A six-gene signature related with tumor mutation burden for predicting lymph node metastasis in breast cancer

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Background: Breast cancer (BC) is one of the most common cancers worldwide and patients with lymph node metastasis always suffer from a worse prognosis. Tumor mutation burden (TMB) has been reported as a potential predictor for tumor behaviors. However, the correlation between TMB and lymph node metastasis of BC remains unclear. This study aimed to explore TMB-related biomarkers to predict the lymph node metastasis in BC patients.

Methods: A total of 949 BC patients with RNA-seq data, mutation data and clinical data were obtained from The Cancer Genome Atlas (TCGA) database. We visualized mutation data by “maftools” package. We calculated TMB of each patient and investigated its association with lymph node metastasis. BC patients were divided into lymph node positive and negative groups and we respectively identified TMB-related and lymph node-related differentially expressed genes (DEGs) to figure out intersected genes. Functional enrichment analysis and protein-protein interaction (PPI) network were performed to observe relevant biological functions. We constructed a TMB-related signature for predicting lymph node metastasis through Logistic regression analysis. A validation database (GSE102484) from the Gene Expression Omnibus (GEO) database was downloaded to verify the accuracy.

Results: Single nucleotide polymorphism (SNP) occupied the highest proportion in variant types while C>T appeared most frequently in single nucleotide variant (SNV), TMB was regarded as negatively correlated with lymph node metastasis in BC (P=0.003). We identified 125 common DEGs through venn diagram, which were enriched in vesicle localization, calcium signaling pathway and salmonella infection. A TMB-related signature based on six genes (BAHD1, PPM1A, PQLC3, SMPD3, EEF1A1 and S100B) had reliable efficacy for predicting lymph node metastasis in BC and was proven as an independent predictive factor. The accuracy of this signature was further validated by GSE102484 database.

Conclusions: Our results indicated that TMB was associated with lymph node metastasis of BC. We built a TMB-related signature consisting of six genes which might function as a novel biomarker for predicting lymph node metastasis in BC.

Keywords: Breast cancer (BC); tumor mutation burden (TMB); lymph node metastasis; predictive signature; bioinformatics

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**Introduction**

Breast cancer (BC) is one of the leading malignancies among females worldwide, with 279,100 estimated new cases and 42,690 estimated deaths in 2020 (1). In spite of the development of medical technologies, large amounts of BC patients were still diagnosed with lymph node metastasis (2). Lymph node status was identified as an important prognostic factor in BC and patients with lymph node metastasis had a worse survival outcome than those without metastatic status (3,4). Moreover, BC patients with lymph node metastasis often suffer from sentinel lymph node biopsy or axillary dissection, taking a risk of many serious complications, such as lymphedema, chyle leak, seroma formation and so on (5-7). Therefore, it is urgent for clinicians to search for potential biomarkers to predict the lymph node metastasis in BC.

Tumor mutation burden (TMB) refers to the total quantity of non-synonymous point mutations per coding region in the tumor gene (8). Recently, TMB has been regarded as an effective biomarker for predicting response to immune checkpoint inhibitors in multiple cancer types, such as melanoma, non-small cell lung cancer and advanced urothelial cancer (9-11). Besides, an increasing number of researchers begin to explore the relationship between TMB and BC. It was discovered that TMB was closely related with immune-mediated survival in BC (12). According to Park, high TMB was associated with good overall survival and functioned as an independent prognostic factor in HER2-positive metastatic BC (13). Barroso-Sousa suggested that BC patients with high TMB were more likely to benefit from PD-1 inhibitors (14).

Current researches mostly focused on the role of TMB in predicting clinical outcomes of BC, the correlation between TMB and tumor biological characteristics, such as lymph node metastasis, remained unclear. It was reported that tumor progression, such as lymphatic metastasis was regulated by both tumor escape mechanism and dysfunction of immune system, which highlighted the significant role of immune system in tumor biological behaviors (15,16). Moreover, tumor infiltrating lymphocytes (TILs) were observed to participate in the regulation of lymph node metastasis, including CD8(+) T cells and Foxp3(+) Tregs (17). Meanwhile, as to TMB, many researchers considered TMB as an effective biomarker for predicting immune response owing to its potential function of increasing neoantigens and inducing TILs infiltration (18,19). According to Mei, high TMB level in BC patients was positively related with the amount of TILs (20). However, whether TMB can influence the immune cell infiltration and further regulate lymph node metastasis in BC is still lack of evidence, which deserves further investigation.

With the development of sequencing and chip technologies, an increasing number of public databases are emerging, such as The Cancer Genome Atlas (TCGA) database and Gene Expression Omnibus (GEO) database. Researchers around the world upload their research data to these public databases, promoting information sharing and accelerating medical development. In the present study, we obtained mutation data, transcriptome data and clinical data from the TCGA database. The TMB value of each BC sample was calculated and the correlation between TMB and lymph node metastasis was investigated. We identified TMB-related and lymph node-related differentially expressed genes (DEGs) respectively to extract common DEGs. Functional enrichment analysis and protein-protein interaction (PPI) network were performed to explore the biological roles of common DEGs. A TMB-related signature including six genes was constructed for predicting the lymph node metastasis in BC and external verification was further conducted. After a comprehensive analysis, we believed that the TMB-related signature had potential in predicting the lymph node metastasis in BC. We present the following article in accordance with the STROBE reporting checklist (available at http://dx.doi.org/10.21037/tcr-20-3471).

**Methods**

**Data acquisition**

We obtained the BC mutation data, transcriptome data and clinical data from TCGA database (https://portal.gdc.cancer.gov/), comprising 460 lymph node negative samples and 489 lymph node positive samples, which served as the training dataset. We employed “maftools” R package to analyze the Masked Somatic Mutation data, which were processed via VarScan software. Transcriptome data were acquired from HTseq-FPKM platform while clinical data contained age, gender, pathological stage and AJCC-TNM stage. We also obtained a gene expression profile (GSE102484) from the Gene Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/geo/), comprising 300 lymph node negative samples and 383 lymph node positive samples, which functioned as the validation dataset.
The platform for GSE102484 was the GPL570 [HG-U133_Plus_2] Affymetrix Human Genome U133 Plus 2.0 Array. All data were publicly available and patients with data deficiency were excluded. The clinical characteristics of BC patients in training and validation datasets are summarized in Table 1. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

**Table 1: Clinical characteristics of breast cancer patients in training and validation datasets**

| Variable | TCGA database | GSE102484 database |
|----------|---------------|---------------------|
| Age      |               |                     |
| ≤58 years| 481 (50.7)    | 541 (79.2)          |
| >58 years| 468 (49.3)    | 142 (20.8)          |
| Gender   |               |                     |
| Female   | 938 (98.8)    | 683 (100.0)         |
| Male     | 11 (1.2)      | 0 (0)               |
| Stage    |               |                     |
| I        | 162 (17.1)    | 175 (25.6)          |
| II       | 550 (58.0)    | 328 (48.0)          |
| III      | 208 (21.9)    | 174 (25.5)          |
| IV       | 16 (1.7)      | 6 (0.9)             |
| T        |               |                     |
| T1       | 248 (26.1)    | 276 (40.4)          |
| T2       | 557 (58.7)    | 377 (55.2)          |
| T3       | 109 (11.5)    | 24 (3.5)            |
| T4       | 34 (3.6)      | 6 (0.9)             |
| N        |               |                     |
| N0       | 460 (48.5)    | 300 (43.9)          |
| N1       | 317 (33.4)    | 214 (31.3)          |
| N2       | 112 (11.8)    | 87 (12.7)           |
| N3       | 60 (6.3)      | 82 (12.1)           |
| M        |               |                     |
| M0       | 802 (84.5)    | 582 (85.2)          |
| M1       | 18 (1.9)      | 101 (14.8)          |

TMB was calculated by the ratio of total mutation amounts to human exon length (38 Mb) and was regarded as the frequency of tumor gene mutations, consisting of insertions or deletions, coding errors and base substitutions. We conducted correlation analysis between TMB and different lymph node status. Wilcoxon rank-sum test and Kruskal-Wallis test were respectively used for two or more groups.

**TMB value calculation and correlation with lymph node metastasis**

We divided BC samples into lymph node positive and negative groups to identify lymph node-related DEGs.

**Differential analysis, functional enrichment analysis and PPI network construction**

We divided BC samples into lymph node positive and negative groups to identify lymph node-related DEGs.
Furthermore, based on the median TMB value, TMB-related DEGs between high and low TMB groups were respectively acquired in lymph node positive and negative groups. Differential analysis above was performed through “limma” package with the criterion of P value <0.05 and false discovery rate (FDR) <0.05, which was further visualized by volcano plots. Subsequently, the intersection between lymph node-related and TMB-related DEGs was extracted as common DEGs through “VennDiagram” package and further visualized via “UpSetR” package. Besides, in order to explore biological functions of common DEGs, Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis were applied via “org.Hs.eg.db”, “clusterProfiler”, “ggplot2” and “enrichplot” packages. Moreover, the PPI network of common DEGs was conducted by STRING database (https://string-db.org/) and presented via Cytoscape software.

**TMB-related signature construction and external verification**

We firstly conducted univariate Logistic regression analysis to screen for common DEGs associated with lymph node metastasis with P value <0.001. Then, we brought selected DEGs into multivariate Logistic regression analysis to calculate regression coefficient ($\beta$) and establish a TMB-related signature through the formula: risk score = $\Sigma (\beta \times$ the expression of gene). We applied the formula above to calculate risk score of each sample and divided all samples into high and low risk groups according to the median risk score. The correlation between risk score and clinical variables was explored via Chi-square test and the expression of six hub genes between high and low risk groups was displayed via “pheatmap” package. Meanwhile, the receiver operating characteristic (ROC) curve was applied to assess the predictive value via “pROC” package and the evaluation of independent predictive factors was also performed through Logistic regression analysis. Furthermore, we brought all independent predictive factors into a nomogram through “rms” package and assessed its predictive accuracy via the calibration plot. A validation dataset (GSE102484) was further utilized for external verification.

**Expression level and relapse-free survival (RFS) analysis of hub genes between patients with different lymph node status**

We investigated comparative analysis between six hub genes and different lymph node status via Wilcoxon rank-sum test for two groups and Kruskal-Wallis test for more groups. Moreover, we performed RFS analysis of six hub genes respectively in lymph node positive and negative groups via the Kaplan-Meier Plotter mRNA BC database (http://kmplot.com/analysis/index.php?p=service). We divided patients into high and low expression groups based on the best cutoff and calculated relevant log-rank P values respectively in lymph node positive and negative groups. P value <0.05 was identified as statistically significant.

**Correlation of hub genes with TMB and cBioPortal analysis**

We performed correlation analysis between six hub genes and TMB via Spearman correlation analysis and visualized the results via the “ggplot2”, “ggpubr” and “ggExtra” packages. Furthermore, we used cBioPortal database (http://www.cbioportal.org/) to acquire gene alteration frequencies and types of six hub genes in different BC studies.

**Statistical analysis**

We utilized R software (Version 3.6.3) and SPSS software (Version 24.0) to perform data analysis. The construction of TMB-related signature was conducted by univariate and multivariate Logistic regression analysis. Survival analysis was performed by Kaplan-Meier method and log-rank test. We applied differential analysis via “limma” package and conducted correlation analysis through Spearman correlation analysis. We investigated comparative analysis of continuous variables through Wilcoxon rank-sum test for two groups and Kruskal-Wallis test for more groups. Meanwhile, comparative analysis of categorical variables was conducted by Chi-square test. A P value <0.05 was identified as statistically significant.

**Results**

**Visualization of BC mutation profiles**

We utilized waterfall plot to exhibit high-frequency mutated genes in BC, such as TP53 (34%), PIK3CA (33%), TTN (16%), CDH1 (13%) and GATA3 (12%) (Figure 1A). Besides, missense mutation ranked first in variant classification, single nucleotide polymorphism (SNP) appeared more frequently than insertion or deletion and
Figure 1 Summary of mutation information in BC. (A) Visualization of mutation profiles in BC samples. Waterfall plot exhibiting mutation types of each gene in each sample, with barplot representing mutation burden. (B) Mutation information distinguished by different classifying standards and variant burden, classification in inclusive samples. (C) Co-occurance and mutual exclusion among top 25 mutated genes. BC, breast cancer.
C>T was considered as the most common single nucleotide variant (SNV). Meanwhile, the median variation was estimated to be 29 and variant types were also shown via box plot (Figure 1B). Furthermore, the co-occurrence and mutual exclusion among top 25 mutated genes were also displayed (Figure 1C).

**TMB correlated with lymph node metastasis**

Clinical information was acquired and merged with TMB value to further explore the correlation between TMB and different lymph node status of BC. We discovered that TMB was negatively correlated with lymph node metastasis in BC (P=0.003) (Figure 2A). Moreover, N1 group had lower TMB level than N0 group (P<0.001) and N2 group (P=0.018) (Figure 2B).

**Identification of DEGs**

We performed differential analysis to screen for lymph node-related and TMB-related DEGs between lymph node positive and negative groups. According to TMB-related DEGs, 8,751 DEGs (2,522 up-regulated and 6,229 down-regulated) and 9,354 DEGs (3,636 up-regulated and 5,718 down-regulated) were respectively identified in lymph node positive and negative groups, while 763 lymph node-related DEGs (455 up-regulated and 308 down-regulated) were respectively identified in lymph node positive and negative groups, while 763 lymph node-related DEGs (455 up-regulated and 308 down-regulated) was confirmed. Differential expression was displayed through volcano plots (Figure 3) and 125 common DEGs were extracted as the intersection of three datasets above via Venn diagram (Figure 4A).

**Functional enrichment analysis and PPI network of common DEGs**

After acquiring 125 common DEGs, we utilized the Upset diagram for further detailed analysis. Based on median TMB value, 16 simultaneously up-regulated and 104 simultaneously down-regulated common DEGs were identified in lymph node positive and negative groups, while 5 other common DEGs were regulated reversely (Figure 4B). We conducted functional enrichment analysis to further investigate significant pathways of 125 common DEGs. GO analysis indicated that common DEGs were enriched in vesicle localization, the membrane of lysosome, vacuole and lytic vacuole, GTPase regulator activity among biological process (BP), cellular component (CC) and molecular function (MF) categories respectively (Figure 4C). KEGG analysis demonstrated that calcium signaling pathway and salmonella infection were mainly enriched (Figure 4D). Subsequently, we searched 125 common DEGs in STRING database with the interaction score of 0.150 and constructed a PPI network with disconnected nodes hided, which was further visualized via Cytoscape software (Figure 4E). The PPI network of common DEGs contained 101 nodes and 498 edges, with simultaneously up-regulated, simultaneously down-regulated and reversely regulated common DEGs marked in pink, blue and green respectively (Figure 4F).

**Construction and verification of the TMB-related signature**

We firstly performed univariate Logistic regression analysis
to select 29 genes associated with lymph node metastasis among 125 common DEGs, with P value <0.001. We further brought selected 29 genes into multivariate Logistic regression analysis and utilized “Forward: LR” to establish a TMB-related signature of six genes for predicting lymph node metastasis (Table 2). The risk score of TMB-related signature was calculated via regression coefficient(β) and expression level of six hub genes, which was identified as follows: risk score= exp_{BAHD1}*(0.054)+ exp_{PPM1A}*(0.069)+ exp_{PQLC3}*(0.024)+ exp_{SMPD3}*(0.104)+ exp_{EEF1A1}*(-0.001)+ exp_{S100B}*(-0.010) (Figure 5). Subsequently, we calculated risk score of each sample and divided all samples into high and low risk groups. As shown in the heat map, risk score was significantly associated with age (P<0.05), stage (P<0.001) and lymph node status (P<0.001) in TCGA database. Meanwhile, the expression of BAHD1, PPM1A, PQLC3 and SMPD3 was higher in high risk group while EEF1A1 and S100B presented opposite trends (Figure 6A,B). Furthermore, the ROC curve indicated reliable predictivity of six-gene signature with the area under curve (AUC) of 0.656 in TCGA database and 0.561 in GSE102484 database (Figure 6C,D). Univariate and multivariate Logistic regression analysis proved the TMB-related signature as an independent predictive factor for lymph node metastasis in BC (Table 3, Table 4). Besides, a nomogram containing independent predictive factors was constructed and the calibration plot confirmed its effective predictivity in TCGA database and GSE102484 database (Figure 6E,F,G).

**Differential expression of six hub genes between patients with different lymph node status**

We performed comparative analysis between six hub genes and different lymph node status in BC. The results demonstrated that four hub genes were positively correlated with lymph node metastasis, including BAHD1 (P<0.001), PPM1A (P<0.001), PQLC3 (P<0.001) and SMPD3 (P<0.001), while EEF1A1 (P<0.001) and S100B (P<0.001) exhibited negative correlation (Figure 7A). Furthermore, more detailed comparative analysis was conducted. The

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Figure 3 Identification of TMB-related and lymph node-related DEGs. (A) The volcano plot of TMB-related DEGs in lymph node positive and negative groups. (B) The volcano plot of lymph node-related DEGs. TMB, tumor mutation burden; DEGs, differentially expressed genes.
Figure 4 Identification, functional enrichment analysis and PPI network construction of common DEGs. (A) The venn diagram showing 125 common DEGs extracting from TMB-related and lymph node-related DEGs. (B) Based on median TMB value, the intersection relation of 125 common DEGs between lymph node positive and negative groups was displayed via the Upset diagram. (C) Gene Ontology (GO) functional analysis of 125 common DEGs, including biological process (BP), cellular component (CC) and molecular function (MF) categories. (D) Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways of 125 common DEGs. (E) The PPI network of 125 common DEGs was constructed in STRING database. (F) The PPI network was further visualized via Cytoscape software with simultaneously up-regulated, simultaneously down-regulated and reversely regulated common DEGs marked in pink, blue and green respectively. PPI, protein-protein interaction; TMB, tumor mutation burden; DEGs, differentially expressed genes.
Table 2 Construction of a TMB-related signature including six genes in TCGA database (Logistic regression analysis)

| Gene      | Univariate analysis | Multivariate analysis |
|-----------|---------------------|----------------------|
|           | OR (95% CI)         | P value              | OR (95% CI)         | P value | Coefficient |
| BAH1      | 1.088 (1.041–1.137) | <0.001               | 1.055 (1.007–1.105) | 0.023   | 0.054       |
| PPM1A     | 1.106 (1.059–1.155) | <0.001               | 1.072 (1.024–1.122) | 0.003   | 0.069       |
| PQLC3     | 1.028 (1.014–1.042) | <0.001               | 1.024 (1.010–1.038) | 0.001   | 0.024       |
| SMPD3     | 1.160 (1.077–1.249) | <0.001               | 1.110 (1.032–1.193) | 0.005   | 0.104       |
| EEF1A1    | 0.999 (0.999–1.000) | <0.001               | 0.999 (0.999–1.000) | 0.009   | -0.001      |
| S100B     | 0.980 (0.968–0.991) | <0.001               | 0.990 (0.980–0.999) | 0.039   | -0.010      |

OR, odds ratio; CI, confidence interval.

Figure 5 Construction of TMB-related signature. (A) Forest plot of univariate Logistic regression analysis for six hub genes. (B) Forest plot of multivariate Logistic regression analysis for six hub genes. (C) Regression coefficients of six hub genes. TMB, tumor mutation burden.

expression levels of BAH1 (P<0.001), PPM1A (P<0.001), PQLC3 (P<0.001) and SMPD3 (P<0.001) were found lower in N0 group than in N1 group, while EEF1A1 (P=0.008) and S100B (P=0.006) presented the opposite expression. Meanwhile, PPM1A (P<0.001), PQLC3 (P<0.001) and SMPD3 (P=0.002) had lower expression in N0 group than in N2 group, while EEF1A1 (P=0.005) and S100B (P=0.002) presented the opposite expression. Besides, BAH1 (P=0.002) had lower mRNA expression in N0 group than in N3 group (Figure 7B).

RFS analysis of six hub genes between patients with different lymph node status

Based on the Kaplan-Meier Plotter, we divided patients into high and low expression groups via the best cutoff and conducted RFS analysis of six hub genes respectively in lymph node positive and negative BC. The results indicated that higher expression levels of BAH1 (P=0.020), PPM1A (P=0.049), PQLC3 (P=0.009), SMPD3 (P=0.037) and EEF1A1 (P=0.034) were associated with better RFS in lymph node negative group while S100B (P=0.260) showed no association with RFS (Figure 8A). Meanwhile, higher expression levels of BAH1 (P=0.009), PPM1A (P<0.001), PQLC3 (P=0.043), SMPD3 (P=0.021) and EEF1A1 (P=0.015) were associated with better RFS in lymph node positive group while higher S100B (P=0.033) expression had worse prognosis (Figure 8B).

Correlation of six hub genes with TMB and cBioPortal analysis

We investigated the correlation between six hub genes and TMB. The results demonstrated that the expression levels of BAH1 (P<0.001), PPM1A (P<0.001), PQLC3 (P<0.001), SMPD3 (P<0.001) and EEF1A1 (P<0.001) were...
Figure 6 Verification of TMB-related signature. (A) Correlation analysis between risk score and clinical variables in TCGA database. (B) Correlation analysis between risk score and clinical variables in GSE102484 database. (C) Receiver operating characteristic (ROC) curve analysis of the TMB-related signature in TCGA database. (D) ROC curve analysis of the TMB-related signature in GSE102484 database. (E) A nomogram constructed for predicting lymph node metastasis, which contained independent predictive factors. (F) The calibration plot in TCGA database. (G) The calibration plot in GSE102484 database. *, P<0.05; **, P<0.01; ***, P<0.001. TMB, tumor mutation burden.
Table 3 Univariate and multivariate Logistic regression analysis of predictive factors for lymph node metastasis in TCGA database

| Variable | Univariate analysis | Multivariate analysis |
|----------|---------------------|----------------------|
|          | OR (95% CI)         | P value              | OR (95% CI)         | P value              |
| Risk Score | 2.638 (2.068–3.364) | <0.001*              | 2.664 (2.013–3.526) | <0.001*              |
| Age (≤58 years) | 1.442 (1.093–1.901) | 0.010*              | 1.947 (1.376–2.753) | <0.001*              |
| Gender (female) | 0.426 (0.109–1.661) | 0.219               | –                  | –                   |
| Stage (Stage I-II) | 0.018 (0.008–0.042) | <0.001*              | 0.019 (0.008–0.047) | <0.001*              |
| T (T1) | 0.416 (0.301–0.576) | <0.001*              | 0.619 (0.424–0.903) | 0.013*               |
| M (M0) | 0.065 (0.009–0.492) | 0.008*              | 1.943 (0.208–18.151) | 0.560                |

Age was classified as ≤58 years and >58 years; Gender was classified as female and male; Stage was classified as Stage I-II and Stage III-IV; T was classified as T1 and T2-T4; M was classified as M0 and M1; OR, odds ratio; CI, confidence interval; *, P value <0.05 has statistical significance.

Table 4 Univariate and multivariate Logistic regression analysis of predictive factors for lymph node metastasis in GSE102484 database

| Variable | Univariate analysis | Multivariate analysis |
|----------|---------------------|----------------------|
|          | OR (95% CI)         | P value              | OR (95% CI)         | P value              |
| Risk score | 35.579 (2.865–441.816) | 0.005*              | 21.034 (1.167–379.142) | 0.039*              |
| Age (≤58 years) | 1.268 (0.876–1.837) | 0.208               | –                  | –                   |
| Stage (Stage I-II) | 0.012 (0.004–0.037) | <0.001*              | 0.015 (0.005–0.048) | <0.001*              |
| T (T1) | 0.242 (0.175–0.335) | <0.001*              | 0.372 (0.258–0.538) | <0.001*              |
| M (M0) | 0.285 (0.172–0.473) | <0.001*              | 0.934 (0.478–1.825) | 0.842                |

Age was classified as ≤58 years and >58 years; Stage was classified as Stage I-II and Stage III-IV; T was classified as T1 and T2-T4; M was classified as M0 and M1; OR, odds ratio; CI, confidence interval; *, P value <0.05 has statistical significance.

negatively correlated with TMB while S100B (P=0.18) showed no significant correlation with TMB (Figure 9A). Besides, we utilized cBioPortal database to acquire gene alteration status of six hub genes in different BC studies. The results suggested that gene alteration frequency of PQLC3 and SMPD3 was less than 2% while PPM1A and S100B had less than 4.5% gene alterations. Meanwhile, BAHD1 had less than 3% gene alterations while the gene alteration frequency of EEF1A1 was less than 5%. Moreover, deep deletion was the most common alteration type in BAHD1, SMPD3 and EEF1A1, while amplification ranked first in PPM1A, PQLC3 and S100B (Figure 9B).

Discussion

With the rapid development of sequencing technology, an increasing number of researchers begin to investigate the potential role of TMB in the occurrence and development of BC. Voutsadakis reported that the TMB expression was higher in HER2-positive subtype than in luminal and triple-negative subtypes (21). Similar to Voutsadakis, Xu suggested that elevated TMB expression was identified in HR-negative or HER2-positive BC (22). Besides, many sequencing researches were performed in triple-negative breast cancer (TNBC) and discovered the association between high TMB expression and better prognosis. Karn indicated that early TNBC patients with high TMB were more likely to achieve pathological complete response (pCR) and TMB was regarded as an independent predictive factor for pCR (23). According to Barroso-Sousa, higher TMB expression was associated with better progress free survival in metastatic TNBC patients treated with anti-PD-1/L1 (24).

However, the majority of current studies paid attention to the predictive function of TMB for clinical outcomes in BC, the correlation between TMB and lymph node
Figure 7 Correlation between six hub genes and lymph node status in BC. (A) BAHD1, PPM1A, PQLC3 and SMPD3 were positively correlated with lymph node metastasis while EEF1A1 and S100B were negatively correlated with lymph node metastasis. (B) The expression levels of six hub genes had statistical difference in different lymph node groups. BC, breast cancer.
Figure 8 Relapse-free survival (RFS) analysis of six hub genes in lymph node positive and negative groups. (A) Among the lymph node negative group, the higher expression levels of BAH1, PPM1A, PQLC3, SMPD3 and EEF1A1 were associated with better RFS while S100B had no association with RFS. (B) Among the lymph node positive group, the higher expression levels of BAH1, PPM1A, PQLC3, SMPD3 and EEF1A1 were associated with better RFS while higher S100B expression had worse RFS.
Figure 9 Correlation of six hub genes with TMB and cBioPortal analysis. (A) BAHD1, PPM1A, PQLC3, SMPD3 and EEF1A1 were negatively correlated with TMB while S100B had no significant correlation with TMB. (B) According to the cBioPortal database, deep deletion was the most common alteration type in BAHD1, SMPD3 and EEF1A1, while amplification ranked first in PPM1A, PQLC3 and S100B.
metastasis is still ambiguous. Among BC patients, those with lymph node metastasis were under a threat of wider surgical margin, more serious complications and higher chemotherapy toxicity, suffering from a worse prognosis (25-27). Therefore, it is of great significance to further investigate the relationship between TMB and lymph node metastasis in BC.

It was demonstrated that tumor-mediated immune dysfunction played an important role in accelerating tumor progression, including lymphatic metastasis (28-30). According to Zuckerman, the down-regulation of immune-related genes and up-regulation of tumor-promoting genes were observed in BC patients with lymph node metastasis (31). Besides, Kohrt demonstrated that CD1a dendritic cells were lower in tumor-involved axillary nodes than tumor-free axillary nodes in BC and dendritic cells in axillary nodes were closely related with disease-free survival (32). As to TMB, it has been known as a potential predictor for tumor-related immunological response due to the emerging neoantigens from gene alterations and increased infiltration of immune cells (33,34). Based on above researches, we hypothesize that TMB may take part in the regulation of lymph node metastasis in BC through influencing immune cell infiltration and more detailed researches should be performed for further verification.

In our study, we downloaded mutation, transcriptome and clinical data from TCGA database and calculated TMB value of each patient. We merged TMB value and clinical data and discovered that TMB was negatively associated with lymph node metastasis in BC. Subsequently, TMB-related and lymph node-related DEGs were identified respectively and we extracted common DEGs from them. We further conducted functional enrichment analysis and constructed a PPI network of common DEGs. The membrane of lysosome and vacuole, vesicle localization and GTPase regulator activity were mainly enriched in GO analysis, while calcium signaling pathway and salmonella infection were mainly enriched in KEGG analysis. The roles of vesicle localization, calcium signaling and salmonella infection in regulating BC metastasis have been explored in some researches (35-37). Furthermore, we established the TMB-related signature of six genes, including BAHD1, PPM1A, PQLC3, SMPD3, EEF1A1 and S100B, via univariate and multivariate Logistic regression analysis. The predictive accuracy of this signature was assessed reliable in both TCGA database and GSE102484 database.

The function of six hub genes in BC was previously investigated in many researches. Singh verified that SMPD3 participated in the encoding of neutral sphingomyelinase 2 (nSMase2) enzyme and nSMase2 regulated BC invasion via enhancing the exosome-mediated secretion of miR-10b (38). Lin discovered that EEF1A1 had gene alterations in 27% BC patients and could protect tumor cells from proteotoxic injuries via enhancing heat shock responses (39). Li identified that the promoter regulatory element of EEF1A1 was regulated by MALAT1 and over-expressed MALAT1 played an important role in the metastasis of BC (40). In addition, the decreased EEF1A1 expression via curcumol attributed to the suppression of BC metastasis (41). The function of S100B for inhibiting tumor migration was found in ER-negative BC and high expression of S100B was associated with better distant metastases-free survival in BC (42). Over-expressed PPM1A was demonstrated to restrain the progression of triple negative BC through suppressing cell cycle and reducing the phosphorylation of CDK and Rb (43). The researches of BAHD1 in BC remained rare while Goryca identified that the mutation of BAHD1 took part in promoting the metastasis of colorectal cancer (44). The role of PQLC3 in tumorigenesis and progression was still unclear and deserved further investigation.

In this study, we discovered that TMB was negatively correlated with lymph node metastasis and constructed a TMB-related signature based on six genes for predicting lymph node metastasis in BC, which might provide novel sights for clinicians. With the further investigation of TMB, we would like to explore other potential roles of TMB-related signature in BC, such as predicting immune response, survival status and so on. However, there are still some limitations in our study. On one hand, the samples involved in our study is limited and clinical trials of large samples should be further conducted to evaluate the predictivity of the TMB-related signature. On the other hand, our study was lack of experiments in vitro or in vivo, which should be performed to verify the biological functions of six hub genes.

Conclusions

In summary, our study suggested that TMB was negatively correlated with lymph node metastasis in BC. A TMB-related signature including six genes was further constructed and validated for predicting lymph node metastasis in BC, which may provide guidance for clinicians.
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Footnote

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/tcr-20-3471). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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