N6-methyladenosine-associated prognostic pseudogenes contribute to predicting immunotherapy benefits and therapeutic agents in head and neck squamous cell carcinoma

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Abstract

Rationale: N6-methyladenosine (m⁶A) is involved in critical cancerous processes. Pseudogenes play various roles in carcinogenesis and progression. However, the functional roles of m⁶A-associated pseudogenes in head and neck squamous cell carcinoma (HNSCC) are largely unknown.

Methods: We systematically analyzed the mRNA profile of 24 m⁶A regulators and 13931 pseudogenes from The Cancer Genome Atlas HNSCC dataset and ultimately identified 10 m⁶A-associated prognostic pseudogenes, which were validated in the Gene Expression Omnibus and our hospital datasets. Based on the risk score of m⁶A-associated pseudogenes, comprehensive analytical frameworks and experimental validation were implemented among pseudogene-defined low-/high-risk subtypes.

Results: Here, we found expression pattern of m⁶A-associated pseudogenes was significantly associated with infiltrating immune cell compositions, and the expression of antitumor immune response markers, including T cell exhaustion, antigen presentation, interferon, and kinase genes. The m⁶A-associated pseudogenes, which had dramatic m⁶A peaks and higher m⁶A levels, could regulate the expression of targeted immune-involved genes through miRNAs. We experimentally validate the oncogene PDIA3P1, and tumor-suppressor RRN3P3, which promote the RNA and protein expression of their targeted immune-involved genes AKT1 and EZH2 via miR-34a-5p and miR-26b-5p, respectively. Moreover, HNSCC patients in the high-risk subtype could benefit more from immune checkpoint inhibitors therapy. Furthermore, doxorubicin and topotecan were considered to hold the most promising therapeutic potential robustly in silico evidence and in vitro experiments for HNSCC patients in the high-risk subtype.

Conclusions: Our discovery revealed that the 10 m⁶A-associated prognostic pseudogenes significantly contribute to predicting immunotherapy benefits and therapeutic agents, which might bring some potential implications for both immunotherapy and chemotherapy in HNSCC.

Key words: N6-methyladenosine, Pseudogene, Immunotherapy benefits, Therapeutic agents, Head and neck squamous cell carcinoma
**Introduction**

Head and neck squamous cell carcinoma (HNSCC) accounts for over 900,000 cases and approximately 400,000 deaths per year worldwide [1], which is characterized by aetiological, phenotypic, and clinical heterogeneity. Smoking causes the rise of HNSCC in developing countries, and human papillomavirus (HPV) is emerging as an important factor in the rise of oropharyngeal tumors of non-smokers in developed countries [2]. Although advances in surgery, radiation, and chemotherapy, around half of all patients, will die of the disease. Despite risk stratification for HNSCC by age, gender, anatomic site, TNM stage, HPV status, laterality, and histological characteristics, numerous molecular biomarkers that have been investigated have limited clinical utility. Therefore, seeking new promising prognostic biomarkers at the intrinsic molecular level is necessary, which contributes to identifying high-risk subtypes and making precise therapeutic strategies.

Except for surgery, radiotherapy, chemotherapy, and targeted therapy based on molecular subtypes and the TNM stage of HNSCC, immunotherapy is an emerging treatment modality due to the promising therapeutic effect of selective immune checkpoint inhibitors (ICI), including monoclonal antibodies against programmed death 1 (PD-1), programmed death-ligand 1 (PD-L1) and cytotoxic T lymphocyte-associated protein-4 axes (CTLA-4) [3-5]. These expression levels of PD-1, PD-L1, and CTLA-4 in tumor tissue are currently used as predictive markers for immune response. And these predictive results were still not satisfactory, indicating that immune modulation in tumor tissue is a complex process and needs many more functional predictors [6]. Thus, it is necessary to identify new robust predictive biomarkers for the immune response process when executing clinical trials of immunotherapy.

Pseudogenes played a pivotal role in many human diseases, including tumorigenesis and tumor progression [7, 8]. As non-functional homologs of protein-coding genes, Pseudogenes are commonly caused by the accumulation of multiple nonsense mutations within genes. Based on the unique biogenesis mechanisms, pseudogenes are divided into three categories: unitary pseudogenes, unprocessed pseudogenes, and processed pseudogenes [9]. Although pseudogenes were once considered “genetic fossils” due to their lack of protein-coding ability or cellular gene expression, increasing evidence indicates that some pseudogenes have been identified to participate in multiple biological functions and regulate their parental transcripts by acting as competitive endogenous RNAs (ceRNA) [10]. What’s more, dysregulation of pseudogenes is associated with diseases, which was immensely attributed to the discovery that the PTENP1 [11] and BRAFPI [12] could upregulate their corresponding parental gene PTEN and BRAF via the ceRNA mechanism.

N6-methyladenosine (m^6A) RNA modification is pervasively a reversible internal RNA modification in most kinds of RNAs covering mRNAs and long noncoding RNAs (lncRNAs), which has been confirmed to participate in regulating RNA transcription, processing, and so on [13]. It is installed by m^6A methyltransferases (termed as “writers”) complicated with METTL3 as the catalytic subunit and reversed by demethylases (termed as “erasers”) FTO and ALKBH5 [14, 15]. In addition, m^6A can be specifically regulated through a series of RNA binding proteins (termed as “readers”) and co-transcriptionally through a variety of transcription factors [14, 15]. As m^6A readers, YTH domain-containing proteins can specifically read the m^6A and regulate diverse post-transcriptional processes of host mRNAs [14, 15]. Accumulating evidence demonstrated that critical roles of m^6A have been identified in the development of multiple cancers [16].

According to the genome-wide profiling of m^6A, pseudogenes are also modified by m^6A in non-tumor cell lines such as GM12878 and H1 [17]. Our previous research found that pseudogenes and m^6A were significantly correlated with host antitumor immune response and might serve as potential biomarkers for immunotherapy in breast cancer [18, 19]. Furthermore, we investigated the function of the m^6A sites on pseudogenes in non-tumor cell lines, which indicated a new evolutionary role of m^6A in cleaning up the unnecessary processed pseudogenes to mitigate their interference of expression of cognate protein-coding genes [20]. However, little is known about what the function of m^6A-associated pseudogenes is and whether they can predict immunotherapy benefits and therapeutic agents in HNSCC.

In this study, as shown in the workflow (Figure S1), we systematically analyzed the RNA sequencing data of HNSCC patients and eventually identified and validated 10 m^6A-associated prognostic pseudogenes, which were independently prognostic indicators in HNSCC patients. We further found a close association between the m^6A-associated prognostic pseudogenes and the antitumor immune response from different aspects. Our convergent evidence showed that HNSCC patients could benefit more from immune checkpoint inhibitor therapy and the most promising potential therapeutic agents (doxorubicin and topotecan) for HNSCC patients. Our study indicated the 10 m^6A-related prognostic pseudogenes could
predict immunotherapy benefits and potential therapeutic agents in HNSCC.

Results

Identification and validation of 10 m6A-associated prognostic pseudogenes in TCGA and GEO datasets

To study the functions of m6A-related pseudogenes in HNSSC, we first ask whether m6A regulators are expressed differently between HNSSC tissues with normal tissues. We collected 24 m6A regulators from studies that have been publicly reported (List S1) and explored their expression between 500 tumor tissues and 44 normal tissues in 500 HNSSC patients with different clinicopathological features. Our research results showed that the expressions of m6A regulators differed enormously across subtypes (Figure 1A). For example, compared to normal tissues, the expressions of METTL3, METTL14, WTAP, VIRMA, ZCCHC4, FTO, ALKBH5, YTHDC1, YTHDF1, YTHDF2, YTHDF3, HNRNPC, HNRNP D, RBM15, RBM27, ZC3H7B, YWHAG, CAPRIN1 and PCIF1 increased significantly in tumor tissues, while the expressions of YTHDC2, ZC3H7B, TRA2A, GNL3, and MS12 decreased (Figure S2). Besides, the expression pattern of m6A regulators was inconsistent among different clinicopathological features, which indicated the crucial but complicated roles of m6A regulators in tumorigenesis and the development of HNSSC (Figure 1A). On account, genes do not perform a function in isolation, and increasing evidence has manifested that collaboration among writers, erasers, and readers exists in the context of cancer. Thus the co-expression among 24 m6A regulators by Pearson correlation analysis was investigated. We found not only that m6A regulators within the same functional class showed significant co-expression and highly correlated expression patterns, but that a close correlation also existed among writers, erasers, and readers (Figure 1B). For instance, the reader YTHDF2 was significantly correlated with writers, such as METTL14 (Figure 1B, R = 0.66 and P-value < 2.2e-16). Through the String database, we found that these 5 writers, 2 erasers, and 17 readers interacted with each other frequently in protein-protein interaction networks (Figure 1C). The high degree of co-expression and strong interaction among 24 m6A regulators suggested that m6A RNA modification played an important and complex role in HNSSC.

Then, a total list of 13931 pseudogenes was compiled from GENCODE, Vega, and psiCube databases (List S2), of which 6218 pseudogenes were obtainable in TCGA HNSSC datasets and thus used in the subsequent analyses. Firstly, we took advantage of the Pearson correlation analysis of expression between candidate pseudogene with 24 m6A regulators to filter a list of 2682 one-to-one pairs of pseudogenes and m6A regulators at | R | ≥ 0.3 and P < 0.05 (Table S1), including 842 pseudogenes with different numbers of m6A regulators. Secondly, through the univariate Cox proportional hazard regression, 53 pseudogenes were screened at P < 0.05 (Table S2). Thirdly, based on the second step analysis above, the multivariate Cox proportional hazard regression was implemented. Using the above three methods, overlapping 10 m6A-associated prognostic pseudogenes (PDIA3P1, LDHAP4, LDHAP7, EEF1AIP6, EEF1AIP11, SDHAP1, SDHAP3, DDX12P, CLUH3P, RRN3P3) were ultimately identified (Figure 1D).

To further interpret the molecular mechanisms by which m6A regulators and pseudogenes are involved in cancer, we examined the Pearson correlation between the expression of individual m6A regulators and the 10 m6A-associated prognostic pseudogenes. We found that there were significantly higher correlations between YTHDC1 and WTAP with 10 prognostic pseudogenes, which implies that YTHDC1 and WTAP play important roles in the function of m6A modification on these 10 pseudogenes (Figure 1E). Forest plot of 10 m6A-associated prognostic pseudogenes showed that the hazard ratio of PDIA3P1, LDHAP4, LDHAP7, EEF1AIP6, EEF1AIP11 was greater than 1, which indicated that these 5 pseudogenes may be factors with poor prognosis, while the hazard ratio of SDHAP1, SDHAP3, DDX12P, CLUH3P, RRN3P3 was less than 1, hinting these 5 pseudogenes may be factors with good prognosis (Figure 1F). What’s more, these 10 m6A-associated prognostic pseudogenes were further verified in the GEO GSE65858 dataset (Figure S3A). To raise the predictive effect of pseudogenes in the clinical outcomes of HNSSC, we applied the least absolute shrinkage and selection operator (LASSO) Cox regression algorithm to the 10 m6A-associated prognostic pseudogenes and established a risk signature based on the minimum criteria using TCGA HNSSC data as the training set (Figure S3B-C) and GEO GSE65858 data as the validation set. Based on the coefficients of the 10 pseudogenes (Table S3), the risk score was calculated on account of the survival risk score model formula. Then, according to the median risk score, the HNSSC patients were dichotomized into low or high-risk groups. We found that patients in the high-risk subtype displayed remarkably shorter overall survival than those in the low-risk subtype (TCGA dataset, log-rank test, P < 0.001, Figure 1G; GEO...
dataset, log-rank test, \( P = 0.035 \), Figure 1H). The patients in high-risk group were significantly associated with female (\( P = 0.008 \)), higher Pathologic T stage (\( P = 0.003 \)) and higher Pathologic M stage (\( P = 0.040 \)) (Table S4). In addition, to better understand whether m\( ^6 \)A-associated pseudogenes be able to effectively predict the prognosis of HNSCC patients, the receiver operating characteristic curve (ROC) curve analysis was performed. Results showed that the risk score based on m\( ^6 \)A-associated pseudogenes was a good predictor of survival rates with an area under the curve (AUC) value of 0.722 in the training set (Figure 1I) and 0.773 in the validation set (Figure 1J).

Figure 1. Identification and validation of 10 m\( ^6 \)A-associated prognostic pseudogenes in TCGA and GEO datasets. (A) Heatmap showing the expression of 24 m\( ^6 \)A regulators across different clinicopathological features between normal tissues and tumor tissues. The expression value between normal and tumor tissues were compared through the Wilcoxon test. ns denotes no significance, * denotes \( P < 0.05 \), ** denotes \( P < 0.01 \), *** denotes \( P < 0.001 \) and **** denotes \( P < 0.0001 \). (B) Triangle heatmap of Pearson correlation among the expression of m\( ^6 \)A regulators. The scatter plot shows the correlation of the expressions between \textit{METTL14} and \textit{YTHDF2}. (C) Circular arc diagram of the protein-protein interactions among m\( ^6 \)A regulators. (D) Venn diagram revealing the overlapping m\( ^6 \)A-associated prognostic pseudogenes screened by three methods. (E) Heatmap showing Pearson correlation of the expression between 24
Screened 10 m6A-associated pseudogenes are an independent prognostic factor in HNSCC patients

To intuitively understand the prognostic effect of m6A-associated pseudogenes, the distribution of the risk scores based on pseudogenes, overall survival of HNSCC, and corresponding pseudogene expression profiles in the TCGA dataset were displayed (Figure 2A). The composite plot indicated that SDHAP1, SDHAP3, DDX12P, CLUHP3, and RRN3P3 demonstrated high expressions in the low-risk subtype, which were categorized as tumor-suppressor pseudogenes in our study. However, the residual pseudogenes (PDIA3P1, LDHAP4, LDHAP7, EEF1A1P6, and EEF1A1P11) displayed high expressions in the high-risk subtype and therefore were classified as oncogene pseudogenes in our study (Figure 2A). Besides, our study also revealed that the risk score and prognostic pseudogenes were significantly related to different clinicopathological features of HNSCC patients. Compared to HPV negative, patients with HPV positive had significantly higher expressions of oncogene pseudogenes (SDHAP1, SDHAP3, DDX12P, CLUHP3, and RRN3P3), but there was no prominent difference in the expressions of tumor-suppressor pseudogenes (PDIA3P1, LDHAP4, LDHAP7, EEF1A1P6, and EEF1A1P11) (Figure S4A). In addition, patients with higher grades (such as G4) had significantly higher expressions of PDIA3P1, EEF1A1P6, EEF1A1P11, SDHAP1, SDHAP3, DDX12P, CLUHP3, and RRN3P3, but lower expressions of LDHAP4 and LDHAP7 than those with lower grades (Figure S4B). On the contrary, there were no striking differences in the expressions of the 10 m6A-associated prognostic pseudogenes among patients with different TNM (Figure S5A-C).

On account of a series of factors, such as HPV status and different clinical features, which can affect the prognosis of HNSCC patients, we couldn't help asking whether m6A-associated prognostic pseudogenes are independent prognostic factors. To address the doubt, univariate and multivariate Cox regression analyses are simultaneously executed. Then our findings demonstrated that risk score, age, Gender, pathology T stage, and pathology N stage were all correlated with the overall survival by univariate Cox regression analysis. What’s more, when including these factors in the multivariate Cox regression, our analysis revealed that risk score ($P < 0.001$), age ($P = 0.005$), and pathology N stage ($P = 0.007$) remained closely associated with the prognosis (Figure 2B), which proved that the risk score derived from these 10 m6A-associated pseudogenes was able to independently predict the clinical outcome in HNSCC patients.

To further demonstrate the prospect of m6A-associated pseudogene clinical application, 32 HNSCC tissues (16 oral squamous cell carcinoma tissues and 16 thyroid cancer tissues) from the Tumor Resource Bank of Sun Yat-sen University Cancer Center were used to verify the relative RNA expression of these 10 m6A-associated by qPCR. Results show oncogene pseudogenes (PDIA3P1, LDHAP4, LDHAP7, EEF1A1P6, and EEF1A1P11) displayed high expressions in the low-OS subtype and that tumor-suppressor pseudogenes (SDHAP1, SDHAP3, DDX12P, CLUHP3, and RRN3P3) demonstrated high expressions in the high-OS subtype (Figure 2C-D), which was consistent with our above results (Figure 2A).

Since the above screened 10 m6A-associated pseudogenes are an independent prognostic factor in HNSCC patients, we would expect to see a higher expression of oncogene pseudogenes in companions with worse survival outcomes, as well as higher expression of tumor-suppressor pseudogenes in companions with better survival outcomes. To test this, we examine the associations between the expression of m6A regulators and m6A-associated pseudogenes with overall survival in HNSCC patients from the TCGA dataset. We found that patients with high expression of three m6A regulators in TCGA dataset (Figure 2E) and nine m6A regulators in the GEO dataset (Figure S6A) had a significantly worse outcome than those with low expression, suggesting that high expression of m6A regulators might have accelerated the progression of the tumor. It’s worth noting that WTAP, YTHDC1, and YWHAG represented significant differences both in TCGA and in the GEO dataset, implying the critical function of m6A regulators in HNSCC, in line with the pivotal roles of YTHDC1 and WTAP in the previous findings (Figure 1E). Patients with high expression of oncogene pseudogenes had strikingly shorter survival than those with low expression (log-rank test; PDIA3P1, $P = 0.046$; LDHAP4, $P = 0.004$; LDHAP7, $P = 0.014$; EEF1A1P6, $P = 0.025$; EEF1A1P11, $P = 0.007$; Figure 2F), manifesting that high expression of oncogene pseudogenes might correlate with high malignancy of the tumor. In contrast, patients with high expression of tumor-suppressor pseudogenes had significantly
better outcome than those with low expression (log-rank test; SDHAP1, \( P = 0.023 \); SDHAP3, \( P = 0.036 \); DDX12P, \( P = 0.007 \); CLUHP3, \( P = 0.017 \); RRN3P3, \( P = 0.014 \); Figure 2G), which explained that high expression of tumor-suppressor pseudogenes was associated with high benignity of tumor. What’s more, this conclusion has been further verified in the GEO dataset (Figure S6B-C).

| Clinical features | No. of Patients (%) | Hazard Ratio (95% CI) | P-value |
|-------------------|---------------------|-----------------------|---------|
| Risk score       | 500 (100)           | 8.05 (3.10 to 13.0)   | \(< 0.001\) |
| Age               | 500 (100)           | 1.00 (1.00 to 1.00)   | 0.002   |
| Gender            | 500 (100)           | 0.78 (0.56 to 1.00)   | 0.041   |
| Pathology_T Stage | 500 (100)           | 1.20 (1.10 to 1.30)   | 0.003   |
| Pathology_N Stage | 500 (100)           | 1.20 (1.10 to 1.30)   | 0.001   |
| Histology_grade   | 500 (100)           | 1.99 (1.99 to 2.10)   | 0.089   |
| Laterality        | 500 (100)           | 1.34 (0.93 to 2.30)   | 0.230   |
| HazardRatio       | 500 (100)           | 1.06 (0.93 to 2.30)   | 0.460   |

Figure 2. Screened 10 m6A-associated pseudogenes are an independent prognostic factor in HNSCC patients. (A) The composite plot of the distribution of risk score, vital status, and the expression pattern of 10 m6A-associated prognostic pseudogenes in 500 HNSCC patients. The risk scores are arranged in ascending order from left to right. (B) Forest plot of univariate and multivariate Cox regression analyses of the association between clinicopathological factors (including the risk score) and overall survival (OS) of HNSCC patients. ns denotes no significance, * denotes \( P < 0.05 \), ** denotes \( P < 0.01 \), *** denotes \( P < 0.001 \) and **** denotes \( P < 0.0001 \). (C) The heatmap of the relative RNA expression (by qPCR) of 10 m6A-associated pseudogenes and OS of the corresponding patient in 32 HNSCC tissues from the Tumor Resource Bank of Sun Yat-sen University Cancer Center. (D) Barplot of the mean relative RNA expression of oncogene pseudogenes in 16 low-OS subtype tissues and tumor-suppressor pseudogenes in 16 high-OS subtype tissues. (E) Kaplan-Meier curves of association between the expression levels of m6A regulators and overall survival in patients with HNSCC from TCGA dataset. (F) Kaplan-Meier curves of association between the expression levels of oncogene pseudogenes and overall survival in patients with HNSCC from TCGA dataset. (G) Kaplan-Meier curves of association between the expression levels of tumor-suppressor pseudogenes and overall survival in patients with HNSCC from TCGA dataset.
To better show, the correlation between the expression of pseudogenes and the clinical outcome of patients, nomograms for evaluating the risk of HNSCC were developed for HNSCC patients based on risk factors identified by the multivariate logistic regression analysis. According to the results of the multivariate analysis, we included risk scores based on m6A-associated pseudogenes and some important clinical features to build a nomogram for predicting the 1-year, 3-year, and 5-year prognoses of HNSCC patients. Although our analysis did not identify Gender, Pathology_T_Stage, Pathology_M_Stage, Hpv_status, Laterality, and Histologic_grade as independent predictive factors, we took into account these variables in the nomogram. The total point was calculated according to the Oxford classification recommendations and principle, which was further transformed into probability (see the bottom scale) (Figure S7A). The calibration curves expressed a good consistency between the nomogram-predicted progression probability and the actual progression probability. And the calibration curves also showed our nomogram performing well in predicting the 1-year, 3-year, and 5-year prognosis of HNSCC patients (Figure S7B). The above evidence strongly proved that expressions of 10 m6A-associated pseudogenes are significantly associated with survival outcomes of HNSCC patients.

**The expression pattern of m6A-associated prognostic pseudogenes was significantly correlated with antitumor immune response**

To investigate and make clear the correlation between the expression pattern of m6A-associated pseudogenes and antitumor immune response in HNSCC, we compared the expressions of m6A-associated pseudogenes, estimated the immune cell infiltration through CIBERSORT, and assessed the expressions of T cell exhausted, antigen presentation, interferon activity, kinase, cytolytic and integrin genes in tumor tissues between low- and high-risk subtypes. The expression pattern of the 10 m6A-associated pseudogenes across low- and high-risk subtypes indicated that the low-risk subtype had lower expressions of oncogene pseudogenes as well as higher expressions of tumor-suppressor pseudogenes, while the high-risk subtype showed the opposite trends (Figure 3A). Studying the important role of immune-infiltrating cells in tumor immunity helps us to specify the best immunotherapy regimen. Through assessing the immune cell infiltration, we found that the low-risk subtype had a prominently higher number of tumor-infiltrating B cells, CD8+ T cells (as known as cytotoxic T cells), helper T cells, regulatory T cells, and a lower fraction of activated natural killer cells, M1 macrophage cells, M2 macrophage cells than high-risk (Figure 3B), suggesting an enhanced immunosurveillance in the low-risk subtype. Of note, CD4+ T cells, a renowned member of the helper T cell population, showed a significantly higher quantity in high-risk than in the low-risk subtype (Figure 3B). Evaluating the expression level of immuno-modulatory genes in tumor immunity contributes to helping us to target specific immunotherapy targets. Then comparisons of expressions of immuno-modulatory genes were conducted. In terms of T-cell exhausted genes, the low-risk subtype was dramatically associated with higher expressions of PD-1 than the high-risk subtype (Figure 3C), which is a key gene of T-cell exhaustion markers. However, the high-risk subtype had remarkably higher expressions of PD-L1 and PD-L2, which also account low-risk subtype having a longer Survival period (Figure 3C). Concerning antigen-presenting genes, our findings revealed that high-risk had significantly higher expressions of HLA-A, HLA-B, HLA-C, HLA-E, TAP1, and B2M than low-risk (Figure 3D), which can activate cytotoxic T cells. Regarding interferon activity genes, we surprisingly found that the expressions of CXCL9, CD24, and STAT1 were also significantly higher in high-risk than the low-risk subtype, while the expressions of CD27 and IRF3 show the opposite trend (Figure 3E). In addition, the low-risk subtype was also associated with higher expressions of AKT1, E2F2, MECP2, HOXA1, and HOXA10 (Figure 3F), several important regulatory genes for kinase activity. Besides, there is a significant difference in the expressions of cytolytic genes (CYTH1, CYTH2, CYTH3) and integrin genes (ITGA family genes and ITGB family genes) (Figure S8A-C) between low- and high-risk subtype.

In consideration of the large amounts of m6A-associated prognostic pseudogenes, we performed consensus clustering of the 10 m6A-associated prognostic pseudogenes through dimensionality reduction analysis in the subsequent study. Based on the similarity of pseudogenes expression, when clustering stability increased from k = 2 to 10, k = 2 seemed to be the optimal selection in the TCGA dataset (Figure 3G-I). Thus, we divided the 500 HNSCC patients into two subgroups by making 2 the k value, namely, P1 (Patients subgroup 1) and P2 (Patients subgroup 2). We found that patients in the P1 subgroup had a significantly better outcome than those in the P2 subgroup by Kaplan-Meier analysis (median overall survival 7.5 years vs. 3.5 years, log-rank test, P = 0.008, Figure 3J). And our results indicated that the P1 subgroup had higher expressions of tumor-suppressor pseudogenes and lower expressions of oncogene pseudogenes, while...
the P2 subgroup showed the contrary tendency (Figure 3K), which was highly consistent with the low-risk and high-risk subgroups. Specifically, compared with P2, the P1 subgroup had significantly higher expressions of 5 tumor-suppressor pseudogenes and significantly lower expressions of 5 oncogene pseudogenes (Figure S9A).

Figure 3. Expression pattern of m6A-associated prognostic pseudogenes was significantly correlated with antitumor immune response. (A) Boxplot revealing comparisons of expression levels of oncogenes (PDIA3P1, LDHAP4, LDHAP7, EEF1A1P6, EEF1A1P11) and tumor-suppressor genes (SDHAP1, SDHAP3, DDX12P, CLUHP3, RRN3P3) between low-risk and high-risk subtypes. (B) Boxplot showing comparisons of cell composition fraction of B cells, CD8+ T cells, CD4+ T cells, helper T cells, regulatory T cells, activated natural killer (NK) cells, M0 macrophages, M1 macrophages, M2 macrophages, monocytes, mast cells, and activated dendritic cells between low-risk and high-risk subtypes. (C) Boxplot displaying comparisons of expressions of PD-L1, PD-L2, LAG3, TIGIT, and CTLA4 between low-risk and high-risk subtypes. (D) Boxplot manifesting comparisons of expressions of HLA-A, HLA-B, HLA-C, HLA-E, TAP1, and TAP2 between low-risk and high-risk subtypes. (E) Boxplot comparing the expressions of kinase genes (AKT1, FOXM1, E2F2, MECP2, HOXA1, HOXA10) between low-risk and high-risk subtypes. The P-value of comparisons between the two subtypes was calculated through the Wilcoxon test. Purple represents P-value < 0.05. (G) Plot of consensus clustering cumulative distribution function (CDF) for k = 2 to 10. (H) Line graph showing the relative change in area under CDF curve for k = 2 to 10. (I) The plot of consensus clustering of 500 HNSCC with k = 2, indicates P1/2 subgroups were identified by consensus clustering of the 10 m6A-associated pseudogenes in the TCGA dataset. (J) Kaplan-Meier survival curve reveals that the HNSCC patients in the P1 subgroup displayed significantly longer overall survival than those in the P2 subgroup (P = 0.008). (K) Heatmap displaying the expression pattern of m6A-associated pseudogenes between P1 and P2 subgroups.

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The correlation results between the expression pattern of m6A-associated pseudogenes and immune response across P1 and P2 subgroups were also consistent with the low-/high-risk subtype (Figure S9B-I). Besides, the P1 subgroup was significantly associated with few HPV infections ($P = 0.009$) and lower Pathologic M stage ($P = 0.047$) than P2 (Table S5). By comparing the difference between P1/P2 with the high/low-risk subtype, we found that the low-risk subtype contains 222 patients in the P1 and 28 patients in the P2 subgroup, and the high-risk subtype contains 95 patients in the P1 and 155 patients in the P2 subgroup. The $P$-value of the correlation between the low/high-risk subtype and Pathologic T stage, Pathologic M stage, and Vital status was more significant than the P1/P2 subgroup (Table S4 and S5). These observations were highly in line with those presented a higher fraction of active immune cell infiltrations. Therefore, convergent evidence supported that m6A-associated prognostic pseudogenes played an important role in the antitumor response, which might serve as potential biomarkers for immunotherapy.

**m6A-associated pseudogenes can regulate targeted immune-involved genes via miRNAs**

To illuminate the potential mechanism of how m6A-associated pseudogenes regulated anti-tumor immune response, we constructed a pseudogene-miRNA-targeted immune-involved gene regulatory network. Underlying miRNAs binding to the 10 pseudogenes were identified using the dreamBase and miRNA target genes were extracted by the miRTarBase, which were verified by at least two strong experiments (Table S6). We calculated expression correlations between each pseudogene and its miRNA target genes using Pearson correlation analysis. Target genes with $|R| \geq 0.3$ and $P < 0.05$ were picked up (Table S7). Ultimately, 4 tumor-suppressor pseudogenes (SDHAP1, SDHAP3, DDX12P, RRN3P3) together with 26 microRNAs and 138 targeted genes, and 4 oncogene pseudogenes (PDIA3P1, LDHAP7, EEF1A1P6, EEF1A1P11) together with 28 microRNAs and 58 targeted genes, were used to build the pseudogene-miRNA-target gene regulatory networks and visualized using Sankey diagram (Figure 4A; Figure S11A-F). Oncogene pseudogene PDIA3P1, acting as a decoy of hsa-miR-34a-5p, hsa-miR-10b-5p, hsa-miR-199a-3p, and hsa-miR-19a-3p downregulated the expression of AKT1 and then downregulated the infiltrations of some immune cells (including CD4+ T and M1 macrophages cells) through calcium signaling and signal transduction (Figure 4A, left panel) in the low-risk subtype. Tumor-suppressor pseudogene RRN3P3 upregulated the expression of EZH2 by competitively binding hsa-miR-26a-5p and hsa-miR-26b-5p, which explained the higher expression of EZH2 in the low-risk subtype (Figure 4A, right panel). Other pseudogenes also played regulatory roles in signaling pathways coving oncogenic transformation, cell proliferation, and cell migration as ceRNAs (Figure S11A-F). The pseudogene-miRNA-targeted immune-involved gene regulatory networks partially clarified the mechanism of how pseudogenes were involved in regulating the immune response in HNSCC.

To experimentally test whether the m6A-associated pseudogenes affect the expression of their targeted immune-involved genes, we selected representative oncogene pseudogene PDIA3P1 and tumor-suppressor pseudogene RRN3P3 for further validation in ARO and Tca8113 cell lines, which were two representatives HNSCC cell lines. First, we tested whether oncogene pseudogene PDIA3P1 can regulate its targeted AKT1. We found knockdown of PDIA3P1 using siRNA significantly down-regulated gene expression as well as the protein expression of AKT1 (Figure 4B-D), consistent with our observation that their gene expressions were positively correlated (Figure 3A and 3F). To test whether PDIA3P1 modules AKT1 via ceRNA mechanism, we first verified that knockdown of PDIA3P1 was able to promote the degradation of AKT1 (Figure 4E). Then
we selected miR-34a-5p for experimental validation. Our findings revealed that inhibition of miR-34a-5p significantly increased the mRNA and protein expression of both PDIA3P1 and AKT1, indicating that miR-34a-5p targets both PDIA3P1 and AKT1 (Figure 4F-G). Next, we tested whether Tumor-suppressor pseudogene RRN3P3 can regulate its targeted EZH2. We found knockdown of RRN3P3 using siRNA significantly down-regulated gene expression as well as the protein expression of EZH2 (Figure 4H-J), in line with our observation that their gene expressions were positively correlated (Figure 3A and 3F). Then to test whether RRN3P3 modules EZH2 via ceRNA mechanism, we first confirmed that knockdown of RRN3P3 could promote the degradation of EZH2 (Figure 4K). Then we selected miR-26b-5p for experimental validation. We found inhibition of miR-26b-5p significantly increased the mRNA and protein expression of both RRN3P3 and EZH2, indicating that miR-26b-5p targets both RRN3P3 and EZH2 (Figure 4L and 4M).

To better study the function of m6A-associated pseudogenes, we utilized m6A-LAIC-seq to quantify the m6A levels on pseudogenes on a transcriptome-wide scale and used m6A seq technology to identify m6A peaks and m6A sites on pseudogenes. Here, we took advantage of our previously published m6A levels of pseudogenes [20], which were calculated from m6A-LAIC-seq data [17], to study the m6A levels of pseudogenes. And we used our previously published m6A peaks of pseudogenes, which were calculated from the m6A-seq data [21], to test whether the elevation of m6A levels on pseudogenes was due to de novo formation of m6A peaks on pseudogenes or promoted methylation on ancestral m6A sites. As shown in Figure 4N, m6A-seq identified two m6A peaks on oncogene pseudogene PDIA3P1 (left panel) and one m6A peak on tumor-suppressor pseudogene RRN3P3 (right panel), and the m6A-LAIC-seq data showed that full-length RNAs of pseudogenes were significantly enriched in m6A positive fraction. Similar results were also observed for other m6A-associated pseudogenes. The above results suggest that the m6A levels of m6A-associated pseudogenes are increased attributed to the de novo formation of m6A sites. Our previous discovery revealed that m6A on processed pseudogenes plays a novel evolutionary role in removing the unnecessary processed pseudogenes to mitigate their interference in the expression of protein-coding genes [20]. To determine the m6A methylation levels at each site on pseudogenes, we used absolute quantification in the m6A modification assay, which is a new and ultrasensitive quantitation assay for the accurate determination of m6A at single-nucleotide resolution. We found that METTL3 could significantly methylate PDIA3P1 mRNA on the Chr1:146650342 site and RRN3P3 mRNA on the Chr16:22431201 site (Figure 4O).

**HNSCC patients in the high-risk subtype could benefit more from immune checkpoint inhibitors therapy**

To elucidate the benefits situation of ICI therapy in different low-/high-risk subtypes, we then used tumor immune dysfunction and exclusion (TIDE) to evaluate the potential clinical efficacy of immunotherapy in different low-/high-risk subtypes. A higher TIDE prediction score indicated a higher potential for immune evasion, which represented that the patients were less likely to benefit from ICI therapy. Here, we found that the high-risk subtype had a lower TIDE score than the low-risk subtype, suggesting that high-risk patients could benefit more from ICI therapy than low-risk patients (Figure 5A). Since a higher TIDE prediction score was associated with a worse outcome, the high-risk subtype with a low TIDE score might have a better prognosis than the low-risk subtype with a high TIDE score from immune checkpoint inhibitors therapy, which was consistent with that PD-L1 expression was positively correlated with the efficacy of immune checkpoint inhibitors. Moreover, we found that the low-risk subtype had a higher microsatellite instability (MSI) score, T cell exclusion, and tumor-associated macrophages M2 (TAM_M2) (Figure 5A), but there was no difference in T cell dysfunction (Figure 5B), myeloid-derived suppressor cell (MDSC) and cancer-associated fibroblasts (CAF) (Figure 5C) between the two subtypes. In addition, the tumor inflammation signature (TIS) score based on the 18 signature genes (List S6) was calculated as an average value of the log2-scale normalized expression. And we also found that the high-risk subtype with a high TIS score than the low-risk subtype with a low TIS score, which suggested that the high-risk subtype might have a better outcome by estimating the TIS score (Figure 5D; Table S8). Besides, we evaluated the purity of the tumor through several methods including ABSOLUTE, IHC, CPE, and ESTIMATE, convergently suggesting that the low-risk subtype might have a higher purity than the high-risk subtype (Figure 5E), indicating a higher percentage of cancer cells, which might explain the reasons for the worse prognosis of patients in low-risk subtype after ICI therapy.
Figure 4. m6A-associated pseudogene can regulate targeted immune-involved genes via miRNAs. (A) Sankey showing pseudogenes together with binding miRNAs and target genes with $| R | \geq 0.3$ and $P < 0.05$ were used to construct the pseudogene-miRNA-target gene regulatory networks by subtypes of oncogene pseudogene PDIA3P1 and tumor-suppressor pseudogene RRN3P3. The column on the left represented pseudogenes, which are located at the cores of the networks. The column in the middle and the column on the right stand for binding miRNAs and target genes, respectively. (B-G) Experimental validation of PDIA3P1 affects the expression of AKT1 after PDIA3P1 knockdown using siRNA. (B) Relative gene expression of PDIA3P1 after PDIA3P1 knockdown using siRNA. (C) Relative gene expression of AKT1 after PDIA3P1 knockdown using siRNA. (D) Western blot comparing the protein levels of AKT1 in control and PDIA3P1 knockdown cells. (E) The relative expression of AKT1 at different time points after transcription inhibition in control and PDIA3P1 knockdown cells respectively. Error bars represent standard errors. (F) The relative expression of PDIA3P1 and AKT1 after adding control inhibitor versus miR-34a-5p inhibitor. (G) Western blot comparing the protein levels of AKT1 after adding control inhibitor versus miR-34a-5p inhibitor. (H-M) Experimental validation of RRN3P3 affects the expression of EZH2 after RRN3P3 knockdown using siRNA. (H) Relative gene expression of RRN3P3 after RRN3P3 knockdown using siRNA. (I) Relative gene expression of EZH2 after RRN3P3 knockdown using siRNA. (J) The relative expression of EZH2 at different time points after transcription inhibition in control and RRN3P3 knockdown cells respectively. Error bars represent standard errors. (K) Western blot comparing the protein
levels of EZH2 in control and RRN3P3 knockdown cells. (L) The relative expression of RRN3P3 and EZH2 after adding control inhibitor versus miR-26b-5p inhibitor. (M) Western blot comparing the protein levels of EZH2 after adding control inhibitor versus miR-26b-5p inhibitor. (N) UCSC genome browser tracks m6A-seq and m6A-LAIC-seq data indicating m6A peaks and m6A levels of oncogene pseudogene PDIA3P1 and tumor-suppressor pseudogene RRN3P3. Read-coverage tracks of input, m6A-negative, and m6A-positive fractions of m6A-LAIC-seq shown along with overlay tracks of m6A-seq (cyan for input and red for RIP; predicted m6A sites in m6A peaks are indicated by arrows). Read coverage (y-axis) of m6A negative and m6A positive are normalized as previously described [17] to reflect the calculated m6A levels (i.e., equal signals in m6A positive (eluate) versus m6A negative (supernatant) suggest m6A levels of 50%), while input and IP tracks of m6A-seq are shown for optimal viewing at the top panel. * P < 0.05; ** P < 0.01; *** P < 0.001 (two-tailed t-test). (O) The m6A methylation level of the pseudogenes at specific modification sites (Chr1:146650342, GG(m6A)CA on PDIA3P1; Chr16:22431201, GG(m6A)CG on RRN3P3) using SELECT in control and METTL3 knockdown ARO and Tca8113 cell.

Figure 5. HNSCC patients in the high-risk subtype could benefit more from immune checkpoint inhibitors therapy. (A) Violin illustration showing comparisons of TIDE, MSI_Expr_Sig, Exclusion, and TAM_M2 values in different low-/high-risk subtypes, which represent TIDE, MSI, T cell exclusion dysfunction, and TAM score respectively. (B)Violin illustration comparing dysfunction value in different low-/high-risk subtypes, which stand for T cell dysfunction score. (C)Violin illustration comparing MDSC and CAF values in different low-/high-risk subtypes, which stand for MDSC and CAF scores respectively. (D)Violin illustration comparing TIS score in different low-/high-risk subtypes, which stand for TIS score. (E) Violin illustration indicating comparisons of purity calculated by four methods (ABSOLUTE, IHC, CPE, and ESTIMATE) in different low-/high-risk subtypes. The P-value of comparisons between the two subtypes was calculated through the Wilcoxon test. (F) Kaplan-Meier survival curve revealed that the UC patients from the Snyder cohort in the low-risk subtype displayed significantly longer overall survival than those in the high-risk subtype (P = 0.025). (G) Kaplan-Meier survival curve revealed that the UC patients from the Mariathasan cohort in the low-risk subtype displayed significantly longer overall survival than those in the high-risk subtype (P = 0.032). (H) The ROC curve shows AUC for the predictive value of m6A-associated pseudogene in the UC cohort from Snyder. (I) The ROC curve shows AUC for the predictive value of m6A-associated pseudogene in the UC cohort from Mariathasan.

Since the high-risk subtype was able to benefit more from ICI therapy than the low-risk subtype, we would expect to see the high-risk subtype in companion with better survival outcomes after the treatment of ICI than the low-risk subtype. To test this, we assessed the prognostic value of m6A-associated pseudogenes in two urothelial cancer (UC) cohorts with anti-PD-L1 therapy [22, 23]. Surprisingly, we could find that high-risk patients had better OS than low-risk patients (Figure 5F-G). We could also find that the performance of m6A-associated pseudogenes was consistent in the UC cohort from
Snyder [22] and the UC cohort from Mariathasan [23], at 12 months of follow-up (Figure 5H-I). Thus our study indicated that the predictive value of the expression pattern of m6A-associated pseudogenes was comparable with TIDE and TIS for OS.

**Identification and experimental validation of potential therapeutic agents with higher drug sensitivity for HNSCC patients in the high-risk subtype**

In addition to HNSCC patients with m6A-associated pseudogene expression patterns who may benefit from immunotherapy, we also expect these patients to benefit from therapeutic agents. To solve this issue, we took advantage of two powerful datasets, which could quickly screen thousands of drugs from hundreds of human cancer models on an unprecedented scale. Specifically, the Cancer Therapeutics Response Portal (CTRP) and Profiling Relative Inhibition Simultaneously in Mixtures (PRISM) datasets, which contain the gene expression profiling and drug sensitivity profiling of hundreds of cancer cell lines (CCLs), can be used to build a prediction model of drug response. 160 of these compounds are shared between the two datasets. The CTRP has unique 322 compounds and the PRISM has unique 1288 compounds. Results showed that there were 1770 compounds both in the CTRP and PRISM datasets after removing duplication (Figure S12A). Compounds with NAs in more than 20% of the samples were excluded. Ultimately, 654 CCLs with 354 compounds in the CTRP dataset, as well as 439 CCLs with 1291 compounds in the PRISM dataset were utilized for subsequent analysis.

To identify potential therapeutic agents in HNSCC patients, two different methods were used to confirm candidate drugs with higher drug sensitivity in high-risk score patients (Figure S12B). The analyses were conducted using CTRP and PRISM datasets, respectively. Firstly, differential drug response approaches between high-risk score (top decile) and low-risk score (bottom decile) groups were adopted to identify compounds with lower calculated AUC values in the high-risk score group with log2FC > 0.10 (Table S9; Table S10). Secondly, Spearman correlation calculation between AUC value and risk score was performed to screen compounds through a negative correlation coefficient (Spearman’s R < -0.30 for both CTRP and PRISM). The above analyses yielded seven CTRP-derived compounds (including paclitaxel, doxorubicin, gemcitabine, vincristine, SB-743921, clofarabine, and rigosertib) (Figure 6A-B) and nine PRISM-derived compounds (including paclitaxel, docetaxel, NVP-AUY922, dasatinib, epothilone-b, talazoparib, topotecan, rubitecan, and vinblastine) (Figure 6C-D). These screened compounds had lower calculated AUC values in the high-risk score subtype, as well as a negative correlation with a risk score for CTRP (Figure 6B) and PRISM datasets (Figure 6D), which might be potential therapeutic agents in HNSCC patients with a high-risk score.

Despite the 16 candidate therapeutic agents identified showing a higher drug sensitivity in high-risk score HNSCC patients, the above screening method alone was not able to support the conclusion that these agents had a therapeutic effect on HNSCC. Thus, multiple perspective analyses were subsequently performed to investigate the therapeutic potential of these compounds in HNSCC. Firstly, fold-change values in the expression levels (including mRNA-level and protein-level) of candidates’ drug targets between tumor and normal tissue were calculated. In the calculation results, a higher fold-change value manifested a greater potential and better efficacy of the candidate compound for HNSCC treatment (Table S11). Secondly, the CMap analysis approach was adopted to identify compounds in which gene expression patterns were contrary to the HNSCC-specific expression patterns (in other words, gene expression increased in tumor tissues but decreased by treatment of certain agents). Four compounds, including doxorubicin, clofarabine, NVP-AUY922, and topotecan, had CMap scores < -95 (Table S12), on behalf, that these compounds might have a potential therapeutic effect on HNSCC (Figure 6E-F). Thirdly, comprehensive analyses including drug data query in DrugBank and literature search in PubMed were conducted to look for the experimental validation and clinical evidence of candidate agents in treating HNSCC (Figure 6E-F). The above three results were presented on the middle, right, and left of the panel respectively (Figure 6E-F). Finally, doxorubicin, and topotecan, which had robust multi-level evidence including in vitro and in silico, were deemed to hold the most promising therapeutic potential in HNSCC patients with a high-risk score.

To experimentally test whether potential therapeutic agents with higher drug sensitivity for HNSCC patients in high-risk subtype affect the tumor cell viability, we performed IC50 (in other words, the concentration of drug which causes 50% cell viability) assay of doxorubicin and topotecan in head and neck cell lines (ARO and Tca8113) and other corresponding sensitive cell lines (HeLa, GSCs-U251, and MDA-MB-231). The effect of doxorubicin and topotecan on cell viability after 24 h treatment was evaluated using a colorimetric MTT assay in both cell lines. The two therapeutic agents have their distinct effects on cell viability at 10 different concentrations. Our findings revealed that the cell survival rate was observed to
generally decrease with an increase in drug concentration, suggesting a dose-dependent behavior. The IC$_{50}$ value of ARO cells for doxorubicin and topotecan were 1.6 nM, and 3.5 nM, respectively (Figure 6G-H). And the IC$_{50}$ value of Tca8113 cells for doxorubicin and topotecan were 2.2 nM, and 4.6 nM, respectively (Figure 6I-J). The IC$_{50}$ value of doxorubicin and topotecan for ARO and Tca8113 cells were less than those for HeLa, GSCs-U251, and MDA-MB-231 cells, as well as in previous studies, further verifying that HNSCC patients in the high-risk subtype could benefit more from doxorubicin and topotecan.

Figure 6. Identification and experimental validation of potential therapeutic agents with higher drug sensitivity for HNSCC patients in the high-risk subtype. (A) Horizontal column diagram showing the correlation coefficient and P-value of Spearman’s correlation analysis of seven CTRP-derived compounds. (B) Boxplots displaying estimated AUC.
value of differential drug response analysis of seven CTRP-derived compounds between high-risk score patients with low-risk score patients. Note that lower values on the y-axis of boxplots imply greater drug sensitivity. The value between the two subtypes was compared through the Wilcoxon test. ns denotes no significance, * denotes $P < 0.05$, ** denotes $P < 0.01$, *** denotes $P < 0.001$ and **** denotes $P < 0.0001$. (C) Horizontal column diagram showing the correlation coefficient and $P$-value of Spearman’s correlation analysis of nine PRISM-derived compounds. (D) Boxplots displaying estimated AUC value of differential drug response analysis of nine PRISM-derived compounds between high-risk score patients with low-risk score patients. Note that lower values on the y-axis of boxplots imply greater drug sensitivity. The value between the two subtypes was compared through the Wilcoxon test. ns denotes no significance, * denotes $P < 0.05$, ** denotes $P < 0.01$, *** denotes $P < 0.001$ and **** denotes $P < 0.0001$. (E) Identification of most promising therapeutic CTRP-derived agents for high-risk score patients according to the evidence from multiple sources. Combined heatmap showing seven CTRP-derived agents. mRNA or protein expression was compared by fold change (FC) differences of drug targets between tumor and normal tissue (FC > 0 represents up-regulated in tumor tissue). (F) Identification of most promising therapeutic PRISM-derived agents for high-risk score patients according to the evidence from multiple sources. Combined heatmap showing nine PRISM-derived agents. mRNA or protein expression was compared by fold change (FC) differences of drug targets between tumor and normal tissue (FC > 0 represents up-regulated in tumor tissue). (G-H) MTT assay to determine the IC50 value of the different drug (doxorubicin and topotecan) and analyze their effect on head and neck cancer cell lines (ARO and Tca8113) and other sensitive cell lines (HaLa, GSCs-US1, and MDA-MB-231) viability. There are three biological replicates of cell viability per drug concentration. Line graph showing the different drug concentrations used and the corresponding cell viability between ARO with other cells treated with doxorubicin (G), between ARO with other cells treated with topotecan (H), between Tca8113 with other cells treated with doxorubicin (I) and between Tca8113 with other cells treated with topotecan (J). The IC50 values of ARO cells for doxorubicin and topotecan were 4.6 nM, and 4.6 nM, respectively. The IC50 values of Tca8113 cells for doxorubicin and topotecan were 2.2 nM, and 4.6 nM, respectively.

**Discussion**

In this study, 10 m6A-associated pseudogenes were confirmed as promising prognostic indicators for HNSCC by a comprehensive analytical framework and classified into oncogene pseudogenes (PDIA3P1, LDHAP4, LDHAP7, EEFA1AP6, EEFA1AP11) and tumor-suppressor pseudogenes (SDHAP1, SDHAP3, DDX12P, CLUHP3, RRN3P3) owing to their different effects in prognosis in TCGA dataset. Then a risk score model based on the 10 m6A-associated pseudogenes was constructed and found very good in predicting clinical outcomes in HNSCC, which was further validated in the GEO dataset, as well as our clinical tissues. More importantly, we found that the expression pattern of these 10 pseudogenes was dramatically associated with the immune response in terms of some aspects. Then, m6A-associated pseudogene-miRNA-targeted immune-involved regulatory networks were further performed to elucidate the underlying mechanisms that pseudogenes with m6A RNA modification could regulate antitumor immune response via miRNAs. Up to now, this is the first research to systemically clarify the prognostic value of m6A-associated pseudogenes and their regulatory roles in the host antitumor immune response of HNSCC. The novel discovery in this study also unveiled that the therapy of immune checkpoint inhibitors and doxorubicin and topotecan could enable HNSCC patients with the specific expression pattern of m6A-associated pseudogene to obtain good therapeutic effects.

Pseudogenes, as non-coding RNAs, are prevalently transcribed in the genome [24] and are nonfunctional and deleterious. A large proportion of unprocessed and processed pseudogenes can be removed in time in the RNA surveillance system. For example, most unprocessed pseudogenes can be degraded by Nonsense Mediated Decay (NMD) [25], and major processed pseudogenes can be cleaned by m6A on the RNAs of processed pseudogenes in our previously published articles [20]. However, there are still some pseudogenes that have been nonremovable and still highly expressed, which can play increasingly important regulatory roles in diverse human diseases [26] as well as contain miRNA-binding elements and therefore increase their parental and other targeted genes by acting as ceRNA [27]. Our results revealed that 6218 out of 13931 pseudogenes were expressed in TCGA HNSCC, which were our following research targets. So what role do these expressed pseudogenes play in cancer? A series of previous studies have depicted the crucial roles of pseudogenes in tumorigenesis and tumor development. For instance, PTPN1 could suppress the progression of clear-cell renal cell carcinoma by acting as a ceRNA [28]. PKP3, AC027614.4, HILS1, RPS11-H153, and HSPB1P1 were found as prognostic predictors for lower-grade gliomas [29]. ANX2P2, EEF1A1P9, FER1L4, HILS1, and RAET1K were identified to be dramatically correlated with the survival of glioma [30]. RNA55P141 could strongly enhance the RIG-I-mediated antiviral immune response to herpes simplex virus 1 [31]. However, there has been no focus on the role of pseudogenes in HNSCC, and our research will focus on this question.

m6A RNA modification is installed by m6A methyltransferases METTL3 and uninstalled by m6A demethylases FTO and ALKBH5 [14, 15]. A variety of RNA binding proteins can modulate diverse post-transcriptional processes of host mRNAs and non-coding RNAs by reading different m6A on these RNAs [14, 15], such as facilitating the cytosol degradation [13] and accelerating nuclear export of mRNAs [32]. Increasing critical roles of m6A have been reported in many kinds of physiological and pathological processes of various cancers [33], including bladder cancer [35], gastric cancer [33], acute myeloid leukemia [33], and our published breast cancer [18]. However, recent research mainly focuses on the function of m6A in mRNA, there are very few studies on the roles of m6A on pseudogenes. We firstly screened 842 pseudogenes by Pearson correlation analysis, and ultimately screened 10 pseudogenes by univariate and multivariate Cox proportional hazard regression. Exploring the role of these 10 pseudogenes, defined as m6A-associated prognostic pseudogenes, in HNSCC will be the main
line of our next research. So far, two recent articles pointed out that the m6A-modified pseudogene HSPA7 could be a new immunotherapy target for GBM patients [34], and aberrant m6A modification of the pseudogene WTAPP1 results in increased translation of its protein-coding counterpart to promote pancreatic cancer progression [35]. However, there has been no focus on the role of m6A on pseudogenes in HNSCC. Our findings revealed the prognostic function of screened 10 m6A-associated pseudogenes in HNSCC patients. To investigate whether 10 m6A-associated pseudogenes are an independent prognostic factor, we analyze the prognostic effects of 10 pseudogenes both in the TCGA and GEO datasets. Convergent evidence supports the 10 m6A-associated pseudogenes as an independent prognostic factor. To further demonstrate the prospect of m6A-associated pseudogene clinical application, we verified the correlation of pseudogene expression with clinical overall survival in a relatively small cohort including 32 HNSCC tissues from our hospital. Unfortunately, pseudogenes are non-coding proteins, so pseudogenes cannot express proteins, so we cannot verify the expression of pseudogenes at the protein level, which limits its potential clinical application value.

Our current study indicated that 10 m6A-associated pseudogenes were identified as promising prognostic indicators for HNSCC. Curiously, the oncogene pseudogenes happened to be processed pseudogenes, however, the tumor-suppressor pseudogenes were unprocessed pseudogenes, which was worthy of further study. Carcinogenic roles of PDLA3P1 in HNSCC were in close agreement with that a higher expression of PDLA3P1 was closely associated with a poorer recurrence-free survival of human hepatocellular carcinoma [36]. The anticancer effect of SDHAP1 was accordant with that SDHAP1 upregulated EIF4G2 level by sponging miR-4465 and therefore promoted the PTX-induced apoptosis in ovarian cancer [37]. SDHAP3 plays an important role in carcino genesis in our study, which also displays strong involvement in neurodevelopmental disorders, and cancer susceptibility [38]. The role of the remaining 7 m6A-associated pseudogenes (LDHAP4, LDHAP7, EEF1A1P6, EEF1A1P11, DDX12P, CLUHP3, and RRN3P3) has not been reported in the previous literature and has been coming up in our research, which was indispensable and identified as a good prediction of outcome in HNSCC. In summary, our study provides prospective prognostic predictors for HNSCC, which can better fulfill the principle of precise medicine.

Our findings firstly revealed that the expression pattern of the 10 m6A-associated pseudogenes was dramatically associated with tumor-infiltrating B cells, CD8+ T cells, helper T cells, and regulatory T cells, as well as the expressions of T cell exhausted markers including PD-1, PD-L1, PD-L2, LAG3, TIGIT, and CTLA4. What’s more, antigen presentation genes, interferon activity genes, cytolytic genes, integrin genes, and kinase genes were also significantly associated with pseudogenes levels. To explain the mechanism of how m6A-associated pseudogenes regulated immune response, pseudogene-miRNA-targeted immune-involved gene regulatory networks were further constructed and validated experimentally to explain the underlying mechanisms and demonstrated that m6A-associated pseudogene can regulate antitumor immune-involved target genes via plenty of miRNAs. We found that the oncogene PDIA3P1, and tumor-suppressor RRN3P3, promote the RNA and protein expression of their targeted immune-involved genes AKT1 and EZH2 via miR-34a-5p and miR-26b-5p, respectively. Existing research indicates that AKT inhibition reduces PD-L1 expression in tumor cells, enhances activation and tumor infiltration of CD8+ T cells, and reduces tumor growth, accompanied by prolonged mouse survival [39]. The previous study has shown that suppressing EZH2 activity resulted in increased numbers of myeloid-derived suppressor cells (MDSC) and fewer CD4+ and IFNγ+CD8+ T cells, which are involved in antitumor immunity [40]. However, another study suggests that cell cycle-related kinase (CCKR) activated the EZH2/NF-κB/IL-6 cascade, which lead to the accumulation of polymorphonuclear MDSCs, downregulated PD-L1 expression, and decreased intratumoral CD8+ T cells [41]. Inconsistent conclusions from two research teams further reveal the complexity of EAH2’s role in tumors. What’s more, we quantified the m6A levels and identified m6A peaks and m6A sites on m6A-associated pseudogenes, and verified the m6A level of the pseudogenes at specific modification sites in HNSCC cells by m6A-qPCR.

Our current study also firstly suggested the clinical application of m6A-associated pseudogenes in HNSCC, which can effectively predict treatment outcomes of immune therapy and drug therapy. By integrating with public datasets about immune therapy and drug sensitivity of CCLs, HNSCC patients in the high-risk subtype could benefit more from immune checkpoint inhibitors and promising potential therapeutic agents including doxorubicin and topotecan. Our study indicates that the high-risk subtype with higher PD-L1 expression, fewer infiltrating CD8+ T cells, and Low-MSI might have a better response from ICI treatment, which was
possibly attributed to the effects of PD-L1 on apoptosis of CD8+ T cells [42]. Our results of drug targets and compounds complemented each other, indicating a comprehensive view of a potential personalized treatment strategy, were further confirmed in vitro experiments. Overall, our study indicated that m6A-associated prognostic pseudogenes have not only provided new insights into personalized prognostication approaches but also thrown light on integrating tailored risk stratification with precision therapy.

Conclusions

Our discovery revealed that the 10 m6A-associated prognostic pseudogenes significantly contribute to predicting immunotherapy benefits and therapeutic agents, which might bring some potential implications for both immunotherapy strategy and medical treatment in HNSCC. It provided new insights into personalized prognostication approaches and precision therapy.

Materials and methods

Data sources

The list of pseudogenes were compiled from GENCODE (https://www.encodeproject.org/) [43], Vega (http://vega.archive.ensembl.org/index.html), and Pseudogene.org databases (http://pseudogene.org/) [44]. The gene expression profiles and corresponding detailed clinical information of HNSCC were downloaded from the TCGA data portal (http://firebrowse.org/) and GEO dataset (https://www.ncbi.nlm.nih.gov/geo/) by access number (GSE65858). Altogether, 544 samples (including 500 tumor tissues and 44 normal tissues) from TCGA and 270 tumor samples from GEO with pseudogene expression profiling and corresponding clinical data were included. The immune cell fraction data were obtained through CIBERSORT (https://cibersort.stanford.edu/) [45, 46]. The antigen-presenting genes and immunomodulatory genes were obtained from TCIA (https://tcia.at/home) [47]. The miRNAs binding to pseudogenes were extracted from the dreamBase database (http://rna.sysu.edu.cn/dreamBase/index.php) [48]. The miRNA-targeted genes were identified using the miRTarBase (https://mirtarbase.cuhk.edu.cn/~miRTarBase/miR TarBase_2022/php/index.php) [49]. The m6A-LAIC-seq data of non-tumor cell lines (GM12878 and H1 hESC) were obtained from our previous publication [17, 20]. The m6A-seq data of non-tumor cell lines (GM12878 [21] and H1 hESC [50]) were also acquired from previous publications. Expression profiles, clinical files, and anti-PD-L1 therapy information about urothelial cancers were accessed from previous publications [22, 23]. Expression profiles of human cancer cell lines (CCLs) were downloaded from the Broad Institute Cancer Cell Line Encyclopedia (CCLE) project (https://portals.broadinstitute.org/ccle/) [51]. Drug sensitivity data of CCLs were achieved from the Cancer Therapeutics Response Portal (CTRP v.2.1, https://portals.broadinstitute.org/ctrp.v2.1/) [52-54] and PRISM Repurposing dataset (PRISM 19Q4, https://depmap.org/portal/prism/) [55-58].

Processing of sequencing data

The RNA-seq, m6A-LAIC-seq, and m6A-seq raw reads were subject to quality control with Fastqc, removed the adapter with Cutadapt, and dismissed low-quality bases with Trimmomatic according to the standard protocol of sequencing data with default parameters. Then preprocessed reads are conventionally aligned to the hg19 human genome using Hisat2 with default parameters [59]. The proper paired and uniquely mapped reads with perfect match except for mismatches at SNPs were used for the downstream analyses. RNA expression was evaluated by Transcripts Per Kilobase of exon model per Million mapped reads (TPM). TPMs of genes were calculated using StringTie2 [60]. Differential gene expression analyses were performed using the DESeq2 R package. Genes with an adjusted P-value < 0.05 (detected by DESeq2 soft) were differentially expressed. Since most of the pseudogenes were not expressed, we first obtained available pseudogenes in all datasets and excluded those with a TPM value less than 1.

Collation of m6A RNA methylation regulators and pseudogenes

We first compiled a whole list of m6A RNA methylation regulators from previously published literature, and then restricted the list to the genes with available RNA expression data in the TCGA HNSCC dataset, which ultimately resulted in a total of 24 m6A regulators in the current study. These regulators were stratified into 3 categories based on their functions (List S1). Above all, we systematically compared the expression levels of m6A regulators between tumor tissues and normal tissues, as well as among different pathological characteristics. Then, the protein-protein interaction (PPI) network based on the STRING database of studied m6A regulators was constructed to explore whether there is an interaction between m6A regulators. Hereafter, to investigate whether there is co-expression and correlation between m6A regulators, a Pearson correlation analysis among m6A regulators was executed. Afterward, we compiled a
list of pseudogenes from GENCODE, Vega, and Pseudogene.org databases, which were used for the downstream analyses (List S2).

Screening for m⁶A-associated prognostic pseudogenes by three methods

The m⁶A-associated prognostic pseudogenes were screened using the following three methods, (1) carry out the Pearson correlation analysis of expression between pseudogene in this study with m⁶A regulators, and filter m⁶A-associated pseudogenes under the conditions | correlation coefficient (referred to as R) | ≥0.3 and P < 0.05; (2) perform the univariate Cox proportional hazard regression to screen prognostic pseudogenes; (3) conduct the multivariate Cox proportional hazard regression based on the second step. Finally, overlapping candidate m⁶A-associated prognostic pseudogenes were identified.

Construction of the risk score model

Based on the LASSO Cox regression algorithm [61], an L1-penalized regression on the strength of the highest lambda value selected utilizing 1,000 cross-validations (‘1-se’ lambda) was implemented to further identify the regression coefficients of the candidate m⁶A-associated prognostic pseudogenes. Then we established a survival risk score model through the LASSO coefficients (β) as follows:

Risk score = \sum_{i=1}^{n} GeneExpr \times \beta_i.

The HNSCC patients were dichotomized into high-risk or low-risk groups based on the median risk score. The receiver operating characteristic (ROC) curve and area under the curve (AUC) was performed to assess the prediction accuracy of the risk score model. Each m⁶A-associated prognostic pseudogene was divided into low or high expression levels, with the cut-off values defined as the median expression value. And Kaplan-Meier plots and Log-rank tests were carried out to estimate and compare the survival rate between subtypes. Then univariate and multivariate Cox regression analyses were utilized to determine the prognostic value of the risk score and various clinical characteristics. Nomograms for evaluating the risk of HNSCC were developed for HNSCC patients based on risk factors. All the analyses mentioned above were conducted using TCGA data as the training set and GEO data as the validation set.

RNA extraction and real-time quantitative PCR (RT-qPCR) in HNSCC tissues

We applied and collected tumor tissues from 32 HNSCC patients with clinical features, including 16 oral cancer and 16 thyroid cancer tissues, which were from the Tumor Resource Bank of Sun Yat-sen University Cancer Center. Total RNA derived from HNSSC tissues was extracted using the NucleoZol RNA reagent (MACHEREY-NAGEL). And 1 µg of DNA-free RNA was then reverse-transcribed using HiScript III-RT SuperMix for qPCR (+gDNA wiper) (Vazyme, R232). Then qPCR was performed using the ChamQ Universal SYBR qPCR Master Mix (Vazyme, Q711) and carried out in an LC480 Real-Time PCR System (Roche). Ultimately, the fold-change value was calculated using the 2⁻∆∆CT method. Besides, the clinical features of HNSCC patients and the primers of ten pseudogenes had been appended in List S3 and List S4, respectively.

Consensus clustering analysis and functional enrichment analysis

To investigate the biological functions of m⁶A-associated prognostic pseudogenes in HNSCC patients, we clustered the patients into different subgroups by the R package “ConsensusClusterPlus” (50 iterations, resample rate of 80%, and Pearson correlation) based on the expression levels of the m⁶A-associated prognostic pseudogenes in TCGA dataset [62]. To better understand and interpret the association between m⁶A-associated prognostic pseudogenes and malignancy of HNSCC, GO pathway analysis and GSEA [63] were performed to functionally annotate genes that are differentially expressed in different subgroups by using the R package “clusterProfiler” [64].

Immune cell infiltration analysis and antitumor immune response analysis

CIBERSORT [45], a bioinformatic deconvolution algorithm to estimate immune cell composition based on related-gene expression profiles, was utilized to calculate tumor-infiltrating cell compositions in HNSCC. The immune cell fractions, expressions of T cell exhausted genes, antigen presentation genes, interferon activity genes, cytolytic genes, integrin genes, and kinase genes were compared in different subtypes and subgroups by Wilcoxon signed-rank test.

m⁶A-associated pseudogene-miRNA-targeted immune-involved gene regulatory networks

Potential miRNAs binding to pseudogenes were obtained from the dreamBase database [48]. Potential miRNA-targeted genes with at least two solid experimental methods (reporter assay and western blot) were extracted using the miRTarBase [49]. Pearson analysis was carried out to calculate the expression correlation between m⁶A-associated pseudogenes and miRNA-targeted genes. Then these
targeted genes conforming to $|R| \geq 0.3$ and $P < 0.05$ were identified and applied to build pseudogene-miRNA-targeted immune-involved gene regulatory networks.

**Cell culture**

ARO, Tca8113, and HeLa cells were cultured in Roswell Park Memorial Institute (RPMI) Medium 1640 (Corning), supplemented with 10% fetal bovine serum (FBS) (Gibco) at 37 °C with 5% CO₂. GSCs-U251 and MDA-MB-231 cells were cultured in Dulbecco’s Modification of Eagle’s Medium (DMEM) (Corning), supplemented with 10% fetal bovine serum (FBS) (Gibco) at 37 °C with 5% CO₂. All cells were tested to ensure that they are free from Mycoplasma infection.

**siRNA and miRNA inhibitor transfection**

Cells were seeded in TC-untreated plates and transfected with siNC and specific siRNA or inhibitor NC and specific miRNA inhibitor using RNAiMAX (Invitrogen) according to the manufacturer’s instructions. RNA samples and protein samples were harvested at 72 h after transfection for qRT-PCR or western blotting. The siRNA pools targeting PDIA3P1 and RRN3P3, as well as control siRNA (NC), were synthesized by RIBOBIO (Guangzhou, China). The siRNA duplexes used in the current study are listed in List S5.

**RNA extraction and RT-qPCR in HNSCC cells**

Total RNA from HNSCC cells was extracted using the NucleoZol RNA reagent (MACHEREY-NAGEL). And the specific steps of RT-qPCR are the same as in Part “RNA extraction and real-time quantitative PCR (RT-qPCR) in HNSCC tissues”. In addition, the primers of pseudogenes and their targeted immune-involved genes had been appended in List S4.

**Western blotting**

Proteins were extracted from cells after incubating with RIPA buffer (Cell Signaling Technology, Cat. 9806) on ice for 10 min, and then the insoluble fraction was removed by centrifugation. 20 μg of extracted protein was separated on 15% SDS-PAGE and transferred to the PVDF membrane. The membranes were blocked in 5% BSA in Tris-Buffered Saline with 0.01% Tween 20 (TBS-T) at room temperature for 1 h and incubated overnight with primary antibodies diluted in 1% BSA/TBS-T at 4 °C, followed by incubating with goat anti-rabbit HRP conjugated secondary antibody diluted in TBS-T for 1 h at room temperature and visualized using Clarity™ Western ECL Substrate (Bio-Rad). The following antibodies were used for immunoblotting: AKT1 (1:1000, Affinity Biosciences, AF5150-50), and EZH2 (1:1000, Affinity Biosciences, AF5150-50).

**RNA stability assay**

The ultimate concentration of 5 μg/mL Actinomycin D (Sigma, A9415) was added to control and PDIA3P1/RRN3P3 knockdown cells to assess RNA stability. The cells were collected after incubation for indicated time points, and RNA samples were extracted for reverse transcription and qPCR. And 18S was utilized as the reference gene and the fold-change value was calculated using the $2^{-\Delta\Delta CT}$ method.

**m6A analyses about m6A-associated pseudogene**

We recalculated the m6A levels of all compiled annotated pseudogenes based on the reprocessed m6A-LAIC-seq profiles of non-tumor cells (GM12878 and H1 cells) according to the method described in our previously published paper [17, 20] and identified the m6A peaks based on the reprocessed m6A-seq profiles of non-tumor cells (GM12878 [21] and H1 [50]). In addition, the single-nucleotide m6A sites were determined by combining the m6A sites predicted by sequence-based m6A site predictors SRAMP [65] and Whistle [66] within m6A peak regions.

**Absolute quantification of m6A modification on pseudogene mRNA**

We adopted the Epi-SELECT™ m6A fraction quantification kit (Epibiotek) to detect absolute m6A levels on pseudogenes in ARO and Tca8113 cells. The experimental schematic and protocol of SELECT (single base elongation- and ligation-based qPCR amplification method) were performed as previously described [67]. In brief, 9.8 μl total RNA was mixed with 1.6 μl Up Primer (1 μM), 1.6 μl Down Primer (μM), and 2 μl dNTP in a 2 μl 10× Reaction Buffer. The RNA and primers were annealed by incubating the mixture at a temperature gradient: 90 °C for 1 min, 80 °C for 1 min, 70 °C for 1 min, 60 °C for 1 min, 50 °C for 1 min, and then 40 °C for 6 min. Subsequently, 3 μl of a mixture containing 0.3 μl SELECT™ DNA polymerase, 0.47 μl SELECT™ ligase, and 2.23 μl ATP was added to the former mixture to a final volume of 20 μl. The final reaction mixture was incubated at 40 °C for 20 min and denatured at 80 °C for 20 min. The qPCR reaction was performed using ChamQ Universal SYBR qPCR Master Mix (Vazyme, Q711) with the Roche Lightcycler 480 Instrument II system. According to the above methods, the experiments of the control and METTL3 knockout group were carried out. SELECT primer sequences and shMETTL3 sequences are listed in List S6.
Comprehensive analysis of ICI therapy benefit

To explore the prognostic value of m6A-associated pseudogene in HNSCC patients after immunotherapy, we compared the immune dysfunction and exclusion (TIDE) score, MSI, T cell dysfunction, TAM_M2, T cell exclusion score, MASC, and CAF, which were calculated online (http://tide.dfci.harvard.edu/) among different subtypes [68]. Besides, the tumor inflammation signature (TIS) score [69] was calculated as an average value of log2-scale normalized expression of the 18 signature genes (List S7) and compared in different subtypes. Moreover, four methods (ABSOLUTE, IHC, CPE, and ESTIMATE) [70-72] were performed to estimate the purity of the tumor, which can better understand ICI therapy’s benefit. To validate the prognostic value of m6A-associated pseudogene in patients after immunotherapy, we performed survival analyses in two urothelial cancer (UC) cohorts treated with PD-L1 blockade [22, 23]. Moreover, we performed time-dependent ROC curve analyses to obtain AUC which is used to evaluate the prognostic value of m6A-associated pseudogene with the R package of timeROC.

Predictive analysis of potential therapeutic agents for patients with high-risk scores HNSCC

Drug sensitivity profiles of cancer cell lines (CCLs) were downloaded from the Cancer Therapeutics Response Portal (CTRP v.2.0, https://portals.broadinstitute.org/ctrp/) and Profiling Relative Inhibition Simultaneously in Mixtures dataset (PRISM 19Q4, https://depmap.org/portal/prism/). At first, compounds with more than 20% of missing data were excluded. After imputation, the K-nearest neighbor (k-NN) imputation was used to impute the missing AUC values. Based on CTRP and PRISM-derived drug response profiles respectively, the two different approaches were determined to identify candidate agents with higher drug sensitivity in high-risk score patients (Figure S12B). Then the differences in the mRNA and protein expression levels of candidates’ drug targets between tumor and normal tissue were calculated, and a higher differential (fold change) value indicated a greater potential and better efficacy of the candidate agent. The CMap analysis was utilized to select agents in which gene expression patterns were contrary to the HNSCC-specific expression patterns [73]. The drug data query was performed in DrugBank (https://go.drugbank.com/) and the literature search was conducted in PubMed (https://www.ncbi.nlm.nih.gov/pubmed/), which was used to seek the experimental validation and clinical evidence of candidate agents in treating HNSCC. The code for the predictive analysis of this section was available at https://github.com/tlqsysu/Predictive-analysis-of-potential-therapeutic-agents-for-patients-with-high-risk-scores-HNSCC.

MTT assay

Cell viability was estimated by using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; Sigma, St Louis, USA) colorimetric assay. In short, cells were seeded in 96-well plates and incubated for 24 h, and cells were treated with the drug at different concentrations. After the treatments, cells were washed with phosphate-buffered saline (PBS) and then incubated in MTT solution (Sigma) for 4 h. After dimethyl sulfoxide (DMSO) was added to each well, the absorbance value was measured at 490 nm to decide the cell viability with a microplate reader (BioTek, Winooski, USA). The cell viability was plotted in a graph and the IC50 was calculated accordingly to determine the optimum dosage of the drugs for further studies.

Statistical Analysis

One-way ANOVA and t-test were conducted to compare the expression levels of m6A-associated prognostic pseudogenes in different subtypes and subgroups differentiated by different clinical characteristics. A Chi-square test was utilized to estimate the differences in clinicopathological characteristics in different subtypes and subgroups identified by consensus clustering of pseudogenes. All statistical analyses were carried out using R4.0 software (http://www.r-project.org/) and Bioconductor (http://bioconductor.org/). A two-sided P value < 0.05 was considered statistically significant in all analyses.

Abbreviations

TCGA: The Cancer Genome Atlas; GEO: Gene Expression Omnibus; LASSO: the least absolute shrinkage and selection operator; ROC: the receiver operating characteristic curve; AUC: area under the curve; PDIA3P1: protein disulfide isomerase family A member 3 pseudogene 1; LDHAP4: lactate dehydrogenase A pseudogene 4; LDHAP7: lactate dehydrogenase A pseudogene 7; EEF1A1P6: eukaryotic translation elongation factor 1 alpha 1 pseudogene 6; EEF1A1P11: eukaryotic translation elongation factor 1 alpha 1 pseudogene 6; EEF1A1P11: eukaryotic translation elongation factor 1 alpha 1 pseudogene 11; SDHAP1: succinate dehydrogenase complex flavoprotein subunit A pseudogene 1; SDHAP3: succinate dehydrogenase complex flavoprotein subunit A pseudogene 3; DDX12P: DEAD/H-Box Helicase 12 pseudogene; CLUHP3: clustered mitochondria homolog pseudogene 3; RRN3P3: homolog, RNA polymerase I transcription factor
pseudogene 3; P1: patient subgroup 1; P2: patient subgroup 2; ceRNA: competitive endogenous RNA; PD-1: programmed cell death 1; PD-L1: programmed cell death 1 ligand 1; PD-L2: programmed cell death 1 ligand 2; LAG3: lymphocyte-activation gene 3; TIGIT: T cell immunoreceptor with Ig and ITIM domains; CTLA4: cytotoxic T-lymphocyte-associated protein 4; HLA: major histocompatibility complex, class I; TAPI1: transporter 1, ATP binding cassette subfamily B member; B2M: beta-2-microglobulin; CCL5: C-C motif chemokine ligand 5; CXCL9: C-X-C motif chemokine ligand 9; CD24: CD24 molecule; CD27: CD27 molecule; STAT1: signal transducer and activator of transcription 1; IRF3: interferon regulatory factor 3; GZMA: Granzyme A; PRF1: Perforin 1; CYTH: cytohesin; ITGA: integrin subunit alpha; ITGB: integrin subunit beta; AKT1: AKT serine/threonine kinase 1; FOXM1: forkhead box M1; E2F2: E2F transcription factor 2; MECP2: methyl-CpG binding protein 2; MageA: Mage family member A; HOXA: homeobox A; EZH2: Enhancer Of Zeste 2 Polycomb Repressive Complex 2 Subunit; GO: Gene Ontology; GSEA: Gene Set Enrichment Analysis; TIDE: tumor immune dysfunction and exclusion; MSI: microsatellite instability; TAM: tumor-associated macrophages; MDSC: myeloid-derived suppressor cell; CAF: cancer-associated fibroblasts; TIS: tumor inflammation signature; IHC: immunohistochemistry; CPE: consensus measurement of purity estimations; ESTIMATE: Estimation of Stromal and Immune cells in Malignant Tumours using Expression data; CCLs: cancer cell lines; CTRP: Cancer Therapeutics Response Portal; PRISM: Profiling Relative Inhibition Simultaneously in cell lines; CTRP: Cancer Therapeutics Response Portal; PRISM: Profiling Relative Inhibition Simultaneously in Mixtures.

Supplementary Material

Supplementary figures, list and table legends. https://www.thno.org/v12p7267s1.pdf
Supplementary lists. https://www.thno.org/v12p7267s2.xlsx
Supplementary tables. https://www.thno.org/v12p7267s3.xlsx

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Author contributions

LT and MC conceived the study. LT and YQ carried out its design, analyzed, and interpreted the data. LT, RX, TX, XD, and LP performed experiments. LT wrote and edited the paper with YQ, RX, TX, XD, LP, RY, YL, XZ, MZ, and ML contributing to providing feedback. LT and MC revised the paper. All authors read and approved the final manuscript.

Availability of data and materials

All data used in this study are publicly available as described in the Materials and methods section. The web links or unique identifiers for public datasets are described in the paper. Other requests for data are available upon reasonable request from the lead contact. If you have any questions about the use of the database, please contact us or the author of the database directly.

Competing Interests

The authors have declared that no competing interest exists.

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