based on the DEGs or DMCpGs revealed two distinctive sample clusters, indicating the possibility to distinguish between OSCC samples and adjacent tissue samples (Fig. 2A,B). Of the 8074 DEGs, 4327 were upregulated and 3747 were downregulated (Fig. 2C). Of the 10,084 DMCpGs, 5937 were hypermethylated DMCpGs (HyperCpGs) and 4147 were hypomethylated DMCpGs (HypoCpGs). Both HyperCpGs and HypoCpGs were correlated with the upregulation and downregulation of genes (Fig. 2D). These results indicate that the regulation of gene expression by methylation may be complex and multimodal.

Relationship between distribution pattern of methylation alterations and gene expression. DMCpGs across different genomic regions were not randomly distributed across the genome. Most DMCpGs are found in the gene body and intergenic regions (IGR). HyperCpGs had a higher proportion located 200 bp upstream of transcriptional start sites (TSS200), 5′-untranslated regions (UTR), and 1st exon, whereas HypoCpGs were located in the gene body and intergenic regions (Fig. 3A). The results indicated that there was a high density of methylated CpGs in several kb regions upstream and downstream of the transcriptional start sites (TSS). The distribution pattern around the CGI differed significantly between HyperCpGs and HypoCpGs. The distribution of HyperCpGs was significantly enriched within CGI (62.9%), whereas HypoCpGs were mostly enriched in the open sea regions (72.6%) (Fig. 3B), indicating that CpG methylation in CGI and non-CGI was potentially functional in gene expression. To further integrate the distribution pattern around TSS, CGI, and gene expression, we plotted the CGI and non-CGI methylation levels of every gene within four expression quartiles grouped by distance to the TSS (Figs. 3C and S1). Then, to check the potential bias owning to unpaired samples, we divided the samples into paired normal samples (pNormal), paired Tumor samples (pTumor) and unpaired single Tumor samples (sTumor) and found that the methylation pattern of pTumor was similar to that of sTumor instead of pNormal, showing the robustness of results (Fig. S2). HypoCpGs proximal to the TSS (approximately ±1 kb) were observed in both OSCC and adjacent samples in the highly expressed CGI genes. CGI CpGs with low gene expression levels exhibited a higher number of HyperCpGs around TSSs in OSCC samples than in adjacent tissues, indicating that hypermethylation-induced silencing of tumor suppressor genes was more evident in transcriptionally silent genes with CGI. Moreover, HypoCpGs were observed in non-CGI CpGs with low gene expression and in those with high gene expression away from the TSS, suggesting a potential role of non-CGI HypoCpGs in the regulation of oncogenes and tumor-suppressor genes.

Correlation between methylation levels and gene expressions. We recognized that HyperCpGs and HypoCpGs might function differently in genes with CGI and non-CGI CpGs in OSCC samples. To further understand the correlations between CpG alterations and gene expression, we defined CpGs as positively-correlated CpGs (PosCpGs), wherein HyperCpGs and HypoCpGs were positively correlated with upregulation and downregulation of gene expression, respectively, and vice versa, as negatively-correlated CpGs (NegCpGs). We plotted the correlation between methylation and gene expression levels grouped by the genomic distribution of methylated CpGs. NegCpGs were highly concentrated around CGIs and within the promoter, while PosCpGs were highly distributed within the gene body, and the majority of CpGs were open sea (Fig. 4A). The correlation modes were gradient from CGI to the open sea. The results indicated that the correlation between methylation and gene expression levels was diverse but organized across the genomic distribution. However, not all correlations between methylation and gene expression levels were statistically significant. We further divided the
correlation into significantly positive, significantly negative, and non-significant. Significant associations were observed more frequently in the DMCpGs than in the non-DMCpGs. CGI CpGs located in the promoter tended to have significantly negative associations with gene expression, whereas CpGs away from CGI and located in the gene body had significantly positive associations (Fig. 4B).

**Screening of prognostic DEGs with significant methylation correlation.** To further screen the CpGs significantly correlated with gene expression, we intersected DMCpGs with PosCpGs or NegCpGs and
obtained 522 and 384 DMCpGs that exhibited significant positive and negative correlations with gene expression, respectively (Fig. 5A). We then intersected DEGs with genes containing selected PosCpGs or NegCpGs and found 121 DEGs with positively related DMCpGs, 96 DEGs with negatively related DMCpGs, and 24 DEGs with both positively and negatively related DMCpGs (Fig. 5B). The intersection results further indicate the complexity of methylation regulation. It was difficult to predict the final gene expression, particularly when multiple methylation sites were altered. Here, we defined DEGs with significantly correlated DMCpGs as methylation related DEGs (mrDEGs). To determine the prognostic value of the 241 mrDEGs in OSCC, a multivariate Cox regression analysis was performed with a cutoff of \( P < 0.01 \). We found that three mrDEGs (ESRRG, CCNA1, and SLC20A1) were associated with a poor prognosis and significantly increased hazard ratio (HR), whereas five mrDEGs (COL6A6, FCGBP, CDKN2A, MEI1, and ZNF43) served as protective genes with HR < 1 (Fig. 5C). However, MEI1 did not reach a significant level in the Least Absolute Shrinkage and Selection Operator (LASSO) and was thus abandoned.

**Establishment and validation of the prognostic signature.** We then built an mrDEGs predictive signature (mrDEGPS) using seven survival-relevant mrDEGs. The mrDEGPS for each patient was calculated using the following formula:

Figure 2. Alterations of transcriptome and methylome profiling in oral squamous cell carcinoma (OSCC) samples. (A) Unsupervised hierarchical clustering of normal and OSCC samples using the top 100 differentially expressed gene (DEG) probes according to F score. The heatmap shows DEGs arranged in rows (upregulation colored in red and downregulation in dark blue) and samples in columns (OSCC samples colored in pink and adjacent normal samples in turquoise). (B) Volcano plot of DEGs using the cutoff of \( |\log FC| \geq 1 \) and the \( p_{adj} < 0.05 \). LogFC \( \leq \) were downregulated DEGs colored in blue and LogFC \( \geq \) were upregulated DEGs colored in red. (C) Unsupervised hierarchical clustering of normal and OSCC samples using the top 100 differentially methylated probes according to F score. The heatmap shows differentially methylated CpGs (DMCpGs) arranged in rows (hypermethylation colored in red and hypomethylation in dark blue) and samples in columns (OSCC samples colored in pink and adjacent normal samples in turquoise). (D) Scatter plot of DMCpGs and related DEGs using the cutoff of \( |\log FC| \geq 1 \), \( |\Delta of Beta| \geq 0.3 \), and the \( p_{adj} < 0.05 \). LogFC \( \leq 1 \) were downregulated DEGs colored in blue and LogFC \( \geq 1 \) were upregulated DEGs colored in red. The delta of Beta \( \geq 0.3 \) were hypermethylated DEGs in the upper region and delta of Beta \( \leq 3 \) were hypomethylated DEGs in the lower region. This heatmaps was generated by the R (version 4.1.0, https://www.r-project.org).
We divided the patients with OSCC into low- and high-risk groups (Fig. 5D). The expressions of *ESRRG*, *CCNA1*, and *SLC20A1* were comparatively higher in the high-risk group than in the other groups (Fig. 5E). Prognosis comparison showed that low-risk patients had significantly higher overall survival (OS) ($P = 4.587 \times 10^{-10}$), disease-specific survival (DSS) ($P = 1.28 \times 10^{-7}$), and progression-free survival (PFS) ($P = 9.588 \times 10^{-6}$) than that in high-risk patients (Fig. 5F). The validation analysis in a small sample size ($n = 97$) demonstrated that patients in

$$mrDEGPS = 0.248 \exp(ESRRG) + 0.047 \exp(CCNA1) + 0.200 \exp(SLC20A1) - 0.138 \exp(COL6A6) - 0.0507 \exp(FCGBP) - 0.045 \exp(CDKN2A) - 0.161 \exp(ZNF43)$$

Figure 3. Complex pattern of DNA methylation in oral squamous cell carcinoma (OSCC) samples. (A) Bar plots of numbers and ratios of hypermethylated and hypomethylated CpGs grouped by gene features. The gene features colored in cold tones were divided into regions 200 bp upstream of transcriptional start site (TSS200), 1500 bp upstream of TSS (TSS1500), 5\textsuperscript{-}untranslated regions (5\textsuperscript{-}UTR), 3\textsuperscript{-}UTR, the first exon (1st exon), and gene body and intergenic regions (IGR). (B) Bar plots of numbers and ratios of hypermethylated and hypomethylated CpGs grouped by gene and CpG island (CGI) features. The CGI features were divided according to the distance to CGI, including island, shore (regions within 2 kb upstream or downstream of island), shelf (regions of 2–4 kb upstream or downstream of island), and open sea (regions outside of island, shore, and shelf). (C) Plots of the median methylation values per 100 bp distance grouped by gene expression quartiles based on the expression levels in either tumor or normal samples ($n = 5493, 5493, 5493, and 5496$ for Q1, Q2, Q3, and Q4, respectively; Q4 is the highest expression) showing methylation ratio at 100 bp segments including genomic loci within and outside CpG islands in genes with promoter associated CpG islands. The curves were colored in red in OSCC samples and in blue in normal samples.
the high-risk group had poorer overall survival ($P = 3.098e^{-2}$) than those patients in the low-risk group (Fig. 5G). The area under the curve (AUC) of 1-year survival was 0.714 and 0.702 in the training and validation datasets, respectively. Compared with the clinical stage, mrDEGPS displayed superior predictive performance.

**Association between mrDEGPS with clinical features and human papillomavirus (HPV).** Next, we explored the association between the mrDEGPS and clinical features (Fig. 5H). We found that the mrDEGPS did not differ between patients with different age groups, gender, alcohol consumption, and surgical margin status ($P > 0.05$). Notably, there was no difference in mrDEGPS in the different clinical stages and histopathological grades ($P > 0.05$), implying the absence of linear correlation between mrDEGPS and these traditional survival stratification factors; thus, mrDEGPS was a potentially independent prognostic factor. We also found that the mrDEGPS was significantly associated with lymphovascular and perineural invasion. As both lymphovascular and perineural invasion were independent prognostic factors, this result was consistent with mrDEGPS as a
prognostic factor. Moreover, the human papillomavirus (HPV) infection was associated with higher mrDEGPS ($P < 0.001$), indicating that HPV might partially promote epigenetic alterations. We further included patients with HPV status not available (NA) in the analysis and found no differences in mrDEGPS between HPV (NA), HPV (+), or HPV (−) (Fig. S1), which indicated that the mixed population neutralized the difference between HPV (+) and HPV (−).

Association between mrDEGPS and TIME. Since the selected mrDEGs were relevant to the immune response, we hypothesized that mrDEGPS might have the capacity to identify alterations in TIME. The results showed that the high-risk group samples had reduced proportions of exhausted T cells, type 1 regulatory T cells, follicular T-helper cells, dendritic cells, B cells, CD4+ T cells, and CD8+ T cells, and increased proportions showed that the high-risk group samples had reduced proportions of exhausted T cells, type 1 regulatory T cells, natural killer T cells, natural killer cells, and neutrophils. (Figs. 6A and S3). It has been reported that DCs, CD8+ T cells, and CD4+ T cells display a beneficial effect on survival, while neutrophils, NK cells, and Tem cells display harmful effects in breast cancer. Our results implied that high-risk patients might have an altered survival–harmful TIME. To further characterize the potential signaling pathways involved in the influence of mrDEGPS, gene set enrichment analysis (GSEA) was performed to enrich the Kyoto Encyclopedia of Genes and Genomes pathways ranked by gene correlation values with mrDEGPS. Half of the enriched pathways were associated with immune processes like “primary immunodeficiency,” “allograft rejection,” “autoimmune thyroid disease,” “intestinal immune network for IgA production,” “T cell receptor signaling pathway,” “antigen processing and presentation,” “natural killer cell-mediated cytotoxicity,” and “B cell receptor signaling pathway” (adi. $P < 0.01$) (Fig. 6B).

Prediction of immunotherapy and chemotherapy by mrDEGPS. T-cell receptor signaling is the essential basis for immunotherapy and may participate in chemotherapy resistance. Current immunotherapy is mainly achieved by antibody blocking of CTLA-4 or PD-1 pathway. Studies indicate that LAG3, LGALS9, HAVCR2, and TIGHT could be the next-generation immunotherapy checkpoints. We found that all these immunotherapy checkpoints were downregulated in high-risk patients ($P < 0.001$) (Fig. 6C). However, there was no significant difference in PD-L1, indicating that mrDEGPS was correlated with TIME rather than with tumor cells (Fig. S4). The mrDEGPS was higher in patients with a response ($P = 0.0005$) and the response rate was significantly lower in high-risk patients ($P = 0.0024$) (Fig. 6D). These results indicate that mrDEGPS low-risk patients might benefit from immunotherapy. For chemotherapy, the estimated half-maximal inhibitory concentration difference between low- and high-risk patients was inconsistent among drugs, which was not significant for gefitinib, lower in low-risk patients for rapamycin, and lower in high-risk patients for cisplatin, docetaxel, sorafenib, erlotinib, and gemcitabine (Figs. 6E and S5). We observed that the result might provide a reference for chemotherapeutic drug selection for individuals, and thus facilitated the survival of patients with OSCC. In summary, mrDEGPS screened from complex transcriptome–methylome interactions could facilitate the prediction of survival and immunotherapeutic efficacy in patients with OSCC (Fig. 7).

Discussion

Several studies have demonstrated a strong relationship between epigenetic and genetic aberrations in tumorigenesis. It is commonly believed that epigenetic changes, such as DNA methylation, can drive abnormal gene expression of crucial genes involved in the development and progression of cancer, including head and neck cancer. Hypermethylation of tumor suppressor genes and hypomethylation of proto-oncogenes at the promoter sites are associated with carcinogenesis and progression of OSCC. For example, several studies have suggested that hypermethylation of PAX1 and ZNF582 genes is associated with aggressive progression and poor survival.

Although the effect of promoter methylation changes has been studied extensively, increasing evidence from genome-wide methylome studies suggests that the methylation patterns are complex and cancer-type-specific. The term “CpG island methylator phenotype” has been repeatedly used over decades to describe widespread CpG island promoter methylation. However, only around 4–8% CGIs exhibit tissue-specific methylation, while approximately 70% of annotated gene promoters are associated with a specific CGI. Therefore, there must be undisclosed cancer-type-specific methylation outside CGIs. Our results showed that CpGps away from islands, denoted as open sea CpGs, may facilitate gene expression of oncogenes in OSCC.

Despite CGI features, cumulative evidence indicates that the transcriptome–methylome interaction is not restricted to promoters and TSS. In contrast to the repression of promoter methylation on expression, gene body methylation orchestrates transcription in a complex pattern, which is in contrast to the repression of promoter methylation. Approximately half of CGIs in mammalian genomes are not associated with a known gene promoter and are referred to as orphan CGIs. Orphan CGIs that do not map to promoters of any protein-coding or non-coding transcripts but possess chromatin and transcriptional markers may reflect enhancer activity. Orphan CGIs display most of the evolutionarily conserved methylation differences among tissues, indicating their possible role in tissue specification. Our results indicated that CpGps located in the gene body might have a positive association with gene expression, suggesting that most tissue-specific methylation CGIs are not located at promoter regions in OSCC. These findings may potentially facilitate research on aberrant methylation of these rarely investigated regions.

Another hypothesis regarding the non-promoter DMCGps is that most genes have two or more TSSs; therefore downstream TSS are probably within the bodies of the transcriptional units of the alternative upstream promoters. A large-scale full-length cDNA analysis to explore the budding yeast transcriptome showed that alternative promoters could be located at CGIs or non-CGIs, and combinations of an upstream non-CGI and a
The low- and high-risk patients were divided by the cutoff of mrDEGPS = 1.0256. OS was calculated based on the follow-up time, if patients survived (blue), or time until death (red). (E) Heatmap of high- and low-risk OSCC samples grouped by mrDEGPS according to the simulation result of mrDEGPS model. The heatmap shows mrDEGs arranged in rows (upregulation colored in red and downregulation in dark blue) and samples in columns (high-risk OSCC samples colored in pink and low-risk OSCC samples in turquoise). (F) Kaplan–Meier plots for OS, disease-specific survival, and progression-free survival grouped by low- and high-risk patients in the training set (The Cancer Genome Atlas (TCGA) dataset). (G) Kaplan–Meier plots for OS grouped by low- and high-risk patients in a validation set (GSE41613). (H) Box plots of mrDEGPS grouped by different clinical features including age, gender, clinical stage, alcohol habit, histopathological grade, HPV infection, lymphovascular invasion, surgical margin, and perineural invasion in the training set (TCGA dataset). ** and *** indicate P < 0.05, P < 0.01, and P < 0.001, respectively. (I) Violin plot of mrDEGS grouped by HPV (+), HPV (−), and HPV status not available (NA) in a validation set (GSE87053).
Identification and validation of mrDEGPS. Kaplan–Meier analysis was used to evaluate the relationship between the genes and survival of patients with OSCC. The LASSO binary logistic regression model and multivariate Cox regression were adopted after primary filtration. The linear combination of the regression coefficient \( \hat{\beta} \) derived from the multivariate Cox regression model and multiplied by the corresponding mRNA levels were adopted for the multivariate Cox regression model and multiplied by the corresponding mRNA levels was used to plot the distribution of significant probes around the TSS \( \pm 5 \) kb. The methylation level of each probe was measured by the beta value, which ranged from 0 to 1 (representing unmethylated to fully methylated levels, respectively). Similarly, we removed probes with missing beta values in ≥ 5% samples. The remaining probes with missing beta values were imputed using the k-nearest neighbors algorithm.

We mapped the probes to the promoter regions of the genes, which were defined as \( -1.5 \) to 0 kb regions around the TSS. Next, the DNA methylation level of a gene was defined as the average beta value of the probes that mapped to its promoter region. Finally, samples with paired mRNA expression and DNA methylation profiles were analyzed, which involved 17,481 genes and 384 samples, to determine DMCpGs and DEGs. The Spearman rank correlation of CpG methylation levels with related gene expression was analyzed, and significant associations were based on the criteria of |correlation value|> 0.3 and FDR < 0.05.
 Patients were divided into high- and low-risk groups by setting the median risk score as the cut-off value. The OS, DSS, and PFS of the two groups were calculated using the Kaplan–Meier method with the log-rank test. Receiver operating characteristic (ROC) curves were generated to assess the predictive performance of the prognostic model. The expression patterns of genes in this prognostic model were visualized using the “pheatmap” package. In the validation analysis, we verified the Kaplan–Meier plot and ROC test in GEO using another OSCC cohort, GSE41613. The clinical features of the patients with low or high mrDEGPS were analyzed and compared.

**Analysis of correlation between the survival risk model and tumor immune infiltration.** ImmuneCellAI (http://bioinfo.life.hust.edu.cn/ImmuCellAI#!/) was utilized to analyze the fraction of 24 types of immune cells in high- and low-risk samples with OSCC. These immune cells included 18 subtypes of T cells, namely CD4+ T cells; CD8+ T cells; naïve CD4+ T cells; naïve CD8+ T cells; cytotoxic T cells; exhausted T cells; type 1 regulatory T cells; natural regulatory T cells; induced regulatory T cells; T-helper 1, 2, and 17 cells; follicular T-helper cells; central memory T cells; effector memory T cells; natural killer T cells; mucosal-associated invariant T cells; and gamma-delta T cells, as well as six other types of immune cells including B cells, natural killer cells, monocytes, macrophages, neutrophils, and dendritic cells. Immune Cell Abundance Identifier (ImmuCellAI) built an immune cell-based support vector machine model for the prediction of immunotherapy response (AUC: 0.80–0.91), and the model was used to estimate the response results of OSCC.

GSEA was performed using the R package “clusterProfiler” to determine the enrichment of previously defined biological processes in the ranked correlated gene with risk score using RNA-seq data from TCGA-OSCC cohort. The raw count data of gene expression from the TCGA-OSCC cohort were normalized using the variance stabilizing transformation function in R package “DESeq2,” and then submitted to ImmuneCellAI (http://bioinfo.life.hust.edu.cn/web/ImmuCellAI/) to estimate the abundance of immune cells, particularly the T-cell proportions, and to predict the response of immune-checkpoint inhibitor treatments.
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Author contributions
All the authors contributed the concept. C.S., S.L., and Y.C designed the study, developed methodology, and acquired, analyzed, and interpreted data; Y.C. wrote the manuscript. C.S., S.L., X.T., C.M., R.W, X.M. and X.W. revised the manuscript. X.W. and Y.C. supervised the study.

Competing interests
The authors declare no competing interests.

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