Autophagy regulation patterns characterization identifies immune phenotypes and immunotherapy response in head and neck squamous cell carcinoma (HNSCC)

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Abstract

Background

Autophagy degraded and recycled cytoplasmic components to maintain cellular homeostasis under stress conditions, which was recognized as double-edged sword in oncogenesis and novel target in cancer treatment. However, comprehensive analysis of the relationship between autophagy regulation and immunity has not been reported yet.

Methods

Unsupervised consensus clustering algorithm was used to identify autophagy regulation patterns. LASSO cox regression algorithm was used to build a scoring system (ATGscore) to represent the individual autophagy regulation pattern. Then integrated analysis of autophagy regulation patterns and ATGscore was performed.

Results

We have successful depicted five autophagy regulation patterns and established a scoring system (ATGscore) to represent it, which was shown to be significantly correlated with TIME infiltration, immune phenotypes, molecular subtypes, and genetic variation, etc. in 1165 head and neck squamous cell carcinoma (HNSCC) patients. Moreover, ATGscore was an independent prognostic factor and potent predictor for clinical response to immune-checkpoint inhibitors (ICIs) targeting immunotherapy.

Conclusion

Understanding the molecular characteristics of autophagy regulation patterns in HNSCC could help us to depict the underlying mechanism of tumour immunity and lay a solid foundation on combination of autophagy targeting therapies and immunotherapies for clinical application in HNSCC.

Background

Head and neck squamous cell carcinoma is one of the most common cancers with nearly 500,000 new cases and 350,000 deaths, which ranks 11th cancer-related deaths worldwide each year [1]. HNSCC is heterogeneous disease which not only originates from the different sites of the upper aero-digestive tract including oral cavity, oropharynx, larynx or hypopharynx, but also caused by distinct aetiologies. Smoking and alcohol abuse were recognized as the classical risk factors for HNSCC. Currently, infection with high-risk human papillomaviruses (HPVs) was considered as another vital cause for these tumours, especially in the oropharynx and occurring particularly in the Western world [2]. The patients with HPV-negative (HPV-) and HPV-positive (HPV+) diseases displayed different socioeconomic profiles, clinical presentation, molecular profiles, as well as the prognosis of the tumours [3–6]. Although combination therapy improved the prognosis of HNSCC patients, nevertheless, the five-year survival rate of patients
with HNSCC is still fewer than 50% [7, 8]. Thus, lack of rapidly improvement in patient survival and personalized treatment approaches prompted us to further study the molecular landscape of HNSCC.

Autophagy is a constitutively conserved physiologic catabolic process to harvest energy and nutrients from cellular components by degrading and recycling damaged organelles including macromolecules and organelles [9–11]. Under normal conditions, autophagy actively acts at basal rates to maintain cellular homeostasis, but could respond to survival demands to induce nonselective bulk degradation or selectively target cytoplasmic constituents via cargo-specific autophagy receptors [12, 13]. Thus, autophagy was involved in numerous biological and pathological processes, including neurodegenerative diseases, aging and cancer [14]. However, autophagy plays a dual role in oncogenesis through a context-dependent manner [15]. Under different tumour type, pathological stage, genetic background, and even therapeutic regimen, distinct autophagy patterns will be sustained to either promote or inhibit tumorigenesis [16]. Lack of autophagy could trigger the accumulation of genotoxic cellular waste and induce the genetic and chromosomal alteration, subsequently facilitating precancerous cells transformation and mature cancer cell formation [17]. However, excessive autophagy continuously recycle the remodelling components and refuel energy supply, which allow cancer cells to escape the damage from the immune system or targeting drugs and promoting tumour progression [18, 19]. All of these indicated the autophagy is double-edged sword in tumorigenesis, which has been already verified in preclinical cancer models [20]. In head and neck squamous cell cancers, some reported that increased levels of cytoplasmic p62, which indicated autophagy inhibition, was correlated with reduced overall and disease specific survival [21]. While other groups demonstrated that poor clinical outcome patients exhibited higher levels of LC3-II, which suggested the re-activation of autophagy [22]. However, Zhou et al showed that radiation-induced autophagy could increase the survival of CNE-2 cells, which was further counteracted by autophagy inhibition with chloroquine, resulting in enhanced cell death [23]. We noticed that in HNSCC there also existed an autophagy paradox where played a controversial role in tumorigenesis, but the underlying mechanism has not been reported.

Currently, we are experiencing a strikingly shift from chemotherapy and radiation anti-tumour broadly towards immune-checkpoints inhibitors (ICIs) targeting immunotherapy more precisely that regulate immune response against tumours. Recent studies have reported that autophagy significantly controls the immune response. Autophagy could not only activate receptors such as toll-like receptors (TLRs) and nucleotide oligomerization domain (NOD)-like receptors (NLRs) to induce natural killer T (NKT) activation, cytokine production, and phagocytosis in innate immunity, but also provide an abundance of antigen for MHC class II molecules including HLA molecular toward dendritic cells for cross-priming to CD8 + T cells in adaptive immunity [24, 25]. Moreover, autophagy could facilitate, promote or inhibit the proliferation and differentiation of a variety of immune cells or secretion a wide range of cytokines to modulate the tumour immune microenvironment (TIME) homeostasis. Conversely, certain cytokines and immune cells also exhibit a significant impact on autophagy function [26, 27]. Furthermore, immune-checkpoints including Indoleamine 2,3 dioxygenase(IDO), CTLA-4 and PD-1 were shown to regulate tumour immune tolerance through autophagy pathways. Accumulating evidences have showed that autophagy could interact with TIME immune cells and cytokines to strengthen or attenuate the immunotherapy response,
which has provided us targets for combination of immunotherapy [28]. However, definite correlation between autophagy and tumour immune microenvironment has not been study yet, a comprehensive and systematic analysis of them were urgently needed.

In present study, we integrated the transcriptional and genetic profiles of several cohorts to systematically analyse autophagy regulation patterns and established an autophagy phenotype related signature (ATGscore) to represent it in individual. As a result, we found distinct autophagy regulation patterns were highly correlated with TIME infiltration, immune phenotypes, molecular subtypes, as well as clinicopathological characteristics, and ATGscore was a robust independent prognostic and predictive factor for clinical outcome of ICIs immunotherapy.

**Materials And Methods**

**Data collection and processing**

Publicly available transcriptional datasets for HNSCC were systematically searched. Then eleven datasets including E-MTAB-1328, GSE39366, GSE41613, GSE42743, GSE65858, E-TABM-302, GSE6791, GSE30784, GSE40774, GSE84846 and TCGA-HNSCC were enrolled (Additional file 1: Table S1). After filtering out the samples without complete prognosis information, we finally got six datasets (E-TABM-302, GSE6791, GSE30784, GSE40774, GSE84846 and TCGA-HNSCC) with 1165 samples for further study. The microarray datasets were downloaded from Gene-Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/geo/) and ArrayExpress (www.ebi.ac.uk/arrayexpress/). The raw signal data in microarray were processed via RMA algorithm background correction, log2 transformation, quantile normalization and annotation by the package “Affy” in R [29]. Relative expression of each gene symbol was annotated as highest when several probes mapped to single gene symbol. Then “ComBat” algorithm of package “sva” in R, which reduce the likelihood of batch effects of non-biological technical biases from each dataset [30], was utilized to merge the five microarray datasets (E-TABM-302, GSE6791, GSE30784, GSE40774, and GSE84846) as a meta-HNSCC cohort. The level 3 fragments Per Kilobase per Million (FPKM) data of TCGA-HNSCC dataset were downloaded from the TCGA Genomic Data Commons (GDC) data portal (https://portal.gdc.cancer.gov/). Transcripts per kilobase million (TPM) values were transferred from the FPKM values to represent the relative expression of each gene symbol, which are more similar to gene expression from microarrays and more comparable between samples [31]. Detailed information of clinicopathological characteristics for TCGA-HNSCC dataset could be found in Additional file 2: Table S2. The somatic mutation data processed with MuTect2 algorithm for TCGA-HNSCC cohort was downloaded from the Genomic Data Commons (https://portal.gdc.cancer.gov/) using the package “TCGAbiolinks” in R [32]. The total number of mutations counted in the whole exon territory was set as 38 Mb according to a previous study [33]. Moreover, we also enrolled IMvigor210 (mUC) cohort, which included patients with metastatic urothelial cancer receiving PD-L1 inhibitor atezolizumab, to validate the results we found in HNSCC. The raw gene expression and clinical data were retrieved using the package “IMvigor” in R (http://research-pub.gene.com/IMvigor210CoreBiologies) [34]. The detailed clinical
information of IMvigor210 (mUC) cohort could be found in Additional file 3: Table S3. Data were analysed with the R (version 3.5.3) and Bioconductor packages.

Unsupervised consensus clustering for autophagy regulation patterns

Autophagy related genes (ATGs) were obtained from Human Autophagy Database (HADb, http://www.autophagy.lu/). Then ATGs were submitted to unsupervised consensus clustering algorithm (K-means) based on Euclidean distance and Ward's linkage for analysis to identify distinct autophagy regulation patterns (ATGclusters) [35]. The number of clusters was determined with a consensus clustering method in TCGA-HNSCC cohort and further validated in meta-HNSCC cohort. The package “ConsensusClusterPlus” in R was applied to performed this procedure and repeated 1000 times to guarantee the stability of classification.

Differentially expressed genes (DEGs) between autophagy regulation patterns

Based on the distinct expression of ATGs, patients with HNSCC were successfully classified into five autophagy regulation patterns through consensus clustering algorithm. DEGs were screened out by comparing the patterns with different function annotation by using the package “edgeR” in R [36], which implements an empirical Bayesian approach to evaluate the change of gene expressions in different groups. The significance criteria for determining DEGs were set as a false discovery rate (FDR) < 0.05 and |log2FC| > 1.0.

Estimation of infiltrating immune cells

Single cell gene set enrichment analysis (ssGSEA), which evaluated the variation in pathway and biological process activity in the single sample of gene expression dataset, was utilized to estimate relative amount of infiltration immune cells in HNSCC TIME by using package “GSVA” in R [37]. The basic unit for ssGSEA was gene set, which is consisted of genes that share common biological function, chromosomal location, or genetic regulation [38]. TIME infiltration immune cell types such as innate immune cells (dendritic cell, eosinophil, mast cell, macrophage, natural killer cell, neutrophil, etc.) and adaptive immune cells (B cell, T cell, T helper cell, CD8+ T cell, regulatory T (Treg) cell and cytotoxic cell, etc.) were obtained and from the study of Bindea and Charoentong [39, 40]. The normalized enrichment score (NES) from the ssGSEA was regarded as relative amount of TIME infiltrating immune cell in HNSCC.

Functional annotation and pathway enrichment analyses

The difference on biological process between distinct autophagy regulation patterns was analysed through ssGSEA algorithm. The gene sets of “h.all.v7.1.symbols” and “c2.cp.kegg.v7.1.symbols” were downloaded from MSigDB database of Broad Institute (https://www.gsea-msigdb.org/gsea/msigdb/index.jsp) to evaluate the regular biological pathway enrichment. In spite of HALLMARK and KEGG signalling pathway, a series of gene sets were curated to represent biological processes related with immune activation, stromal activation, DNA damage repair, and immune
checkpoints, which was obtained from Mariathasan et al, including (a) Angiogenesis; (b) Antigen processing machinery; (c) Base excision repair; (d) CD8 T-effector; (e) Cell cycle; (f) Cell cycle regulators (g) DNA damage repair 1 and 2; (h) DNA replication; (i) Epithelial-mesenchymal transition (EMT) 1, 2 and 3; (j) Fanconi anemia; (k) Homologous recombination; (l) Immune-checkpoint; (m) Mismatch repair; (n) Nucleotide excision repair; (o) Pan-fibroblast TGFβ response signature (Pan-F-TBRS); (16) WNT targets [34, 41, 42]

Construction of autophagy phenotype related signature

In order to define autophagy phenotype in individual patient, a set of scoring system was needed. The DEGs between autophagy regulation patterns were submitted to univariate cox analysis with cut-off p-value < 0.01 to generate candidate prognostic DEGs. For dimension reduction, we then performed LASSO cox regression algorithm on prognostic DEGs and construct an optimal autophagy phenotype related signature (ATGscore) through package “glmnet” in R [43]. The optimal values of the penalty parameter λ were determined through 10 cross-validations. The ATGscore of each sample was defined by the relative expression of candidate prognostic DEGs within the model and their cox coefficient. The ATGscore = Σ_(i=1)^n (coei × Expri), where Expri is the relative expression of the DEG in the signature for patient i and coei is the LASSO cox coefficient of DEG i.

Statistical analyses

The statistical significance of variables between two groups or more that two groups were by Wilcoxon tests or Kruskal-Wallis tests respectively. The differences between survival curves for each group were determined by Kaplan-Meier analysis with the log-rank test using the package “survminer” in R. The distance between different parameters was computed by Pearson and distance correlation analyses. Independent prognostic factors were identified through univariate and multivariate cox proportional hazard analysis and visualized with the package “forestplot” in R. ATGscore was integrated with other independent factors to establish the nomogram and calibration curves through packages “rms”, “nomogramEx” and “regplot” in R. According to Iasonos' suggestion, DCA was used to evaluate the clinical utility of our established nomogram [44]. The mutation landscape of patients with TCGA-HNSCC cohort was visualized with waterfall plot through packages “maftools” [45] and “complexheatmap” [46] in R. Contingency tables such as the ICIs targeting immunotherapy response were analysed by two-sided Fisher's exact tests. All statistical analyses were performed with R software 3.5.3. Statistical significance was set at p < 0.05.

Results

Characterization of autophagy regulation patterns in HNSCC

A workflow and design of this study were shown in Fig. 1a. TCGA-HNSCC cohort was used as the training set to identify the autophagy regulation patterns through package “ConsensusClusterPlus” in R. We found that ATGs could successful cluster five autophagy regulation patterns with high stability in TCGA-HNSCC
cohort, including 175 cases in pattern A, 56 cases in pattern B, 143 cases in pattern C, 191 cases in pattern D, and 33 cases in pattern E, which was also termed as ATGclusterA-E respectively (Fig. 1b). Unsupervised hierarchical clustering demonstrated that the ATGs were significantly differential expressed among ATGclusterA-E in TCGA-HNSCC cohort (Fig. 1c). Moreover, Kaplan-Meier survival curves showed ATGclusterA-E displayed a complete different survival beneficial that patients with ATGclusterB and E have a better prognosis than other clusters (Log-rank test, p = 0.0054; Fig. 1d).

**Tumour immune microenvironment (TIME) landscape in distinct autophagy regulation patterns**

As autophagy played a dual role in tumour immune microenvironment (TIME), here we aimed to make clear the association between TIME infiltration and autophagy in HNSCC. The landscape of TIME was calculated via ssGSEA algorithm and showed via cluster heat map (Additional file 4: Table S4). The relative amount of TIME immune cells was strikingly different in distinct ATGclusters as follows (Fig. 2a): ATGclusterA exhibited high infiltration with almost all immune cells; While ATGclusterB was remarkably rich in effector immune cells including activated B, activated CD4, CD8 T cells, activated dendritic cells (DCs) and cytotoxic cells, but less infiltrated with immunosuppressive cells such as regulatory T cells (Treg cells), macrophages and mast cells; ATGclusterC/D displayed low infiltration with all immune cells; ATGclusterE was characterized by high infiltration in activated CD8 and cytotoxic cells but low infiltration in regulatory Treg cells, macrophages and mast cells.

**Characteristics of biological process in distinct autophagy regulation patterns**

The biological process of distinct autophagy regulation patterns was explored. As shown in Additional file 5: Fig. S1a-b and Additional file 6-7: Table S5-6, five distinct ATGclusters were enriched in different HALLMARK and KEGG signalling pathways respectively: ATGclusterA was annotated with pathways associated with immune infiltration and stromal activation such as interferon a/g response, inflammatory response, epithelial mesenchymal transition (EMT) and TGFb signalling pathway, etc.; ATGclusterB was highly featured with immune activation and DNA damage repair (DDR) including antigen processing and presentation, T/B cell receptor signalling pathway, cytokine-cytokine receptor interaction and natural killer cell mediated cytotoxicity (Additional file 5: Fig. S1C); ATGclusterC was enriched in carcinogenic pathways including MTORC1 and TP53 signalling pathways; ATGclusterD/E were both prominently related to DDR related pathways activation.

**Distinct autophagy regulation patterns exhibit different immune phenotypes**

We found that ATGclusterA was highly infiltrated but patients with this pattern showed a worse prognosis. Recent study have determined three immune phenotypes of tumours: desert, excluded, and inflamed. Immune inflamed phenotype was characterized as high infiltration with immune cells, while immune desert phenotype displayed the opposite situation. Moreover, immune excluded phenotype were featured with infiltration of abundant immune cells, but most immune cells were located in the stroma surrounding the core tumour niche rather than penetrate their parenchyma, which was considered cytotoxic T-cell suppressive [47]. Then we enrolled a specific gene sets from Mariathasan et al to
investigate the enrichment of key signalling pathways associated with immune phenotypes. From the TIME landscape and function annotation analysis, ATGclusterA was recognized as immune-excluded phenotype as stromal related signalling pathways including angiogenesis, epithelial-mesenchymal transition (EMT), WNT target, and pan-fibroblast TGFβ response signalling pathways (pan-F-TBRS) were strikingly activated, which could hamper the beneficial effect of highly immune cells infiltration (Fig. 2b and Additional file 8: Table S7). Furthermore, ATGclusterB/E was remarkably induced in immune activation signalling pathway including antigen processing machinery, CD8 T effector, and immune checkpoint but deactivated in stromal related signalling pathways, which was the characteristics of immune-inflamed phenotype (Fig. 2b and Additional file 8: Table S7). The results from TIME landscape and pathway enrichment indicated that ATGclusterC/D was more likely to be immune-desert phenotype (Fig. 2b and Additional file 8: Table S7). Then major markers representing for immune phenotypes related signalling pathways were subsequent curated as follows: immune activation: CD8A, CXCL10, CXCL9, GZMA, GZMB, IFNG, PRF1, TBX2, and TNF; immune-checkpoints: CD274 (PD-L1), CD80, CD86, CTLA4, IDO1, LAG3, PDCD1 (PD-1), PDCD1LG2 (PD-L2), TIGIT, HAVCR2 (TIM-3) and TNFRSF9; MHC molecules: HLA-A/B/C, HLA-DMα/DMβ, HLA-DOA/DOB, HLA-DPA1, HLA-DPB1/DPB2, HLA-DQB1, HLA-DRA, HLA-DRB6, HLA-E/F/G; TGFβ/EMT signalling pathway: ACTA2, COL4A1, PDGFRA, SMAD9, TGFβ1/2/3, TGFβR1/2/3, TWIST1/2, VIM, and ZEB1/2. Surprisingly, the expression of above markers displayed a similar distribution as the biological process and pathways enrichment among distinct ATGclusters, which again validated that distinct autophagy regulation patterns exhibited differential immune phenotypes (Fig. 2c and Additional file 5: Fig. S1d-f).

**Tumour somatic mutation in distinct autophagy regulation patterns**

The relationship between somatic mutation and autophagy was also measured. We found that ATGclusterD exhibited highest tumour mutation burden (TMB), while ATGclusterB was associated with lowest TMB (Fig. 2d). Moreover, top 30 highly variant mutant genes were utilized to plot the somatic mutation landscape among distinct ATGclusters in patients with HNSCC. In Fig. 2e, ATGclusterD displayed highest mutation rate of top 30 mutant genes, especially for TP53, which was identified as key gene in tumorigenesis of HNSCC [48]. But ATGclusterB only showed small amount of TP53 mutation, which was consistent with the TMB calculation. All above results demonstrated that autophagy regulation patterns both correlated with TIME infiltration and tumour mutation landscape, which underlies the indispensable role of autophagy in HNSCC development.

**Validation of autophagy regulation patterns in meta-HNSCC cohort**

Unsupervised consensus clustering also identified five ATGclusters in meta-HNSCC cohort (Fig. 3a). The cluster heat map showed that the transcriptional profile of ATGs were differential distributed among five ATGclusters (Fig. 3b). Moreover, Kaplan-Meier survival curves demonstrated that the prognosis of five distinct ATGclusters were strikingly different, which patients with ATGclusterB/E lived longer and patients with ATGclusterA/C/D was associated with poorer survival (Log-rank test, p = 0.0022; Fig. 3c).
Furthermore, a similar TIME landscape among five distinct ATGclusters of TCGA-HNSCC was determined in meta-HNSCC cohort as follows (Fig. 3d and Additional file 9: Table S8): ATGclusterA/B/E was more infiltrated with immune cells, while ATGclusterC/D was less infiltrated. Compared with effector immune cells (CD4, CD8 T cells, and cytotoxic cells), ATGclusterA exhibited an abundance of immunosuppressive cells (Treg cells, macrophages and mast cells). The opposite situation was found in ATGclusterE that the amount of effector immune cells was robustly higher against immunosuppressive Treg cells, macrophages and mast cells. A similar phenomenon was seen in ATGclusterB which the ratio of cytotoxic T lymphocyte (CTL) including activated CD8 T cells and cytotoxic cells to Treg cells was highest among five ATGclusters. ATGclusterC/D displayed lowest infiltration of effector immune cells among five ATGclusters.

Then function annotation of ATGclusters in meta-HNSCC cohort showed a similar enrichment of KEGG and HALLMARK signalling pathways with TCGA-HNSCC cohort. ATGclusterA was both activated in immune infiltration and stromal activation related pathways, while ATGclusterB/E was only annotated with immune activation signalling pathways. ATGclusterC/D was both remarkably enhanced in DDR and EMT/ TGFβ related signalling pathway (Additional file 10: Fig. S2a-b and Additional file 11-12: Table S9-10). Moreover, ssGSEA of specific gene sets also showed a similar trend. ATGclusterA was both enriched in immune and stromal activation signalling pathways, which could be recognized as immune-excluded phenotype. ATGclusterB/E was more prominently enhanced in immune activation gene sets such as CD8 T effector, and immune checkpoint, which was more likely to immune-inflamed phenotype. And ATGclusterC/D was slightly annotated with DDR related signalling pathways but less enriched in immune infiltration signalling pathways, which was the characteristic of immune-desert phenotype (Fig. 3e and Additional file 13: Table S11). The expression distribution of immune phenotypes related markers displayed a similar trend to function annotation among five ATGclusters (Additional file 10: Fig. S2c-f). All of these again validated that autophagy played an indispensable role in the immune regulation within TIME and might distinguish immune phenotype in HNSCC.

Establishment of autophagy phenotype related signature (ATGscore)

Above findings revealed that there exists distinct autophagy regulation patterns in HNSCC and demonstrated an essential role of autophagy in shaping TIME landscapes, but all analyses were conducted in cohorts with patient population. Due to the heterogeneity of individuals, it was urgent to construct a set of scoring system to quantify the autophagy related pattern in individual patient with HNSCC. By comparing the transcriptomic profiles from main autophagy regulation patterns, we have obtained 5734 phenotype-related meta-DEGs (Additional file 14: Table S12). Then meta-DEGs were submitted to univariate cox regression analyses, 383 of 5734 meta-DEGs which were significantly correlated with prognosis were finally identified as autophagy phenotype candidate genes (Additional file 15: Table S13). Furthermore, LASSO cox regression analysis was utilized for dimension reduction on these genes to construct an autophagy phenotype related signature (ATGscore) which was representative for autophagy related pattern in individuals. At last, 11 genes were selected to establish ATGscore and the formula was as follows: ATGscore = ACTL10*(-0.0729) + C19orf57*(-0.0292) + CHAD*(-0.0801) + FCN2*
\(-0.3865\) + FGB*(0.1961) + GPR174*(-0.0437) + HSF5*(-0.0126) + SERPINA5*(0.1589) + SRPX*(0.0322) + ZNF541*(-0.033) + ZNF831*(-0.3632).

Kaplan-Meier survival curves showed that patients with low ATGscore lived longer than patients with high ATGscore when they were grouped at best cut-off in TCGA-HNSCC cohort (Log-rank test, p < 0.00001; Fig. 4a-b). Then we found that ATGscore was differentially distributed among five ATGclusters that ATGclusterA with poor prognosis showed the highest median score while ATGclusterB with good prognosis exhibited the lowest median score (Kruskal-Wallis test, p < 2.2e-16; Fig. 4c). Recently, a comprehensive molecular landscape has been constructed for HNSCC by The Cancer Genome Atlas Network, which classified HNSCC into atypical, basal, classical, and mesenchymal molecular subtypes [49]. Then we surprised to find that low ATGscore group was concentrated on the atypical subtype, which was correlated with low malignancy and good outcome, while other molecular subtypes with high malignancy and bad outcome, significantly accumulated in high ATGscore group (Fig. 4d-e). The correlation matrix showed that ATGscore and autophagy candidate genes was significantly correlated with immune cells which ATGscore was robustly negative associated with almost immune cells, indicating its role in TIME regulation (Fig. 4f and Additional file 16: Table S14). Then ssGSEA of KEGG and HALLMARK gene sets demonstrated that low ATGscore group was enriched in immune infiltration related pathways, but high ATGscore group was induced in EMT/TGF-b signalling pathways (Additional file 17: Fig. S3a-b and Additional file 6-7: Table S5-6). We also detected the relationship between ATGscore and intrinsic gene expression gene sets closely linked to the stromal activation program and immune activation procedure. Function annotation demonstrated that immune activation gene sets such as antigen processing machinery, CD8 T effector, and immune checkpoint were strikingly induced in low ATGscore group, while stromal relevant signalling pathways were enhanced in high ATGscore group (Fig. 4g). ATGscore was found to be negative correlated with immune activation signature and positive correlated with stromal activation signature through Pearson correlation analysis (Fig. 4h and Additional file 18: Table S15). Then the expression of immune phenotype marker genes showed similar trend with results of ssGSEA (Additional file 19: Fig. S4a-d).

Then the role of ATGscore was validated in EMTAB1328, GSE41613, and GSE42743 meta-cohorts, which contained eleven autophagy phenotype candidate genes. Cluster heat map demonstrated that TIME infiltration distribution among five ATGclusters was similar to meta-HNSCC cohort (Fig. 5a and Additional file 20: Table S16). Moreover, we could clearly see that the amount of effector immune cells including CD8 T cells and cytotoxic cells were robustly accumulated in low ATGscore group when compared with high ATGscore group in ATGclusterA/B/E, which was characterized as high immune infiltration (Fig. 5a). Kaplan-Meier survival curves demonstrated that patients with low ATGscore had a better prognosis than patients with high ATGscore (Log-rank test, p = 0.0196; Fig. 5b). Then function annotation of HALLMARK and KEGG gene sets showed that EMT/TGF-b signalling pathways was highly enriched in high ATGscore group, while low ATGscore group was featured with immune infiltration and activation related signalling pathways enrichment (Additional file 21: Fig. S5a-b and Additional file 22-23: Table S17-18). Furthermore, low ATGscore group exhibited upregulation of immune activation, immune checkpoints, and human leukocyte antigen (HLA) relevant markers and downregulation of TGF-b/EMT relevant markers, while high
ATGscore displayed the opposite distribution (Additional file 24: Fig. S6a-d). We also found that ATGscore was negative correlated with the expression of immune checkpoints (Fig. 5c). Moreover, ATGscore was significantly positive correlated with angiogenesis, EMT, Pan-F-TBRS, and WNT targets signatures, and negative correlated with antigen processing machinery, CD8 T-effector, and immune-checkpoint signatures (Fig. 5d-e and Additional file 25-26: Table S19-20). All above results strongly suggested that ATGscore is a good representative of autophagy regulation patterns and is competent at distinguishing immune phenotypes in HNSCC.

**ATGscore can be utilized as an independent prognostic factor in HNSCC**

We next want to make clear the association between ATGscore and clinical characteristics in HNSCC. The results showed that high ATGscore group were more likely to be patients with more advanced pathological TNM stage, HPV negative, and smoker, as well as treatment outcome of stable disease (SD)/ progression disease (PD), with tumour, and dead status, but did not show a correlation with alcohol history. The opposite patterns could be seen in low ATGscore group (Fig. 6a-f, Additional file 27: Fig. S7a-e, and Additional file 2: Table S2). The distribution changes between ATGscore and clinical features were visualized with alluvial diagram (Fig. 6g).

As gene mutation and TMB were remarkably different in five ATGclusters, we next investigated whether they were still different between low and high ATGscore groups in TCGA-HNSCC cohort. The waterfall plots revealed that TP53 and CDKN2A were differential mutated between high and low ATGscore groups (Fig. 6h). Kaplan-Meier survival curves demonstrated that the prognosis of patients with TP53 mutation were robustly worse than patients with TP53 wild type (WT) (Fig. 6i). We were amazed to find that patients with TP53 mutations were more likely to be higher ATGscore (Fig. 6j). Although mutation of CDKN2A was not correlated with patients’ survival, patients with high ATGscore were also more likely to be CDKN2A mutation (Fig. 6k-l). Unfortunately, we found no difference in TMB between the two groups with high and low ATGscore (Additional file 27: Fig. S7f). This might be caused by the mutation data in TCGA-HNSCC cohort as the high TMB patients were correlated with worse prognosis, which is against our common sense (Additional file 27: Fig. S7g).

As highly associated with the above clinical traits in HNSCC, we sought to determine whether ATGscore was responsible for prognosis prediction independent of these clinicopathological characteristics. Patients were divided into different subgroups according to their features. Stratification survival analyses demonstrated that ATGscore could efficiently predict the prognosis of HNSCC patients in almost all the subgroups from the aforementioned clinical features (Additional file 28-29: Fig. S8-9). Furthermore, univariate and multivariate cox regression analyses were enrolled to figure out independent prognostic role of ATGscore in HNSCC. ATGscore, together with other clinical features, including age, gender, histological grade, HPV status, smoking status, histologic grade, and alcohol history, TMB, pathological T stage, pathological N stage, pathological TNM stage, were enrolled as covariates to conduct the analysis. The results demonstrated that the ATGscore, pathological T stage, pathological N stage, and gender were independent factors that could be utilized to predict the prognosis of HNSCC patients (Additional file 30:...
Fig. S10a-b and Additional file 31: Table S21). Then we constructed a nomogram by integrating above independent prognostic factors to serve as a clinically relevant quantitative method for clinicians to predict mortality in patients with HNSCC (Additional file 30: Fig. S10c). By using the nomogram, each patient would get a total point by adding the points for each prognostic parameter. The patients obtained the higher total points correspond to a worse clinical outcome. The calibration plots and DCA demonstrated that our nomogram had displayed a similar performance to that of an ideal model and had high potential clinical utility (Additional file 30: Fig. S10d-g). All of these demonstrated that ATGscore could not only estimate the autophagy related pattern of individual patients, characterized immune phenotypes, as well as further acted as an independent prognostic factor in HNSCC.

**ATGscore was potent to predict clinical response to ICIs immunotherapy**

Recently, ICIs immunotherapy has emerged as a major breakthrough in the treatment of solid tumours. Above findings demonstrated that ATGscore could not only distinguish the prognosis of patients, but also determine TIME infiltration and immune phenotype, which indirectly confirmed its potential role in predicting clinical response to ICIs treatment. Then IMvigor210 (mUC) cohort, which was consist of the patients with metastatic urothelial cancer receiving PD-L1 inhibitor with atezolizumab, were introduced to study the role of ATGscore in evaluating immunotherapeutic benefits. Kaplan-Meier survival curves showed that patients with low ATGscore exhibited a significantly better clinical outcome compared with patients with high ATGscore (Log-rank test, p < 0.001, Fig. 7a).

Previous study reported that ICIs treatment responders were more likely to be patients with genomically unstable (GU) subtype in Lund classification system and TCGA II subtype TCGA classification system by using the IMvigor210 (mUC) cohort. Here in cluster heat map, we found that low ATGscore group exhibited an abundance of activated B, activated CD4, CD8 T cells, and cytotoxic cells, but less infiltrated with Treg, macrophages and mast cells in patients with GU and TCGA II subtypes (Additional file 32: Fig. S11A). Moreover, ssGSEA of specific gene sets showed antigen processing machinery (APM), CD8 T effector, and immune checkpoints signature which represented the immune activation were strikingly enriched in low ATGscore group (Fig. 7b). The immune activation related markers showed a similar trend to function annotation (Fig. 8e-f and Additional file 32: Fig. S11b). However, we found EMT signature was slightly activated in low ATGscore group, but Pan-F-TBRS and WNT-target signatures were strikingly enriched in high ATGscore group (Fig. 7b and Additional file 32: Fig. S11c). And most EMT/TGF-b signalling pathways related markers were not differential expressed between two groups, which indicated that ATGscore might not distinguish stromal pattern in IMvigor210 (mUC) cohort. The results of the correlation matrix are almost identical to above findings (Fig. 7c). As IMvigor210 (mUC) cohort contained a complete information of immune phenotypes, we were amazed to find that the patients with immune-inflamed phenotype exhibited lowest ATGscore, while patients with immune-desert phenotype displayed high ATGscore (Kruskal-Wallis test, p = 2.2e-6, Fig. 7d). The numbers of patients with immune-inflamed phenotype were nearly two times in low HPXscore group when compared with that in high HPXscore group (Fisher’s exact tests, p < 0.00001, Fig. 7e). All of these definitely confirm our findings and
hypothesis that autophagy was able to shape the TIME infiltration and distinguish the immune phenotype.

Although ATGscore showed no correlation with TMB in TCGA-HNSCC cohort, here we were surprised to find that there existed a negative correlation between TMB and ATGscore in IMvigor210 (mUC) cohort (Wilcoxon test, p = 0.036, Fig. 8a and Fisher's exact tests, p = 0.033886, Fig. 8b). Kaplan-Meier survival curves revealed that survival benefit of patients with high TMB was superior to patients with low TMB (Log-rank test, p < 0.001, Fig. 8c). Moreover, we combine the information of ATGscore and TMB to find patients with low ATGscore as well as high TMB exhibited a tremendous survival advantage to all other subgroups (Log-rank test, p < 0.001, Fig. 8d). Then we comprehensively explore the role of ATGscore in the whole IMvigor210 (mUC) cohort as it contained so many information associated with immunotherapy response. The alluvial diagram was utilized to show distribution changes of these features according to ATGscore of individual patient in IMvigor210 (mUC) cohort (Fig. 7f). We found that GU subtype and TCGA II subtype, which displayed high somatic mutation and more likely responded to ICIs treatment, demonstrated lowest ATGscore when compared with other molecular subtypes in the Lund and TCGA classification system respectively (Kruskal-Wallis test, p = 8e-8, Additional file 32: Fig. S11d; Kruskal-Wallis test, p = 2.6e-5, Additional file 32: Fig. S11f). And numbers of patients with GU subtype and TCGA II subtype in low ATGscore group were over two times of that in high ATGscore group (Fisher’s exact test, p = 0.010413, Additional file 32: Fig. S11e; Fisher’s exact test, p = 0.116173, Additional file 32: Fig. S11g). Furthermore, the correlation between ATGscore and immune checkpoint PD-L1 located on tumour cells (TC) or immune cells (IC) were also measured. Surprisingly, we found that most of patients with IC2, which was correlated with better clinical outcome of immunotherapy, concentrated on low ATGscore group, while IC0 strikingly accumulated in high ATGscore group (Fisher’s exact test, p = 0.000136, Fig. 8g). Moreover, patients with IC2 exhibited lowest ATGscore and patients with IC0 showed highest ATGscore (Kruskal-Wallis test, p = 5.2e-9, Fig. 8h). Then we found no difference in ATGscore among patients with TC0-TC2 (Kruskal-Wallis test, p = 0.34, Fig. S11h; Fisher’s exact test, p = 0.728113, Fig. S11i). In addition, we were delighted to find that patients with low ATGscore were more likely to be ICIs treatment responders (CR/PR), while patients with high ATGscore tended to non-responders (SD/PD), (Wilcoxon test, p = 0.0046, Additional file 32: Fig. S11j; Kruskal-Wallis test, p = 0.00037, Fig. 8j). Moreover, the numbers of patients with ICIs immunotherapy responders (CR/PR) was over two times in low ATGscore group to that in high ATGscore group (Fisher’s exact test, p = 0.0055, Additional file 32: Fig. S11k; Fisher’s exact test, p = 0.006009, Fig. 8i). All our findings implied that there exist distinct autophagy regulation patterns in tumours, which could shape the TIME infiltration and immune phenotypes, as well as predict the clinical outcome of ICIs immunotherapy.

**Discussion**

In recent years, the key topics on cancer research have transferred from focusing on the tumour itself to studying the interaction between the tumour and its surrounding environment, which is commonly referred to tumour microenvironment (TME). Tumour microenvironment, where the tumour cells arise and live, includes not only the tumour cells themselves, but also their surrounding stroma, microvasculature,
and a variety of cells including immune cells, stromal cells, fibroblasts, and various other cells, as well as the biological molecules such as cytokines and chemokines they secreted [50]. They cooperated with each other to form chronic inflammatory, immunosuppressive, and tumour promotion environment so that tumour cells can escape from immune surveillance and survive against attack of effector immune cells [51].

Under normal circumstances, autophagy degraded and recycled cytoplasmic components to maintain protein synthesis and other necessary metabolic functions, which is considered to be an endogenous defence mechanism [52–54]. But autophagy could also promote or inhibit tumour progression through many ways depending different backgrounds, which is thought to be a “double-edged sword” in tumours. Autophagy has been reported to regulate the interaction of tumours cells with various substances within surrounding milieu, especially components from the immune system, containing B and T lymphocytes, dendritic cells (DCs), macrophages, and natural killer (NK) cells, as well as the cytokines and immunoglobulins they released [27]. Conversely, immune cells and their secreted cytokines and antibodies could trigger dysfunction of autophagy, which induced or supressed tumorigenesis. However, the autophagy-mediated regulation of the immune system might strengthen or attenuate the effects of immunotherapy. Autophagy was reported to regulate the interaction of tumours cells with various substances within surrounding milieu, especially components from the immune system, containing B and T lymphocytes, dendritic cells (DCs), macrophages, and natural killer (NK) cells, as well as the cytokines and immunoglobulins they released [27]. Conversely, immune cells and their secreted cytokines and antibodies could trigger dysfunction of autophagy, which induced or supressed tumorigenesis. However, the autophagy-mediated regulation of the immune system might strengthen or attenuate the effects of immunotherapy. Autophagy was reported to enhance the effect of immunotherapy by exposing antigens to antigen-presenting cells (APCs) and CTL to initiate and execute the tumour recognition and elimination. Otherwise, a growing number of studies have demonstrated that autophagy could also attenuate the effect of immunotherapy by inducing the activation of immunosuppressive atmosphere to hamper the effector immune cells to kill tumour cells [55]. These all indicated that appropriate induction or inhibition of autophagy may propose a prospective therapeutic strategy when combined with chemotherapy, radiotherapy, and immunotherapy. But comprehensive and systematic analysis of correlation between autophagy and tumour microenvironment has not been fully identified yet, which impedes the clinical development of autophagy based activators or inhibitors.

In this study, we gathered ATGs to determine autophagy regulation patterns through unsupervised consensus clustering. Five distinct autophagy regulation patterns (ATGclusters) were identified with differential ATGs expression, survival benefit, tumour immune microenvironment (TIME) infiltration, and functional annotation. In TCGA-HNSCC cohort, ATGclusterA/B exhibited high TIME infiltration, while ATGclusterC/D/E was relatively less infiltrated with immune cells. Kaplan-Meier survival curves showed that the prognosis of patients with ATGclusterB/E were robustly better than patients with ATGclusterA/C/D, which was not consistent with the findings in TIME infiltration. We noticed that ATGclusterA was both infiltrated with effector immune and immunosuppressive cells, while ATGclusterB/E was highly infiltrated with activated CD8 T cells and cytotoxic cells and less infiltrated with Treg cells, macrophages, and mast cells. So we inferred that highly infiltrated immunosuppressive cells could counteract and impair effector immune cells to distinguish and eradicate abnormal tumour cells. As the opposite function of CTL and Treg cell in tumour immunity, a combined assessment of CTL and Treg infiltration have comprehensively studied. The ratio of CTL/Treg has finally been recognized as the independent prognostic factor in many tumour types [56, 57]. Here in our study, we found that CTL/Treg ratio was significantly higher in ATGclusterB/E than that in ATGclusterA/C/D, which well
explains the mismatch of immune infiltration and survival analysis. Moreover, function annotation demonstrated ATGclusterA was not only enriched in signalling pathways associated with inflammation, but also induced in angiogenesis, EMT, and pan-fibroblast TGFβ response signalling pathways, which indicated the stromal status was relatively activated in this pattern. The stromal status, which could reversibly shift between “loose” and “dense” status, is the key checkpoint for appropriate localization and migration of T cells into tumour parenchyma. As activated in stromal status, effector immune cells cannot effectively penetrate the stroma surrounding core tumour islets, leaving a large number of CTL trapped in the extracellular matrix, thus failing to execute their antitumor role of immune surveillance and eradication, allowing the tumour to continue to progress in ATGclusterA, which was the characteristic of immune-excluded phenotype [58]. In addition to lack of activated and priming T-cell, ssGSEA of KEGG, HALLMARK and specific signatures all revealed that immune tolerance and ignorance was fully induced in ATGclusterC/D, which was more likely to be to immune-desert phenotype [59]. Moreover, we also found despite highly infiltrated with effector immune cells, the antigen processing machinery, CD8 T effector, and immune checkpoint signature, which were representative for immune activation, was also strikingly enriched in ATGclusterB/E, which was featured as immune-inflamed phenotype. Moreover, the expression of immune activation, stromal activation, MHC molecules, and immune checkpoints was distributed as the same trend to function annotation. Recently, many teams have defined the non-inflamed or immune-suppressed tumours with immune-excluded and immune-desert phenotypes as “cold” tumour and tumours with immune-inflamed phenotype as “hot” tumour, which might be responsible for clinical response of immune-checkpoints inhibitors (ICIs) immunotherapy [60, 61]. Next, we systematically collected transcriptome datasets of HNSCC across GEO and ArrayExpress database and merged them as the meta-HNSCC cohort. We were amazed to find that all above results could be precisely validated in the meta-HNSCC cohort. All of these suggested that distinct autophagy regulation patterns associated with signalling pathway enrichment, TIME infiltration, and immune phenotypes do exist in HNSCC, which provides the possibility of the combination of autophagy activator or inhibitors with ICIs.

Accumulated evidence demonstrated that tumour patients with immune-inflamed phenotype will obtained a durable responses and better overall survival when receiving with ICIs treatment [59]. But we also noticed that not all patients benefit from ICIs-targeting therapy, with an estimated response rate only modestly above the historical 10% response rate to traditional chemotherapies [41]. Immune tolerance to these tumours is still a major impediment in cancer immunotherapy. To improve the efficacy of immunotherapy and prolong survival from immunotherapy, we need to clarify the underlying mechanisms of immune tolerance, which disenabled the role of effector immune cells. Many important factors has been identified to influence the immune tolerance, such as hypo-infiltration of effector immune cells into tumour parenchymal, imbalance between effector immune and immunosuppressive cells, abnormalities in MHC molecules function and expression, and lack of tumour antigen or epitopes exposure leading to failure of antigen processing to T cells [62]. As highly correlated with TIME infiltration and immune phenotypes, the association between autophagy regulation pattern and tumour mutation load was further investigated. Previous study demonstrated that the recognition of neo-antigens exposure, which mainly triggered by somatic nonsynonymous mutations, was essential for initiating the
antigen processing and activation of the adaptive tumour immunity cascade. Moreover, tumour mutation
burden (TMB), which could be easily assessed to replace the overall neo-antigen detection, has been
determined as a potential biomarker for predicting the clinical outcome of ICI treatment [63, 64]. In
TCGA-HNSCC cohort, we found that TMB was differential distributed among five distinct autophagy
regulation patterns, where ATGclusterB showed lowest TMB and ATGclusterD showed highest TMB,
which is not consistence with the results from TIME infiltration and function annotation. But next we were
surprised to find patients with high TMB were associated with worse prognosis than patients with low
TMB in TCGA-HNSCC cohort. This indicated that TMB is harmful in the TCGA-HNSCC cohort which was
against our common sense. However, we then found that DNA damage repairing signalling pathways
were significantly enriched in ATGclusterB/E, which might be another cause of good prognosis in this
pattern.

Then autophagy phenotype related genes, which was exacted from the DEGs among distinct autophagy
regulation patterns, were submitted to LASSO cox regression analysis to establish a set scoring system to
evaluate and quantify autophagy regulation pattern in individuals, which was referred to the autophagy
phenotype related signature (ATGscore). ATGscore was found to be differential distributed among five
distinct autophagy regulation patterns. Moreover, in TCGA-HNSCC, microarray-HNSCC, and IMvigor210
(mUC) cohorts, we were pleasantly to find that low ATGscore group was significantly enriched in immune
activation relevant signalling pathways and deactivated in stromal relevant signalling pathways, which
was characteristic of “hot” tumour. Thus the opposite phenomenon was seen in high ATGscore group,
which referred to “cold” tumour. In addition, we validated these in IMvigor210 (mUC) cohort to find that
immune-infamed phenotype exhibited lowest ATGscore, while immune-desert and immune-excluded
phenotypes displayed higher ATGscore, which indicated the successful of model construction. All above
results revealed that ATGscore was not only a reliable tool to assess autophagy regulation pattern, but
also was potent to effectively evaluate TIME infiltration and immune phenotype in individual patient. In
addition, we found patients with good clinical outcomes, as well as low malignancy clinicopathological
traits and molecular subtypes more likely to accumulate in low ATGscore group, while the opposite
patterns were observed in high ATGscore group. In IMvigor210 (mUC) cohort, we were amazed to find that
patients with GU and TCGA II molecular subtypes, as well as IC2 phenotypes, which was reported that
more likely to have an good response to ICI-targeting immunotherapy, were robustly concentrated in the
low ATGscore group, and rarely observed in with high ATGscore group [34, 41]. Moreover, we found that
TMB was not different between high and low ATGscore groups. But we next found that high ATGscore
group were more likely to be mutation in TP53, which again indicated the role of ATGscore in prediction
the ICI immunotherapy response as mutation status of TP53 could have predictive value in
immunotherapy in patients with HNSCC [65]. Although TMB did not work well in TCGA-HNSCC cohort,
here in IMvigor210 (mUC) cohort we found that ATGscore was negative correlated with TMB and patients
with low ATGscore group were more likely to be patients with high TMB, which was consistent with
findings in GU molecular subtype with high mutation load. Moreover, Kaplan-Meier survival curves
showed that combination of ATGscore and TMB could significantly improve the predictive value when
compared with that of TMB or ATGscore alone where patients with low ATGscore and high TMB exhibited
best prognosis. Finally we found patients with low ATGscore were more likely to benefit from ICIs treatment and ICIs targeting immunotherapy responders exhibited a lower ATGscore. According to all above findings, autophagy regulation patterns and ATGscore was found to be significantly correlated with three main factors: pre-existing activated CTL or immunoreactivity, activation of the EMT/TGFβ signalling pathway or stromal status, and tumour neo-antigen or TMB levels, to influence the clinical outcome of ICIs immunotherapy.

**Conclusion**

In conclusion, we comprehensively and systematically assessed distinct autophagy regulation patterns and established a set scoring system ATGscore which could represent them and was associated with the TIME infiltration, immune phenotypes, molecular subtypes, genetic variation, clinical outcome of ICIs targeting immunotherapy, etc. More importantly, this study has yielded novel insights of combination of autophagy-based inducer or inhibitor with various therapeutic strategies such as immunotherapy for clinical application in HNSCC.

**Abbreviations**

DCA: decision curve analysis; DEGs: differential expression genes; ECM: extracellular matrix; EMT: epithelial-mesenchymal transition; FDR: false discovery rate; GEO: Gene Expression Omnibus; GSEA: Gene Set Enrichment Analysis; GSVA: Gene Set Variation Analysis; HR, hazard ratio; HNSCC: head and neck squamous cell carcinoma; ICIs: immune-checkpoint inhibitors; LASSO: Least absolute shrinkage and selection operator; mUC: metastatic urothelial cancer; ssGSEA: single-sample gene set enrichment analysis; TCGA: The Cancer Genome Atlas; TIME: tumour immune microenvironment; TMB: tumour mutation burden; TME: tumour microenvironment; TNM: Tumour Node Metastasis.

**Declarations**

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**Authors’ contributions**

BM made substantial contributions to conception and design of the research. BM, HL and MZ carried out data collection and analysis. BM, RC and RY wrote the paper. BM, HL, MZ, RC and RY edited the manuscript and provided critical comments. All authors read and approved the final manuscript.

**Data accessibility**

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request. The data that support the findings of this study were derived from the following resource: TCGA GDC data portal (https://portal.gdc.cancer.gov/), GEO (https://www.ncbi.nlm.nih.gov/geo/) and ArrayExpress database (www.ebi.ac.uk/arrayexpress/) as well as package “IMvigor” in R (http://research-pub.gene.com/IMvigor210CoreBiologies).

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Compliance with ethical standards

Conflict of interest

The authors declare that they have no conflict of interest.

Informed consent

All the analysed data used in our research were collected from a public database, such as TCGA, GEO, and ArrayExpress as well as IMvigor210 package and so on; therefore, informed consent was not required for this analysis.

Ethical approval

Since this was a retrospective study and all the data were collected from a public database TCGA, GEO, and ArrayExpress as well as IMvigor210 package and so on, therefore, ethical approval was not required.

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Consensus clustering of autophagy related genes in HNSCC. (a) Overview of workflow and study design. (b) Consensus matrices of patients in TCGA-HNSCC cohort for k=2-5 using 1000 iterations of unsupervised consensus clustering method (K-means) to ensure the clustering stability. (c) Hierarchical clustering of ATGs based on Euclidean distance and Ward linkage in TCGA-HNSCC cohort. The
ATG clusters and vital status were shown as patient annotations. Rows represented ATGs, and columns represented HNSCC samples. Red represented genes were relative upregulated and blue represented genes were relative downregulated. (d) Kaplan-Meier survival curves for distinct autophagy regulation patterns in TCGA-HNSCC cohort using log-rank test (Log-rank test, p = 0.0054).
TIME infiltration and biological process characteristics in distinct autophagy regulation patterns. (a) Hierarchical clustering of TIME landscape in TCGA-HNSCC cohort. Rows represented relative amount of each immune cell, and columns represented HNSCC samples. Red represented immune cells were relative upregulated and blue represented immune cells were relative downregulated. ATGclusters, ATGscore, and clinical features such as vital status, follow-up treatment outcome, primary therapy outcome, new tumour event, neoplasm cancer status, pathological TNM stage, pathological T stage, pathological N stage, smoking status, HPV status, alcohol histology, mutation in TP53, mutation in CDKN2A were used as sample annotations. (b) Difference in the enrichment of specific signatures to represent biological processes related with stromal-activation and immune-activation among five distinct autophagy regulation patterns in TCGA-HNSCC cohort. (c) Difference in the expression of MHC molecules (HLA molecular) among five distinct autophagy regulation patterns in TCGA-HNSCC cohort. In Fig. 2B and C, the upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference among five distinct autophagy regulation patterns was tested by the Kruskal-Wallis test. The asterisks represented the statistical p value (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001). (d) Differences in TMB among different autophagy regulation patterns in TCGA-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value. Kruskal-Wallis test was used to compare the statistical difference between each pattern. (Kruskal-Wallis test, p = 4.3e-11). (e) Distribution of top 30 variant mutated genes among five distinct autophagy regulation patterns in TCGA-HNSCC cohort. The genetic alterations types include frame shift del, frame shift ins, in frame del, in frame ins, missense mutation, multi hit, nonsense mutation and splice site. The upper bar-plots indicated ATGscore, TMB and overall survival (OS) time. The number on the left and right bar-plots showed the mutation frequency of each gene. ATGcluster, and clinical features such as alcohol histology, HPV status, neoplasm cancer status, pathological N stage, pathological T stage, pathological TNM stage, primary therapy outcome, smoking status, and vital status were shown as patient annotations.
Figure 3

Characterization of autophagy regulation patterns in meta-HNSCC cohort. (a) Consensus matrices of patients in meta-HNSCC cohort for k=2-5 using 1000 iterations of unsupervised consensus clustering method (K-means) to ensure the clustering stability. (b) Hierarchical clustering of autophagy related genes based on Euclidean distance and Ward linkage in meta-HNSCC cohort. ATGclusters, vital status, and projects were shown as patient annotations. Rows represented ATGs, and columns represented
HNSCC samples. Red represented genes were relative upregulated and blue represented genes were relative downregulated. (c) Kaplan-Meier survival curves showed the difference in prognosis advantage among five distinct autophagy regulation patterns in meta-HNSCC cohort (Log-rank test, p = 0.0022). (d) Hierarchical clustering of TIME landscape in meta-HNSCC cohort. Rows represented relative amount of each immune cell, and columns represented HNSCC samples. Red represented relative upregulated and blue represented relative downregulated of each immune cell. ATGclusters, vital status, and projects were used as sample annotations. (e) Difference in the enrichment of immune activation, stromal activation, and DNA damage repair (DDR) relevant signatures among five distinct autophagy regulation patterns in meta-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference among five distinct autophagy regulation patterns was tested by the Kruskal-Wallis test. The asterisks represented the statistical p value (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001).
Establishment of autophagy phenotype related signature (ATGscore) in TCGA-HNSCC cohort. (a) Kaplan-Meier survival curves showed the difference in prognosis between high and low ATGscore groups in TCGA-HNSCC cohort (Log-rank test, p < 0.00001). (b) Bar-plot demonstrated that patients with high ATGscore were more likely to be dead and patients with low ATGscore were more inclined to alive. Red indicated patients were dead and Blue indicated patients were alive. Dot-plot indicated that survival time of patients with high ATGscore were less than patients with low ATGscore. Red indicated dead patients.
and Blue indicated living patients. (c) Difference of ATGscore among five distinct autophagy regulation patterns in TCGA-HNSCC cohort. Kruskal-Wallis test was used to compare the statistical difference between each pattern. (Kruskal-Wallis test, $p = 2.2e-16$). (d) Difference of ATGscore among four molecular subtypes in TCGA-HNSCC cohort. Kruskal-Wallis test was used to compare the statistical difference between each pattern. (Kruskal-Wallis test, $p = 2.2e-16$). (e) Kaplan-Meier survival curves showed the difference in prognosis among four molecular subtypes in TCGA-HNSCC cohort (Log-rank test, $p = 0.0326$). (f) Correlation matrix of ATGscore, candidate ATGs and the relative amount of TIME immune cells. The blue indicated positive correlation and red indicated negative correlation. The asterisks represented the statistical $p$ value (*$p < 0.05$; **$p < 0.01$). (g) Difference in the enrichment of specific signatures to represent biological processes related with stromal-activation, immune-activation, and DDR between high and low ATGscore groups in TCGA-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference between high and low ATGscore groups was tested by Wilcoxon test. The asterisks represented the statistical $p$ value (*$P < 0.05$; **$P < 0.01$; ***$P < 0.001$; ****$P < 0.0001$). (h) Correlation between ATGscore and stromal-activation, immune-activation, and DDR relevant signatures in TCGA-HNSCC cohort. Negative correlation was marked with red and positive correlation was marked with blue.
Figure 5

Validation of autophagy phenotype related signature (ATGscore) in microarray-HNSCC cohort (a) Hierarchical clustering of TIME landscape in microarray-HNSCC cohort. Rows represented relative amount of each immune cell, and columns represented HNSCC samples. Red represented immune cells were relative upregulated and blue represented immune cells were relative downregulated. ATGclusters, ATGscore, vital status, and projects were used as sample annotations. (b) Kaplan-Meier survival curves showed the difference in prognosis between high and low ATGscore groups in microarray-HNSCC cohort (Log-rank test, p = 0.0196). (c) The correlation chord chart showed the mutual correlation between ATGscore and several prominent immune-checkpoint-relevant genes (CD274, CD80, CD86, CTLA-4, LAG-3, HAVCR2, PDCD1, and TNFRSF9). (d) Correlation matrix of ATGscore and stromal-activation, immune-activation, and DDR relevant signatures in microarray-HNSCC cohort. (e) Difference in the enrichment of specific signatures to represent biological processes related with stromal-activation, immune-activation, and DDR between high and low ATGscore groups in microarray-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference between high and low ATGscore groups
was tested by Wilcoxon test. The asterisks represented the statistical p value (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001).

Figure 6

Association between ATGscore and clinicopathological characteristics in TCGA-HNSCC cohort. (a-f) Differences of ATGscore among indicated clinical features in TCGA-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value. Wilcoxon test or Kruskal-Wallis test were used to compare the statistical difference of pathological TNM stage (a), HPV status (b), smoking status (c), neoplasm cancer status (d), primary therapy outcome (e), and vital status (f). (g) Alluvial diagram showed the dynamic changes of ATGscore, ATGclusters, and indicated clinical features in TCGA-HNSCC cohort. The waterfall plot showed the distribution of top 30 highly variant mutated genes between high and low ATGscore groups. The genetic alterations types were indicated in the waterfall plot annotation. Each column represented individual patients. The upper bar plots showed ATGscore, TMB and OStime. The number on the left and right bar-plots showed the mutation frequency of each gene. ATGscore, ATGcluster, and clinical features such as alcohol histology, HPV status, smoking status, pathological N stage, pathological T stage, pathological TNM stage, neoplasm cancer status, primary therapy outcome, and vital status are shown as patient annotations. (i)
Kaplan-Meier survival curves for mutation in TP53 and TP53 wild type (WT) in TCGA-HNSCC cohort using log-rank test (Log-rank test, p = 0.0062). (j) Differences of ATGscore between patients with TP53 mutation and TP53 WT in TCGA-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value. The statistical difference was tested by Wilcoxon test. (Wilcoxon test, p = 2.2e-12). (k) Kaplan-Meier survival curves for mutation in CDKN2A and CDKN2A wild type (WT) in TCGA-HNSCC cohort using log-rank test (Log-rank test, p = 0.2827). (l) Differences of ATGscore between patients with CDKN2A mutation and CDKN2A WT in TCGA-HNSCC cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value. The statistical difference was tested by Wilcoxon test. (Wilcoxon test, p = 0.034).

Figure 7
Characteristics of ATGscore in IMvigor210 (mUC) cohort. (a) Kaplan-Meier survival curves showed the difference in prognosis between high and low ATGscore groups in IMvigor210 (mUC) cohort (Log-rank test. p < 0.001). (b) Difference in the enrichment of specific signature between high and low ATGscore groups in IMvigor210 (mUC) cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference between high and low ATGscore groups was tested by Wilcoxon test. The asterisks represented the statistical p value (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001). (c) Correlation between ATGscore and specific signature in IMvigor210 (mUC) cohort. Blue indicated positive correlation and red indicated negative correlation. (d) Differences of ATGscore among different immune phenotypes in IMvigor210 (mUC) cohort. The Kruskal-Wallis test was used to compare the statistical difference between different immune phenotypes (Kruskal-Wallis test, p = 2.2e-6). (e) The proportion of immune phenotypes between high and low ATGscore groups in IMvigor210 (mUC) cohort. The statistical difference was measured with the Fisher’s exact test. (Fisher’s exact test, p < 0.00001). (f) Alluvial diagram showed the dynamic changes of ATGscore, ATGclusters, and indicated immunotherapy related features in IMvigor210 (mUC) cohort.
Figure 8

ATGscore was an efficiency tool to predict clinical response of ICI immunotherapy. (a) Difference of TMB between high and low ATGscore groups in IMvigor210 (mUC) cohort. The TMB was log2 transformed. The statistical difference was measured with the Wilcoxon test. (Wilcoxon test, p = 0.036) (b) The proportion of TMB between high and low ATGscore groups in IMvigor210 (mUC) cohort. The statistical difference was measured with the Fisher's exact test. (Fisher's exact test, p = 0.03386). (c) Kaplan-Meier survival curves showed the difference in prognosis between high and low TMB groups in IMvigor210 (mUC) cohort (Log-rank test. p < 0.001). (d) Kaplan-Meier survival curves showed the difference in prognosis advantage among four groups stratified by ATGscore and TMB in IMvigor210 (mUC) cohort (Log-rank test, p < 0.001). (E-F) Immune activation relevant markers (e) and immune-checkpoints relevant markers (f) were differentially expressed between high and low ATGscore groups in
IMvigor210 (mUC) cohort. The upper and lower ends of the boxes represented interquartile range of values. The lines in the boxes represented median value and the black dots showed outliers. The statistical difference between high and low ATGscore groups was tested by Wilcoxon test. The asterisks represented the statistical p value (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001). (g) The proportion of PD-L1 expression on immune cells (IC) subtypes between high and low ATGscore groups in IMvigor210 (mUC) cohort. The statistical difference was measured with the Fisher’s exact test. (Fisher’s exact test, p = 0.000136). (h) Differences of ATGscore between different PD-L1 expression on immune cells (IC) in IMvigor210 (mUC) cohort. The Kruskal-Wallis test was used to compare the statistical difference between different IC0-IC2 groups (Kruskal-Wallis test, p = 5.5e-9). (i) The proportion of patients with response to ICIs immunotherapy between low or high ATGscore groups in IMvigor210 (mUC) cohort. CR, complete response; PR, partial response; SD, stable disease; PD, progressive disease. CR/PR was identified as responder and SD/PD was identified as non-responder. The statistical difference was measured with the Fisher's exact test. (Fisher’s exact test, p = 0.006009). (j) Differences of ATGscore between different ICIs immunotherapy clinical response groups. The statistical difference was measured with the Kruskal-Wallis test (Kruskal-Wallis test, p=0.00037).

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