Integrative Multi–Omics Analysis Reveals Candidate Biomarkers for Oral Squamous Cell Carcinoma

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Oral squamous cell carcinoma (OSCC) is one of the most common types of cancer worldwide. Due to the lack of early detection and treatment, the survival rate of OSCC remains poor and the incidence of OSCC has not decreased during the past decades. To explore potential biomarkers and therapeutic targets for OSCC, we analyzed differentially expressed genes (DEGs) associated with OSCC using RNA sequencing technology. Methylation–regulated and differentially expressed genes (MeDEGs) of OSCC were further identified via an integrative approach by examining publicly available methylomic datasets together with our transcriptomic data. Protein–protein interaction (PPI) networks of MeDEGs were constructed and highly connected hub MeDEGs were identified from these PPI networks. Subsequently, expression and survival analyses of hub genes were performed using The Cancer Genome Atlas (TCGA) database and the Gene Expression Profiling Interactive Analysis (GEPIA) online tool. A total of 56 upregulated MeDEGs and 170 downregulated MeDEGs were identified in OSCC. Eleven hub genes with high degree of connectivity were picked out from the PPI networks constructed by those MeDEGs. Among them, the expression level of four hub genes (CTLA4, CDSN, ACTN2, and MYH11) were found to be significantly changed in the head and neck squamous carcinoma (HNSC) patients. Three hypomethylated hub genes (CTLA4, GPR29, and TNFSF11) and one hypermethylated hub gene (ISL1) were found to be significantly associated with overall survival (OS) of HNSC patients. Therefore, these hub genes may serve as potential DNA methylation biomarkers and therapeutic targets of OSCC.

Keywords: oral squamous cell carcinoma, multi-omics, DEGs, DMGs, MeDEGs

INTRODUCTION

Oral squamous cell carcinoma (OSCC) is the seventh most common cancer in males under the age of 60, with about 400,000 new cases reported annually worldwide (1). Geographically, the incidences of OSCC in developing countries like India and Brazil could be more than tenfold higher than those in Western countries. Although OSCC is predominantly seen in male over the age of 65, the disease
appears to be increasing in younger patients under the age of 45 and the gender difference has been decreasing over the past few decades. Thus, OSCC represents a major public health problem, especially in poor and developing countries.

Despite recent advances in therapeutic modalities, the five–year morbidity rate of OSCC remains high and the survival rate stays dismal. Under the standard treatment plan, the outcomes of patients with OSCC could be completely different (2, 3). Therefore, uncovering the complex molecular changes leading to the development of OSCC as well as the factors contributing to the differential prognosis among OSCC patients represents a continuously unmet medical need.

The etiology of OSCC is complicated and involves both genetic and environmental factors (4). DNA methylation is one of the most abundant epigenetic modifications with key regulatory functions in gene transcription and genomic stability. Growing evidence suggests that alterations of DNA methylation contribute to carcinogenesis. In OSCC, both global and site–specific DNA hypomethylation have been reported (5–7). Kim et al. found that three out of 14 tumor suppressor genes (TSGs) are hypermethylated and transcriptionally silenced in OSCC (8). Using microarray–based technologies, the methylation alternations in OSCC have been extensively studied and reported by multiple groups (5, 9–15). Considering the fact that the majority of genes that are abnormally methylated in cancers have no corresponding changes in mRNA expression, it becomes important to understand the functional profile of these massive genome–wide methylomic data. Two recent studies have taken a meta–analysis approach and analyzed the impact of DNA methylation on gene expression and biological pathways using publicly available transcriptome and methylome datasets (16, 17). Due to the unknown heterogeneity within and across individual studies with regard to clinical diagnosis, tumor tissue dissection, sample processing, etc., more research efforts are needed to gain a trustworthy assessment of methylomic signatures in OSCC.

In this study, we have characterized the transcriptome of fresh OSCC tumor and paired normal adjacent tissue (NAT) using RNA sequencing technology. Two methylation–profiling datasets, GSE38532 and GSE46802, generated from the same microarray platform and of paired case–control design, were selected and integrated with our transcriptome data for multi–omics analysis. Methylation–regulated and differentially expressed genes (MeDEGs) were identified, hub genes were screened through protein–protein interaction (PPI) network analysis, and further validated in The Cancer Genome Atlas (TCGA) database. Using this integrated approach, we were able to identify novel DNA methylation regulated genes with potential clinical applications in OSCC.

MATERIALS AND METHODS

Tumor Biospecimen Collection and RNA Extraction

In total, 17 paired samples of OSCC and NAT were collected from newly diagnosed patients who underwent surgical resection and had received no prior treatment for the disease at Xiangya Hospital of Central South University (CSU). The samples were collected in accordance with the guidelines issued by the Ethics Committee of School of Life Sciences, CSU (Record#2020–1–42). Written informed consent was obtained to all participants in this study.

All biospecimens from patients were pathologically reviewed and confirmed by two independent pathologists. Among these patients, there were four with stage I, four with stage II, four with stage III, and five with stage IV. Their demographic characteristics are presented in Table 1. After surgical removal, all samples were immediately transferred to the lab on ice using CO2 independent medium (18045088, Gibco, USA). Tissue blocks of about 30 mg were further dissected out and used for extraction of total RNA following the manufacturer’s instruction of the AllPrep DNA/RNA/miRNA Universal Kit (80224, Qiagen, USA).

RNA Sequencing and Data Processing

For library preparation, 1 μg of total RNA from each sample was depleted for rRNA using the Ribo–Zero rRNA Removal Kit (RZHI046, EPICENTRE, USA), followed by construction of sequencing library using the TruSeq RNA Sample Prep Kit v2 (RS–122, Illumina, USA). Individual RNA libraries were pooled and sequenced using paired–end sequencing with 150 bp reads on an Illumina HiSeq3000 instrument.

Adapter and low–quality reads were first trimmed using Trim Galore (version 0.4.4) in paired–end mode with default parameters. Trimmed reads were then mapped to the hg19 reference genome to count mapped reads for all human transcripts. Normalization and analysis of differentially expressed genes (DEGs) were carried out using multifactor analysis (tumor vs. NAT) in the DESeq2 R program package (version 1.30.1). Significant genes were identified with statistical cut–off of [Log2FC] > 2 and false discovery rate (FDR) < 0.05.

Methylation Data Acquisition and MeDEGs Identification

Microarray–based gene methylation profiling datasets GSE38532 and GSE46802 were obtained from the NCBI gene expression omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/). Forty OSCC and 40 matched NAT specimens were obtained in GSE38532, while ten paired OSCC and NAT specimens were included in GSE46802. Both microarrays used the Illumina GPL8490 microarray platform (HumanMethylation 27 BeadChip, HumanMethylation27_270596_v.1.2, Illumina, USA). This methylation chip contains 27,578 individual CpG sites spread across the proximal promoter regions, defined as 1500 bp upstream of the transcription start site (TSS1500), of 14,495 genes.

The GEO2R online software was utilized to analyze these two datasets to identify differentially methylated genes (DMGs). FDR < 0.05 and delta β–value > 0.1 were used as the cut–off standards during analysis as previously described (18). Eventually, Venndiagram package (version 1.6.20) of R program was used to obtain the overlaps between DMGs and DEGs and to identify MeDEGs, including downregulated genes with a hypermethylated pattern and upregulated genes with a hypomethylated pattern.
Table 1: Patient demographics and clinical characteristics.

| Characteristics          | Parameters                  | No. of patients |
|-------------------------|-----------------------------|-----------------|
| Age (years)             |                             |                 |
| <45                     |                             | 4               |
| 46–60                   |                             | 8               |
| >61                     |                             | 5               |
| Gender                  |                             |                 |
| Male                    |                             | 16              |
| Female                  |                             | 1               |
| Stage                   |                             |                 |
| I                       |                             | 4               |
| II                      |                             | 4               |
| III                     |                             | 4               |
| IV                      |                             | 5               |
| Degree of differentiation|                             |                 |
| Well differentiated     |                             | 8               |
| Moderately differentiated|                            | 7               |
| Poorly differentiated   |                             | 1               |
| ND                      |                             | 1               |
| Cancer site             |                             |                 |
| Tongue                  |                             | 8               |
| Gingiva                 |                             | 3               |
| Inner cheek             |                             | 6               |
| P53 expression          |                             |                 |
| Positive                |                             | 5               |
| Negative                |                             | 5               |
| ND                      |                             | 7               |
| Ki67 expression         |                             |                 |
| >60% positive           |                             | 3               |
| 30%–60% positive        |                             | 5               |
| <30% positive           |                             | 3               |
| ND                      |                             | 6               |

ND, Not determined.

Functional Enrichment Analysis

Functional clustering annotation of the DEGs and MeDEGs were conducted via the Metascape online tool (19). For functional clustering, enriched annotations from the Gene Ontology (GO) biological processes and the Reactome gene sets were included. Analyses were carried out with the default parameters of minimal overlap of 3, minimal enrichment ratio of 1.5, and \( p \)-value < 0.01 as previously described (20).

Protein–Protein Interaction (PPI) Network Construction

The PPI networks of downregulated genes with a hypermethylated pattern and upregulated genes with a hypomethylated pattern were constructed using the Search Tool for the Retrieval of Interacting Genes (STRING, version 11.0) online database. PPI with an interaction score > 0.4 was visualized in the final STRING network. Protein networks are clustered to MCL inflation parameter of 2 (21).

Identification of Potential Hub Genes

To identify potential hub genes, the cytoHubba plugin of the Cytoscape (version 3.8.2, http://www.cytoscape.org/) software was used to screen modules within PPI network. Four topological analysis methods were individually utilized (MCC, MNC, Degree, and EPC) to obtain the top ten important genes from each PPI network. Common genes selected from all four methods were regarded as potential hub genes with consensus.

Validation of Hub Genes

The Gene Expression Profiling Interactive Analysis (GEPIA) online platform provides fast evaluation between the survival effect and the expression profile analysis of DEGs in a given cancer type. To validate aforementioned hub genes, the relative expression levels of these genes in head and neck squamous cell carcinoma (HNSC) were identified with statistical cut-off of \( |\text{log}_{2}\text{FC}| > 1 \) and \( p \)-value < 0.05 as previously described (22). In addition, the overall survival (OS) effect of hub genes in HNSC was estimated by calculating the log–rank \( p \)-value and the hazard ratio (HR). Patients were divided into high (higher than the 75th percentile) and low (lower than the 25th percentile) expression groups based on the normalized expression level of the selected gene across all HNSC patients in the database (23). Since there is no OSCC category in both GEPIA and TCGA databases, and OSCC accounts for ~95% of HNSC, we chose HNSC as the category for these validation analyses.

Immune Infiltration Analysis

Infiltrations of immune cells in OSCC tumors and NATs were estimated using TIMER web tool (24, 25). Briefly, RNA–seq data in TPM (transcripts per million) format from the 17 paired samples of OSCC and NAT were calculated. The infiltration abundance of B cell, CD4+ T cell, CD8+ T cell, neutrophil, macrophage, and myeloid dendritic cell in each sample were evaluated using TIMER2.0. Similar procedures were used to analyze data from TCGA-HNSC and paired NAT samples.

RESULTS

Characterization of RNA Sequencing and Mapping

Using RNA sequencing, a total of 1418 million and 1289 million cleaned reads were obtained from pooled OSCC and NAT
libraries with 65–106 million reads per library for all samples. In all 34 libraries, > 92.9% cleaned bases have Phred quality score (Q-scores) above 30, indicating a high base call accuracy. Forty-two to eighty-one million reads were uniquely mapped to the hg19 reference genome for each sample, with an average unique mapping ratio of 92% (Table 2).

**Identification of DEGs in OSCC**

Principal component analysis (PCA) of RNA–seq data showed that deconvoluted data from OSCC samples and NATs were clustered into different patterns (Figure 1A), suggesting the underlying differences between these two conditions. A total of 778 upregulated genes and 2211 downregulated genes were obtained by the DESeq2 package using the cut-off criteria of FDR < 0.05 and |log2FC| > 2, as shown in Figure 1B by a volcano plot. Further functional clustering analysis using the online tools in Metascape shown that upregulated genes were mainly involved in functions such as response to bacterium, formation of the cornified envelope, extracellular matrix organization, antimicrobial humoral response, etc. (Figure 1C), while downregulated genes were mainly involved in muscle system process, muscle contraction, muscle structure development, muscle cell development, etc. (Figure 1D).

**Identification of Aberrantly MeDEGs in OSCC**

To identify potential MeDEGs (i.e., genes that exhibit inverse correlation between their alterations of DNA methylation at promoter regions and gene expression), we performed a multi-omics analysis by integrating our transcriptomic data and publicly available methylomic data (GSE38532 and GSE46802). As shown in the Venn diagrams in Figures 2A, B, 56 upregulated and hypomethylated genes and 170 downregulated and hypermethylated genes (Table 3) were identified from this study. The heatmap of these potential MeDEGs is demonstrated in Figure 2C.

**Functional Enrichment Analyses of Potential MeDEGs**

To evaluate the biological functions of these potential MeDEGs, functional clustering was analyzed using the online tools in Metascape. As shown in Figures 2D, E, genes that were

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**TABLE 2 | RNA sequencing profile.**

| Sample          | Clean reads | Q30 rate | Clean reads (no rRNA) | Unique mapped | Unique mapping ratio |
|-----------------|-------------|----------|-----------------------|---------------|----------------------|
| Oral cancer samples |             |          |                       |               |                      |
| C1              | 86871496    | 93.80%   | 73156876              | 66641472      | 93%                  |
| C2              | 95042470    | 94.45%   | 91337822              | 81376448      | 92%                  |
| C3              | 77878418    | 93.85%   | 6497966               | 48192246      | 91%                  |
| C4              | 94390710    | 94.50%   | 89400168              | 77485932      | 93%                  |
| C5              | 86429488    | 94.05%   | 74290102              | 67319063      | 93%                  |
| C6              | 84077700    | 93.80%   | 77808674              | 60775290      | 94%                  |
| C7              | 89965658    | 93.85%   | 82408242              | 73256107      | 92%                  |
| C8              | 76815890    | 94.10%   | 74169890              | 51400166      | 83%                  |
| C9              | 69672812    | 93.50%   | 60281976              | 51179656      | 88%                  |
| C10             | 73618490    | 93.00%   | 69337410              | 57362728      | 96%                  |
| C11             | 73679338    | 93.00%   | 70965340              | 62125998      | 91%                  |
| C12             | 69570852    | 94.05%   | 67902214              | 61966005      | 93%                  |
| C13             | 71761106    | 93.60%   | 63710848              | 54848328      | 90%                  |
| C14             | 86873658    | 94.25%   | 81753826              | 73884843      | 93%                  |
| C15             | 10602796    | 94.45%   | 91482292              | 81698922      | 91%                  |
| C16             | 91728986    | 94.40%   | 8396796               | 70456741      | 92%                  |
| C17             | 83459218    | 93.65%   | 79658790              | 70201492      | 90%                  |
| Matched adjacent normal tissues |             |          |                       |               |                      |
| N1              | 69157408    | 93.50%   | 67681854              | 63645851      | 96%                  |
| N2              | 65039548    | 94.00%   | 63998054              | 51351866      | 92%                  |
| N3              | 67139172    | 93.70%   | 64235992              | 47105221      | 90%                  |
| N4              | 76287040    | 93.65%   | 65877516              | 53262626      | 93%                  |
| N5              | 7844622    | 93.60%   | 72012974              | 64120834      | 92%                  |
| N6              | 71185918    | 93.85%   | 63433540              | 52672465      | 94%                  |
| N7              | 64770514    | 93.85%   | 63831196              | 52469386      | 97%                  |
| N8              | 83902570    | 94.25%   | 82497918              | 75068818      | 95%                  |
| N9              | 71547434    | 92.95%   | 60248746              | 63107749      | 94%                  |
| N10             | 77368874    | 92.90%   | 67161816              | 59603271      | 94%                  |
| N11             | 87521000    | 93.70%   | 81245060              | 72857823      | 94%                  |
| N12             | 65356658    | 93.65%   | 62147174              | 54454670      | 94%                  |
| N13             | 90320989    | 94.15%   | 88577280              | 58883887      | 94%                  |
| N14             | 100126930   | 93.85%   | 90108998              | 80048283      | 94%                  |
| N15             | 67871766    | 94.15%   | 66695120              | 60492545      | 95%                  |
| N16             | 70708242    | 93.60%   | 67644350              | 42576164      | 90%                  |
| N17             | 81724362    | 93.50%   | 77076054              | 71227418      | 95%                  |

Q30 rate: The percentage of base with a Phred value > 30.

Unique mapped: The number of clean reads that mapped onto the hg19 reference genome.
upregulated and hypomethylated were enriched in biological processes such as leukocyte proliferation, formation of the cornified envelope, response to bacterium, calcium ion homeostasis, etc., while downregulated and hypermethylated genes were linked to neuronal system, signal release, muscle system process, regulation of ion transmembrane transport, etc. To further understand the contribution of DNA methylation to the altered gene expression in OSCC, we evaluated the ratio of DEGs that were inversely affected by methylation changes in each enriched functional cluster. As shown in Figures 1C, D,
differentially upregulated genes involved in functions such as intermediate filament cytoskeleton organization, defense response to Gram−negative bacterium, establishment of skin barrier, and regulation of cell adhesion were often hypomethylated (22.22%, 21.74%, 20.00%, and 19.15%, respectively). For differentially downregulated genes, those involved in neuronal system, potassium ion transport, regulation of membrane potential, and cell junction organization were most likely to be hypermethylated (32.84%, 28.57%, 20.48%, and 20.00%, respectively).

PPI Network Construction and Hub Gene Identification

The PPI network of 56 upregulated MeDEGs and 170 downregulated MeDEGs were constructed using the STRING v11.0 database and visualized via the Cytoscape software. Thirty nodes and 38 edges were identified from the network of upregulated MeDEGs (Figure 3A), and 121 nodes and 207 edges were found from the network of downregulated MeDEGs (Figure 3B). Within each PPI network, the top 10 highly interacted genes were evaluated using four different calculation methods (MCC, MNC, Degree, and EPC) individually. Common genes coming out of all four methods were selected as potential hub genes. Eleven hub genes, including six hypomethylated genes (CTLA4, GRP29, TNFSF11, CD80, CDSN, and PI3) and five hypermethylated genes (ACTN2, MYOD1, ISL1, MYH11, and PAX7) were identified from these networks (Table 4 and Figures 4A, B).

Validation of The Hub Genes

To evaluate the clinical relevance of these potential hub genes in OSCC, we assessed their expression levels in normal and tumor tissues using the GEPIA online tool. Since OSCC accounts for ~95% of all HNSC cases and a specific OSCC category is not included in TCGA database, similar to other studies, TCGA−HNSC data were obtained and used in this analysis (26). As shown in Figures 5A–K, the expression levels of four hub genes (CTLA4, CDSN, ACTN2, and MYH11) were significantly dysregulated in HNSC. The two hypomethylated hub genes, CTLA4 and CDSN, were significantly upregulated, and two hypermethylated hub genes, ACTN2 and MYH11, were downregulated in the HNSC tumor samples. All remaining hub genes, although without statistic strength, showed a correct trend with regard to gene expression in tumor tissues compared to those in NATs.

In addition, we explored the prognostic value of these 11 hub genes by examining the association between the expression level
of each gene and the OS of the HNSC cases using the Kaplan–Meier plotter (Figures 6A–K). The expression levels of four hub genes showed significant correlation with the HNSC clinical outcome. High expression levels of all three hypomethylated and upregulated hub genes (CTLA4, GPR29 and TNFSF11) presented a trend of association with longer OS (Figures 6A–C), while low expression of hypermethylated and downregulated hub gene ISL1 was found to be significantly associated with shorter OS (Figure 6I).

Considering the fact that HNSC datasets were used in these analyses, it may provide more insights if OSCC datasets could be specifically classified in TCGA and GEPIA to allow reexamination of the prognostic value of these genes.

**DISCUSSION**

The purpose of this study is to gain informative insights of the functional and clinically relevant profile of DNA methylation in OSCC. To achieve this goal, we took an integrated multi-omics approach to analyze the impact of DNA methylation on gene expression and biological pathways, then validated the association between the expression level of MeDEGs and the OS of HNSC patients in TCGA. Using this integrated approach, 226 DEGs with inverse corresponding DNA methylation changes and seven hub MeDEGs with potential clinical applications were identified. These findings may contribute to the understanding of the molecular mechanism as well as the development of potential DNA methylation biomarkers for OSCC.

There is strong evidence suggesting the involvement of aberrant DNA methylation in OSCC carcinogenesis. Compared to other epigenetic modification, DNA methylation attracts more attention since it is one of the earliest detectable neoplastic changes and is relatively reversible by chemical treatment. Microarray based methylation evaluation provided a powerful tool for studying global changes of DNA methylation. To the best of our knowledge, eight groups have reported their studies identifying DNA methylation signatures in OSCC using a case–control study design, generating a sufficient amount of publicly available methylomic datasets (5, 9–15). Due to the inherent limitation of the high dimensionality of these huge data, an important question is how to select informative genes which are more biologically relevant to clinical outcomes and transfer these massive data into meaningful patterns with key genes and pathways. Recently, studies from two different groups have integrated transcriptomic and methylomic approaches to examine aberrant MeDEGs and their interrelationships in OSCC (16, 17). Since different inputs of publicly available datasets were used in these two studies, their findings barely overlap with each other. Therefore, more studies are needed to evaluate the power of this multi-omics approach and gain sufficient confidence for the key findings.

In our study, we generated RNA profiling from a stringently controlled collection of 17 paired samples of OSCC and NAT. These samples were similarly collected during the surgery...
procedure and immediately processed for RNA extraction to maximally preserve RNA integrity. A total of 778 upregulated genes and 2211 downregulated genes were identified through transcriptome analysis. Meanwhile, two methylation profiling GSE38532 and GSE46804, generated from same microarray platform and with similar paired case–control design, were obtained from public databases. Integrated multi-omics analysis revealed hypermethylation of 7.69% of under-expressed genes (170 of 2211) and hypomethylation of 7.20% of over-expressed genes (56 of 778) from these datasets. The finding that less than 10% DEGs are inversely associated with methylation modification is consistent with overall cancer associated MeDEGs from other studies (16, 27). The Illumina HumanMethylation 27 BeadChip covers 27,578 CpGs distributed genome-widely but specifically in proximal promoter regions. This supports the current knowledge of how DNA methylation regulates gene expression.

Our functional clustering analysis demonstrated that among all the DEGs, aberrant hypomethylation mainly occurs in genes involved in biological functions such as formation of the cornified envelope, extracellular matrix organization, anchoring fibril formation, regulation of cell adhesion, and intermediate filament cytoskeleton organization, all have been previously suggested to play roles in the development and progression of squamous cell carcinoma. Furthermore, hypomethylated MeDEGs are also enriched in the regulation of immune

![FIGURE 3](https://example.com/image3.png)

**FIGURE 3** | PPI network analysis of MeDEGs. (A) PPI network of upregulated and hypomethylated genes. (B) PPI network of downregulated and hypermethylated genes. Each node represents a protein coded by unique genes, edges indicate interactions, and the color of edge indicates the degree of associations between proteins. Disconnected nodes in the network were hidden. Protein networks are clustered on MCL inflation parameter 2. Nodes from the same cluster are presented in the same color. Edges between different clusters are represented by dashed line.
TABLE 4 | Expression and Methylation Profiles of Eleven Potential Hub MeDEGs.

| Gene symbol | Description | Expression profile | Methylation probe ID | Distance to TSS (nt) | Methylation profile |
|-------------|-------------|--------------------|----------------------|---------------------|---------------------|
|             |             | \(\log_{2}FC\)^a  | FDR^b \(p\)-value  | GSE38532             |                     |
| CTLa4       | Cytotoxic T-lymphocyte associated protein 4 | 2.23       | 6.21E-06 | 4.83E-07 | cg038460026 | 37 | 6.47E-05 | -0.13 | 1.78E-05 | -0.31 |
| GPR29       | C-C motif chemokine receptor 6 | 2.55       | 4.23E-03 | 9.42E-04 | cg05824215 | 211 | 4.37E-15 | -0.22 | 3.44E-04 | -0.19 |
| TNFSF11     | TNF superfamily member 11 | 3.28       | 3.28E-05 | 3.19E-06 | cg08014954 | 326 | 2.58E-15 | -0.22 | 1.53E-02 | -0.18 |
| CD80        | CD80 molecule | 2.54       | 5.13E-06 | 3.89E-07 | cg02422324 | 321 | 1.51E-07 | -0.13 | 9.22E-04 | -0.14 |
| CDSN        | Corneodesmosin | 7.20       | 8.62E-16 | 7.87E-18 | cg08424223 | 35 | 5.07E-12 | -0.16 | 6.61E-05 | -0.24 |
| PI3         | Peptidase inhibitor 3 | 4.28       | 8.81E-14 | 1.23E-15 | cg03962575 | 13 | 5.81E-12 | -0.18 | / | / |
| ACTN2       | Actinin alpha 2 | -8.15      | 8.45E-27 | 8.35E-30 | cg18653982 | 410 | 2.61E-05 | 0.14 | / | / |
| MYOD1       | Myogenic differentiation 1 | -6.09      | 9.18E-13 | 1.49E-14 | cg16519321 | 84 | 1.75E-10 | 0.16 | 1.04E-02 | 0.18 |
| ISL1        | ISL LIM homeobox 1 | -3.74      | 1.09E-04 | 1.28E-05 | cg28896762 | 92 | 4.78E-14 | 0.18 | / | / |
| MYH11       | Myosin heavy chain 11 | -2.92      | 1.63E-07 | 8.18E-09 | cg17880199 | 28 | 2.57E-12 | 0.19 | 1.82E-02 | 0.17 |
| PAX7        | Paired box 7 | -6.31      | 5.45E-15 | 5.90E-17 | cg11428724 | 132 | 6.74E-22 | 0.44 | 7.52E-04 | 0.38 |

\(^{a}\)\(\log_{2}FC\): log-transformed fold change.  
\(^{b}\)FDR: Benjamini & Hochberg adjusted \(p\)-value (False discovery rate).  
\(^{c}\)TSS: transcription start site.

responses, including cellular response to bacterium, antimicrobial humoral response, and cytokine-mediated signaling pathway. This suggests the contribution of infective agents (such as previously reported HPV16 and Candida albicans) and immune dysfunction to OSCC, potentially through chronic inflammation related pathologies. For aberrantly hypermethylated DEGs, functions related to cell junction organization, regulation of membrane potential, and monocarboxylic acid metabolic process are observed to be enriched. Interestingly, multiple muscle related pathways like muscle system process, muscle structure development, muscle cell development, and muscle contraction are also found enriched. Therefore, consistent with previous reports, our study indicates that aberrant muscle function related cytoskeleton remodeling may assist the formation and progression of OSCC (28, 29). Other enriched functions related pathologies. For aberrantly hypermethylated DEGs, the PPI network of 56 upregulated MeDEGs and 170 downregulated MeDEGs were constructed using the STRING v11.0 database. Six highly connected hypomethylated genes (CTLa4, GPR29, TNFSF11, CD80, CDSN, and PI3) and five highly connected hypermethylated genes (ACTN2, MYOD1, ISL1, MYH11, and PAX7) were identified as hub genes from these networks. Hub genes are generally expected to play important roles in biological processes (30). Therefore, we further evaluated these hub genes for their correlations between observed dysregulation of DNA methylation and altered mRNA expression level in the TCGA database. Two hypomethylated hub gene (CTLa4 and CDSN) was found to be significantly upregulated, while two hypermethylated hub genes (ACTN2 and MYH11) showed obviously decreased expression level in the HNSC group. Therefore, our study identified four hub genes whose aberrant expressions were likely regulated by the mechanism of DNA methylation. Additionally, four hub genes showed important value in distinguishing the prognosis of HNSC patients. High expression levels of three individual hypomethylated hub genes (CTLa4, GPR29, and TNFSF11) presented a significant association with longer OS, while low expression of hypermethylated hub gene ISL1 was found to be clearly associated with shorter OS in HNSC patients. Therefore, these aberrantly methylated hub genes have potential diagnostic value and may become biomarkers and therapeutic targets for OSCC. Further studies with more specific OSCC datasets could provide more insights in the future.

CTLa4 (cytotoxic T-lymphocyte associated protein 4) is preferentially expressed in activated CD4+ T cells and constitutively expressed in CD4+Foxp3+ Treg cells (31, 32). It functions as a potent immune inhibitor by reducing the initiation
of T-cell activation mediated by the interactions between antigen presenting cells (APCs) and T cells. Clinically, limiting CTLA4 function by antibody blockade was proved effective as a therapeutical approach to treat various cancers by the mechanism of boosting immune response against malignant tissues (33–36). In our study, we found that CTLA4 was not only significantly upregulated in patients with HNSC, but its high expression level was also associated with patients have better long-term survival. The median survival time was about 91 months for HNSC patients with high expression levels of CTLA4, compared with a median survival time of about 43 months for those patients with low expression levels of CTLA4. The beneficial effect of high expression of CTLA4 in HNSC, at first sight, seems contradictory to the well-known immune suppressive function, in other words, tumor–promoting role of CTLA4. However, the elevated expression of CTLA4 and beneficial effect of upregulated CTLA4 have also been previously noticed in other tumors (37, 38). To address the question of whether the upregulated CTLA4 in tumor tissues was originated from the increased immune infiltration, especially the infiltration of CTLA4 abundant CD4+ T cells, we estimated the abundances of six immune cell types (B cells, CD4+ T cells, CD8+ T cells, macrophages, neutrophils, and dendritic cells) in tumor and paired adjacent tissues by re-analyzing gene expression data using TIMER web tool (Supplementary Figure S1). Although the signals of certain immune cells were found higher in our OSCC samples (neutrophil and myeloid dendritic cells) or TCGA HNSC samples (CD8+ T cell, neutrophil, and myeloid dendritic cells) compared to their corresponding NATs, the counts of CD4+ T cell infiltration were similar between tumor and paired adjacent tissues. Recently, a number of studies suggested that tumor cell-intrinsic CTLA4 might execute different functions, from inducing apoptosis to regulating cell proliferation, than that of T cells (39, 40). Here our results provide additional evidence that, depending on the context, the origin and the involvement of CTLA4 in tumorigenesis is complex.

We also identified two hypomethylated hub genes whose expression levels in tumor tissues showed positive association with the prognosis of HNSC patients, although their overall mRNA levels were not significantly changed in TCGA-HNSC samples compared to those in NATs. GPR29 (G–protein coupled receptor 29) is also known as CCR6 (CC chemokine receptor 6), a chemokine receptor that preferentially expressed by T cells, immature dendritic cells, and B cells. Its exclusive binding partner CCL20 (as known as macrophage inflammatory protein–3α, MIP-3α) is known to be steadily expressed by Th17 cells and secreted by intestinal epithelial cells as a response to local enteropathogenic infection (41). Therefore, CCL20–CCR6 axis plays crucial role in mucosal immunity. Apart from the immune-regulatory function, the elevated expression of CCR6 has been previously shown in various cancers, with complicated anti-cancer or pro-cancer potentials. TNFSF11 (tumor necrosis factor ligand superfamily member 11) is also known as receptor activator of nuclear factor kappa-B ligand (RANKL). It binds to its receptor RANK to regulate the differentiation, activation, and survival of osteoclast cells. While TNFSF11 is primarily known for its function in osteoclast regeneration, it has also been implicated in pathways like innate immune response, cell proliferation, and apoptosis. Here we report the first evidence that aberrant DNA methylation and aberrant expression level of TNFSF11 and GPR29 may play a role in the development of OSCC.

ISL1 (ISL LIM homeobox 1) is a LIM–homeodomain transcription factor. Aberrantly expressed ISL1 has been reported in various cancers, including gastric cancer, non-Hodgkin lymphoma, neuroblastoma, breast cancer, OSCC, etc. (42–44). Functionally, ISL1 has been implicated in many important biological pathways, such as tumorigenesis, cell invasion, apoptosis, and cancer immunity (42, 45). Although many studies indicated a tumor–promoting effect of elevated ISL1, Rajneesh et al. reported that hypermethylated and downregulated ISL1 was correlated with poorer survival in patients with breast cancer (46). In OSCC, Han et al. reported

**FIGURE 4** | Identification of potential hub genes from PPI networks of MeDEGs. (A) Venn diagram of highly connected genes identified by four calculation methods from PPI network of upregulated and hypomethylated genes. Six overlapping genes were identified in the intersection of all lists. (B) Venn diagram of highly connected genes identified by four calculation methods from PPI network of downregulated and hypermethylated genes. Five overlapping genes were identified in the intersection of all lists. List of calculation methods: MCC (red), MNC (green), Degree (blue), and EPC (gray).
that ISL1 is significantly decreased and could be a potential biomarker of the disease (47). Consistently, in the present study, we identify the hypermethylated ISL1 as a key change in OSCC, indicating the prognostic value of ISL1 in OSCC.

Besides CTLA4, our study adds three additional hub genes with a significant and inverse correlation between the methylation pattern in promoter region and the mRNA expression level. The first one is ACTN2, for which the decreased expression level and elevated methylation status has already been reported in OSCC by another group (48). The second is MYH11, which encodes a myosin heavy chain family protein that is selectively expressed in smooth muscle. Similar to our observation, other studies have reported the downregulation of MYH11 in OSCC (49, 50). In colorectal cancer, survival analysis showed that low expression of MYH11
was significantly associated with poor prognosis (51, 52). Although the exact mechanism remains unclear, evidence suggests that the hypermethylation regulated interaction between DNMT3B and MYH11 may contribute to the tumor progression (53). The last is CDSN, which encodes corneodesmosin that is required for corneodesmosomes to maintain properly cornified squamous epithelia. Mutations in CDSN have been described in hereditary skin conditions such as hypotrichosis simplex of the scalp (HSS) and progressive loss of scalp hair (PSD) (54–57). In this study, we are the first to indicate hypomethylated and upregulated CDSN as a key change in OSCC.

**CONCLUSION**

This study analyzed MeDEGs in OSCC using integrated multi-omics analysis and examined their related pathways and functions. From the networks of upregulated and downregulated MeDEGs, we identified the four hub genes (CTLA4, GPR29, TNFSF11, and ISL1) that are highly correlated with the OS of HNSC patients. In addition, we also verified four hub genes (CTLA4, CDSN, ACTN2, and MYH11) that their expression levels in tumor tissue might be regulated by the mechanism of DNA methylation. Therefore, this study provides a list of potential biomarkers and therapy targets for OSCC, as well as insights for unraveling the molecular mechanisms underlying this disease. However, it is worth noting the limitation of these discoveries generated from purely computational approach. Further prospective validation is warranted to characterize the clinical utilities and underlying mechanisms of these genes.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. RNA sequencing data were available in NCBI GEO
The studies involving human participants were reviewed and approved by the Ethics Committee of School of Life Sciences, Central South University. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

Conceptualization, ZW and DW. Methodology, ZW and DW. Software, ZW. Formal analysis, ZW. Investigation, ZW. Validation, ZW, XT, and DW. Resources, DW, HX, TS, and XX. Data curation, ZW. Writing—original draft preparation, ZW and DW. Writing—review and editing, ZW and DW. Visualization, ZW. Supervision, DW and KX. Project administration, ZW and DW. Funding acquisition, ZW and DW. Writing original draft preparation, review and editing, ZW and DW.

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