Research Article

BUBs Are New Biomarkers of Promoting Tumorigenesis and Affecting Prognosis in Breast Cancer

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Background. A tumor occurs because of abnormal cell multiplication caused by many variables like a significant disturbance in the regulation of cell growth and the instability of chromosome mitosis. Budding uninhibited by benzimidazoles 1 (BUB1), BUB1 mitotic checkpoint serine/threonine kinase B (BUB1B), and budding uninhibited by benzimidazoles 3 (BUB3) are key regulators of mitosis, and their abnormal expression is highly correlated with breast cancer (BrCa), sarcoma, hepatic carcinoma, and other malignant tumors. However, the occurrence of BUBs (BUB1, BUB1B, and BUB3) and the development of BrCa have not been systematically explained. Methods. Find out the target gene by looking up literature on PubMed and CNKI. Using the R software, TCGA, GEO, Kaplan-Meier Plotter, TIMER, and other databases, we studied the level of transcription, genetic changes, and physiological functions of BUBs in BrCa patients and their relationship with the origin, development, prognosis, immunity, and drug resistance of BrCa patients. Findings. We found that the high expression level of BUBs in BrCa tissues proposed a poor prognosis. The multivariate Cox regression analysis suggested that BUB1B and BUB3 might be independent prognostic factors of BrCa. In addition, the Metascape functional enrichment analysis showed that BUBs may be involved in the composition of the spindle, chromosome, and other structures and play a role in mitosis, sister chromatid separation, and other processes. Pathway enrichment suggests that BUBs may affect the cell cycle and lead to abnormal proliferation. Meanwhile, we also found that BUB3 can negatively regulate B lymphocytes, and BUB1 and BUB1B inhibit immune responses by promoting the secretion level of checkpoint molecules of the immune system, leading to immune escape of tumor cells. Conclusion. We speculate that BUB1, BUB1B, and BUB3 may be therapeutic targets for BrCa patients and also provide new therapeutic strategies for BrCa treatment.

1. Introduction

According to the latest global cancer data for 2020, released by the International Agency for Research on Cancer, BrCa has overtaken lung cancer as cancer with the highest number of new cases. What is more worrying is that the incidence of BrCa in China has been increasing year by year in recent years. BrCa has become the primary reason for cancer-associated mortality for females in third-world nations. As with other malignancies, aneuploidy is a common feature of BrCa and affects its behavior. Aneuploidy is related to the inappropriate activity of the spindle assembly checkpoint (SAC). SAC is a monitoring mechanism. When the integrity of the genome is impaired, a sister chromatid separation error during mitosis will lead to aneuploidy, which leads to the occurrence of cancer [1, 2]. When it comes to spindle mitosis, the BUB gene family has important functions to perform. BUB1 is necessary for SAC signaling and correct alignment of chromosomes [3]. As a paralogous gene of BUB1, BUB1B suppresses late anaphase-promoting complex or cyclosome (APC/C) activity via interfering with the binding of the cell division cycle 20 (CDC20) to APC/C and monitors centromere protein E- (CENPE-) dependent kinetochore activity [4]. BUB3...
facilitates the establishment of a robust termination on the
dipole association, which is required for BUB1 to be local-
ized to the kinetochore. When SAC is activated, the BUB1/3 complex performs its function in inhibiting late APC/C and inhibits APC/C ubiquitin ligase activity by phosphory-
ating its activator CDC20 [5].

The Genome Project has accumulated a large amount of
biological information, and the task of bioinformatics is to
mine and utilize this information. Therefore, bioinformatics
will play an extremely important role in the “postgenome”
era, which will help to understand the information of the
human genome. Many cancers, including BrCa, have been
deeply explored in terms of genome and transcriptome [6,
7]. Through bioinformatics, the combination of gene expres-
sion profiles and clinical data to mine potential biomarkers
and provide new targets for cancer treatment has become a
hot topic. Therefore, we explored the clinical significance of
BUBs in BrCa patients through public databases such as
TCGA and GEO, to find new targets for BrCa diagnosis
and treatment.

2. Materials and Methods

2.1. Data Acquisition. The level 3 HTSeq-FPKM format
RNAseq data of BrCa was downloaded from TCGA database
(https://portal.gdc.cancer.gov/), and a total of 1109 tumor
samples and 113 normal samples were obtained. Meanwhile,
the clinical information of BrCa patients was downloaded
from TCGA database. We found dataset GSE155478 in the
GEO database (https://www.ncbi.nlm.nih.gov/geo/) and
divided it into two groups according to drug sensitivity: the
drug resistance group (3 cases) and the sensitive group (3
cases).

2.2. Analyses of Gene Expression. We examine the differential
expression of BUBs in BrCa tissues and normal samples in
TCGA database by using the R package ggplot2. Then we
still use the ggplot2 R package, select the level 3 HTSeq-
FPKM format RNAseq data and clinical data in TCGA BrCa
project, transform the FPKM format RNAseq data into log2,
and examine the relationship between the expression of
BUBs and pathological stages. UALCAN is a sophisticated
and dynamic web resource (http://ualcan.path.uab.edu/
analysis.html) website to detect the differential expression
levels of BUBs in different molecular types in BrCa.

2.3. Survival Prognostic Analysis. Using the Kaplan-Meier
Plotter, researchers can determine whether 54000 genes
(mRNA, miRNA, and protein) have an impact on survival
in 21 cancer types, notably lung cancer \( n = 3,452 \), gastric
cancer \( n = 1,440 \), ovarian cancer \( n = 2,190 \), and breast
cancer \( n = 7,830 \). The datasets were compiled by data from
GEO, EGA, and TCGA. The major goal of the tool is to cre-
ate and validate survival biomarkers using meta-analytic
techniques. First, we select Start KM Plotter for BrCa, then
click Use Multiple Genes in the blank column of Gene Sym-
bol to input BUB1, BUB1B, and BUB3 genes, and then select
median to divide patients. In the Probe set options, User
Selected Probe Set was selected, and OS and RFS were
selected successively in Survival to obtain the overall survival
(OS) as well as relapse-free survival (RFS) data of BUB tran-
scription in BrCa patients, respectively. To do additional
research on the possible prognostic significance of BUBs in
BrCa patients, uni- and multivariate Cox regression analyses
were performed first, and the forest plot R tool was used to
create forest plots to represent the hazard ratio, \( p \) value,
and 95% confidence interval for each variable. Because of
the multivariate Cox proportional hazard analysis, it was
necessary to apply the R software package rms to create a
nomogram that could forecast the overall recurrence rate
in one, three, and five years. The nomogram provides a
graphical representation of these factors, and the prognostic
risk of a single patient can be calculated by the points related
to each risk factor.

2.4. Gene Enrichment Analysis in Relation to BUBs. Using
the stat R package, the top 20 genes most associated with
BUBs’ expression in BrCa were obtained from TCGA data-
base. The duplicate genes were removed, and 45 genes were
obtained for further enrichment analysis. The GEPIA online
software (http://gepia.cancer-pku.cn/) was used to analyze
the correlation between BUB1, BUB1B, and BUB3. BUBs
and closely linked genes were investigated by using STRING
(http://string-db.org/), which was used to construct a pro-
tein interaction network for BUBs and closely associated
genes. Subsequently, the Cytoscape software (http://www.
cytoscape.org/) was used for visual analysis of the protein-
protein interaction (PPI) network. It is more important for
the overall stability of the network and if in case the nodes
have a higher degree of connection than others. CytoHubba
is a Cytoscape plugin that may be used to determine the
degree of each protein node in a network of proteins. The
top ten genes found in our study were termed “hub genes.”
In addition, we used the Metascape (http://metascape.org)
website for the Kyoto Encyclopedia of Genes and Genomes
(KEGG) and Genetic Ontology (GO) analyses. The latter is
composed of three components: biological process (BP),
molecular function (MF), and cellular component (CC),
and it can be used to predict the functions of genes that
are closely associated to BUBs, while the former can describe
the gene pathways related to BUBs. Items having a mini-
um count of 5, a \( p \) value less than 0.01, and an enrichment
factor of enrichment more than 2.0 were collected for fur-
ther analysis. We use the GeneMANIA (http://genemania
.org/) database to construct the BUB interactive network.

2.5. Immune-Related Analysis. Using the GSVA package in
the R software and GSEA calculation method, we calculated
the correlation between BUB1, BUB1B, BUB3, and 24 kinds
of common immune cells. Then we use the TIMER (https://
cistrome.shinyapps.io/timer/) database, first for the system-
atric analysis of different cancer types of immune invasion
sites, and then choose the Survival module that analyzes
the 6 kinds of common immune infiltrating cells in their
relationship with prognosis. Then we use the TISIDB
(http://cis.hku.hk/TISIDB/index.php) database to further
analyze the relations between the BUBs and B lymphocytes;
TISIDB is a website of tumor-immune system relationship,
Disease Markers

Figure 1: Plot between BrCa and normal breast tissue; there is a difference in the expression of members of the BUB family. ***p < 0.001.

and it integrates a variety of data types. To verify the above conclusion again, we picked five typical B lymphocyte gene markers from the Systems website of R&D (https://www.rndsystems.com/cn/resources/cell-markers/immune-cells) and used them in our experiments. The correlation between them and BUBs was analyzed by using the Correlation module of TIMER. Then, we selected seven major immune checkpoint molecules. In TISIDB, the expression level of these molecules was at the y-axis, and the abundance of immune checkpoint molecules was at the y-axis. It was decided to utilize scatter plots to demonstrate the relationship between the quantity of each immunological checkpoint molecule and the transcription of BUBs.

2.6. Drug Resistance Analysis. Using the 45 coexpressed genes, most relevant to BUB expression in BrCa obtained above, and the differentially expressed genes in the GSE155478 dataset related to BrCa resistance found from the GEO database (conforming to critical criteria: adjusted p < 0.05 and |logFC| ≥ 1.5 genes are differentially expressed genes) takes the intersection to obtain differential genes coexpressed with BUBs in BrCa resistant cells.

3. Results

3.1. BUBs Expressed Differentially in BrCa and Normal Breast Tissue. BrCa and normal breast tissues were examined for BUB’s expression levels. When compared to normal breast tissue, BrCa tissues had significantly higher levels of BUB1, BUB1B, and BUB3 (Figure 1).

3.2. Expression of BUBs in the Clinicopathology of BrCa Patients. After discussing the expression level of BUBs in BrCa and normal tissues, we continued to study the expression of BUBs in BrCa based on cancer in various stages. The expression level of BUBs in BrCa tissues with different pathological stages was higher than that in negative control sample tissues (Figure 2(a)). We further analyzed the expression of BUBs in BrCa patients with different T/N stages. The expression of BUBs in different T/N stages was significantly greater than the level found in negative control tissue samples (Figure S1). In the four common molecular types of BrCa, Her-2 overexpression, luminal (A, B), and three negative breast cancer (TNBC), the expression level of BUBs was higher than that of normal tissues (Figure 2(b)). BUB1 was statistically significant in all genotyping; therefore, the increase in BUB1 expression can infer that the molecular typing is poor. In conclusion, these findings imply that the high level of BUBs may be related to the progression of BrCa.

3.3. BUBs’ Clinical Prognostic Value in BrCa. To investigate the prognostic significance of BUBs in BrCa, we analyzed the survival data of BrCa patients using the “Kaplan-Meier Plotter database.” We can see the correlation between BUBs differential expression and clinical outcomes of BrCa patients and investigate the prognostic relevance of BUB mRNA expression in patients with BrCa. The high expression of BUB1 and BUB1B resulted in shortened OS and poor prognosis in BrCa patients (Figure 3(a)). The risk ratio of BUB1 was 1.64 (p = 2.3E−07), while the expression of BUB3 did not affect the OS rate (p = 0.49 > 0.05). The RFS of BrCa patients was significantly and negatively correlated with the expression of BUBs. The risk ratio of BUB1 was 1.74 (p < 1E−16), BUB1B was 1.77 (p < 1E−16), and BUB3 was 1.16 (p = 0.0041) (Figure 3(b)).

3.4. Nomograms Predict Survival in BrCa Patients. In the univariate Cox regression analysis, we found that the high expression of BUB1 (hazard ratio (HR) = 1.16125, 95% confidence interval (CI): 1.00955, 1.33575, p = 0.03634) and BUB1B (HR = 1.19911, 95% CI: 1.03011, 1.39585, p = 0.01915) suggested that BrCa patients had poor progression-free survival (PFS). The worse T stage (95% CI: 1.3815, 2.05909, HR = 1.6866, p < 0.0001) and pathological stage (95% CI: 1.6819, 2.65185, HR = 2.11191, p < 0.0001) indicate the worse PFS of BrCa patients (Figure 4(a)). Therefore, BUB1, BUB1B, T stage, and pathological stage are related to PFS. Then, we conducted the multivariate Cox regression analysis and obtained the variables that can be used as nomograms (Figure 4(c)): BUB3...
Pathologic stage
- Normal
- Stage I
- Stage II
- Stage III
- Stage IV

**Figure 2**: The BrCa clinical stage and molecular type were used to determine the expression of BUBs family members. (a) BUBs are expressed during the clinical stages of BrCa. (b) The expression of BUBs in different molecular types of BrCa. ns, $p \geq 0.05$; **$p < 0.01$; ***$p < 0.001$. 

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Disease Markers
other clinical factors. However, the di 

BUB3 and pathological stage are variables independent of 

the multivariate Cox regression analysis suggest that BUB1B, 

Analysis of BUBs and Their Coexpressed Genes. TCGA data-

3.5. Protein Interaction Network and Functional Enrichment 

(Figure 4(d)). 

curve, the more accurate the model 

the patient in the next 1, 3, and 5 years can be inferred. 

then adds the individual scores of each variable to get the 

unique score assigned to each variable at various levels and 

denoted by the point in the illustration. This re 

value range, and the length of the line segment indicates 

effect of BUB3 on PFS is opposite to that of BUB1B. A scale 

the factor 

is used to indicate the line segment corresponding to each 

variable in the nomogram, which denotes the variable 

expression. (a) The e 

ect of BUBs on the OS of patients with BrCa. (b) BUBs and their 

correlations between the BUBs (Figure 5(d)), 

pressed genes (Figures 5(a) 

base was used to find out the coexpression genes with BUBs, 

the correlation coefficients were arranged in descending 

order, the first 20 genes were selected, and 45 coexpression 

genes were obtained by removing duplicate genes. We sepa-

rately studied the correlation between BUBs and their coex-

pressed genes (Figures 5(a)–5(c)). Additionally, we 

examined the correlations between the BUBs (Figure 5(d)), 

which shows that there is a strong correlation among BUB 

gene families. Then we constructed a PPI protein interaction 

etwork between BUBs and 45 coexpressed genes to explore 

the potential relationship among them (Figure S2) using the 

CytoHubba plugin in the Cytoscape software; then we 

the top ten hub genes with the highest 

interactions (Figure 6). It can be seen from the figure that 

BUB1 and BUB1B are at the core of the complex protein 

interaction network. 

To further clarify the functional role of the BUB gene 

family, we used the Metascape online tool to conduct an 

alysis of the function and enrichment of pathways of 

BUBs and their closely related genes. The function of BUBs 

family, we used the Metascape online tool to conduct an 

alysis of the function and enrichment of pathways of 

BUBs and their closely related genes. The function of BUBs 

was evaluated from three aspects: BP, MF, and CC (Figure 7 

) (Figure 4(b)). The results of 

ranks 

P 

HR = 1.74 (1.57 – 1.93) 

logrank 
P < 1E – 16 

HR = 1.74 (1.57 – 1.93) 

logrank 
P < 1E – 16 

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alysis of the function and enrichment of pathways of 

BUBs and their closely related genes. The function of BUBs 

was evaluated from three aspects: BP, MF, and CC (Figure 7 

(a)). We found that BUBs are mainly involved in the composi-

tion of agglutinating the chromosome centromeric region,
<table>
<thead>
<tr>
<th>Variable</th>
<th>$p$-value</th>
<th>Hazard ratio (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Uni_cox</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BUB1</td>
<td>0.03634</td>
<td>1.16125 (1.00955, 1.33575)</td>
</tr>
<tr>
<td>BUB3</td>
<td>0.0755</td>
<td>0.79229 (0.61289, 1.0242)</td>
</tr>
<tr>
<td>BUB1B</td>
<td>0.01915</td>
<td>1.19911 (1.03011, 1.39585)</td>
</tr>
<tr>
<td>Age</td>
<td>0.82444</td>
<td>1.00141 (0.989, 1.01399)</td>
</tr>
<tr>
<td>Gender</td>
<td>0.71099</td>
<td>0.68935 (0.09635, 4.93202)</td>
</tr>
<tr>
<td>Race</td>
<td>0.39337</td>
<td>0.87258 (0.63808, 1.19326)</td>
</tr>
<tr>
<td>$p_T$ stage</td>
<td>&lt; 0.0001</td>
<td>1.6866 (1.3815, 2.05909)</td>
</tr>
<tr>
<td>$p_{TNM}$ stage</td>
<td>&lt; 0.0001</td>
<td>2.11191 (1.6819, 2.65185)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variable</th>
<th>$p$-value</th>
<th>Hazard ratio (95%CI)</th>
</tr>
</thead>
<tbody>
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<td><strong>Mult_cox</strong></td>
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<td></td>
</tr>
<tr>
<td>BUB1</td>
<td>0.63266</td>
<td>0.92314 (0.66502, 1.28143)</td>
</tr>
<tr>
<td>BUB3</td>
<td>0.00861</td>
<td>0.68198 (0.51257, 0.90737)</td>
</tr>
<tr>
<td>BUB1B</td>
<td>0.03457</td>
<td>1.45897 (1.0278, 2.07101)</td>
</tr>
<tr>
<td>Age</td>
<td>0.06600</td>
<td>1.01244 (0.99918, 1.02588)</td>
</tr>
<tr>
<td>Gender</td>
<td>0.54603</td>
<td>0.54152 (0.07393, 3.96674)</td>
</tr>
<tr>
<td>Race</td>
<td>0.29726</td>
<td>0.84478 (0.61518, 1.16009)</td>
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<tr>
<td>$p_T$ stage</td>
<td>0.12247</td>
<td>1.24824 (0.94209, 1.65387)</td>
</tr>
<tr>
<td>$p_{TNM}$ stage</td>
<td>0.00014</td>
<td>1.84021 (1.34445, 2.51876)</td>
</tr>
</tbody>
</table>

**Figure 4:** Continued.
Figure 4: The prognosis of BrCa patients can be predicted using the nomogram model. (a) Regression results from a univariate Cox model are shown in a forest plot. (b) In a forest plot, the results of multivariate Cox regression analysis are displayed. (c) Prediction of BrCa patients’ PFS at one, three, and five years using a nomogram. (d) Evaluation of the nomogram model’s predictive ability.
Figure 5: Continued.
Figure 5: The association between BUBs and coexpressed genes was analyzed. (a) The relationship between BUB1 and the genes that coexpresses. (b) The relationship between BUB1B and the genes that coexpresses. (c) The relationship between BUB3 and the genes that coexpresses. (d) Correlation between BUBs. *** p < 0.001.

Figure 6: The ten hub genes with the greatest degree of linkage are listed.
Figure 7: Continued.
kinetochore, microtubule, spindle pole, and mitotic spindle
and participate in spindle assembly, regulation of mid/late
mitotic transition, regulation of chromosome segregation,
regulation of sister chromatid segregation in mitosis, cell
division, nuclear division, meiotic cell cycle, organelle
fission, and other biological processes. It also possesses
microtubule binding, ATP-dependent activity, protein kinase
binding, and other molecular functions. Pathway enrich-
ment analysis showed that BUBs were involved in the cell
cycle pathway (Figure 7(b)). The GeneMANIA database fur-
ther confirmed the location of the BUB gene family in the
chromosome centromere region and in the chromosome
region and participates in many processes such as chromo-
some separation, sister chromatid separation, and mitosis
(Figure S3).

3.6. Effect of BUB Expression Level on Tumor-Immune
Microenvironment (TIME). Tumor infiltration lymphocyte
(TIL) is closely related to the prognosis of BrCa and subse-
quent immunotherapy [8, 9]. In this study, we investigated
the relationship between BUBs expression and immune cell
infiltration in the BrCa strain (Figures 8(a)–8(c)). In the case
when the correlation coefficient \( |\rho| \geq 0.2 \), BUB1 and BUB1B
were negatively correlated with mast cells, NK cells, cd56
bright NK cells, eosinophils, plasmacytoid dendritic cells,
type 17 T helper cells, and immature dendritic cells, and
BUB1 was positively correlated with type 2 T helper cells,
activated dendritic cells, regulatory T cells, type 1 T helper
cells, and cd56dim natural killer cells; BUB1B was positively
correlated with type 2 T helper cells, activated dendritic cells,
and regulatory T cells; BUB3 was negatively correlated with
dendritic cells, neutrophils, B lymphocytes, macrophages,
cytotoxic cells, type 1 T helper cells, and plasmacytoid den-
dritic cells and positively correlated with type 2 T helper
cells. We also used the TIMER to evaluate the impact of
six common immune infiltrating cells on the prognosis of
BrCa patients (Figure 8(d)). The high expression of B lym-
phocytes is conducive to enhancing patients’ prognosis
\( (p = 0.046) \), and the impact of other immune infiltrating cells
on the prognosis is not statistically significant. Therefore, we
reverified the link between the degree of expression of BUBs
and B lymphocyte infiltration on the TISIDB website
(Figure 8(e)). The negative impact of BUB3 on B lympho-
cytes is greater than the positive impact of BUB1 and
BUB1B. BUB3 may affect the prognosis of patients by affect-
ing B lymphocytes. To further explore the relationship
between BUBs and B lymphocytes, we found five types of
B lymphocytes through the R&D Systems website, selected
their surface markers, and analyzed the correlation between
BUBs and these surface markers (Table 1). BUB1 and
BUB1B had no significant correlation with most markers,
but BUB3 had a negative correlation with these markers,
indicating that BUB3 negatively regulates B lymphocytes in
BrCa. Above, we studied the relationship between BUBs
and TILs.

However, in the TIME, not only TILs but also immune
checkpoint molecules play a role. The immune checkpoint
cell is a regulatory cell that plays an inhibitory
role in the immune system. It is very important to maintain
self-tolerance, prevent autoimmun response, and minimize
tissue damage by controlling the time and intensity of the
immune response. Immune checkpoint molecules expressed
on immune cells will obstruct immune cell function, pre-
venting the body from producing an effective antitumor-
immune response and promoting tumor-immune escape
[10]. To explore whether BUBs have an impact on the
Figure 8: Continued.
Figure 8: Continued.
Figure 8: Continued.
expression of immune checkpoint molecules, we selected seven common immune checkpoint molecules for verification in TISIDB (Figures 9(a)–9(c)). BUB1 and BUB1B are positively correlated with these immune checkpoint molecules, while BUB3 is negatively correlated. BUB1 and BUB1B inhibit the immune response by increasing the expression of immune checkpoint molecules, resulting in immune escape of tumor cells, which is conducive to the occurrence and development of BrCa. In conclusion, BUBs affect TIME through different mechanisms, resulting in a poor prognosis of BrCa patients.

3.7. Expression of BUBs and Their Coexpressed Genes in BrCa Drug Resistance. BrCa is a major public health problem worldwide and is one of the leading causes of death in women. At present, many BrCa treatment drugs have been applied in clinical practice. But long-term use of these drugs can develop resistance, reducing their effectiveness against BrCa and leading to poor survival. In cell development, control of the cell cycle is crucial and plays an important role in the process of tumor drug resistance [9]. Therefore, we searched the GSE155478 dataset in the GEO database to analyze and compare the expression of BUBs and their

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**Table 1: Analysis of correlations between BUBs and a variety of B cell surface markers in BrCa.**

<table>
<thead>
<tr>
<th>B cells</th>
<th>Cell surface markers</th>
<th>BUB1 Cor</th>
<th>BUB1 p value</th>
<th>BUB1B Cor</th>
<th>BUB1B p value</th>
<th>BUB3 Cor</th>
<th>BUB3 p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Follicular B cell</td>
<td>CD19</td>
<td>0.079</td>
<td>8.54E-03</td>
<td>0.023</td>
<td>4.56E-01</td>
<td>-0.172</td>
<td>9.90E-09</td>
</tr>
<tr>
<td></td>
<td>CD20</td>
<td>0.066</td>
<td>2.81E-02</td>
<td>0.027</td>
<td>3.69E-01</td>
<td>-0.124</td>
<td>3.95E-05</td>
</tr>
<tr>
<td></td>
<td>CD23</td>
<td>-0.058</td>
<td>5.58E-02</td>
<td>-0.09</td>
<td>2.93E-03</td>
<td>-0.221</td>
<td>1.40E-13</td>
</tr>
<tr>
<td>Marginal zone B cell</td>
<td>CD19</td>
<td>0.079</td>
<td>8.54E-03</td>
<td>0.023</td>
<td>4.56E-01</td>
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<td>9.90E-09</td>
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<tr>
<td></td>
<td>CD21</td>
<td>0.134</td>
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<td>1.21E-02</td>
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<td>5.69E-10</td>
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<td>CD40</td>
<td>0.04</td>
<td>1.85E-01</td>
<td>-0.021</td>
<td>4.9E-01</td>
<td>-0.231</td>
<td>8.29E-15</td>
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<td>0.076</td>
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<td>-0.185</td>
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<td>-0.185</td>
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<td>1.63E-27</td>
<td>0.251</td>
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Cor: R value of Spearman’s correlation.
coexpressed genes in the Adriaycin- (ADR-) sensitive breast cancer cell line MCF-7 and drug-resistant cell line MCF-7/ADR. The t-test analyzes the difference in BUB gene expression between BrCa MCF-7/ADR and MCF-7 (Figures 10(a)–10(c)). BUBs and most coexpressed genes showed low expression (Table S1), but no high expression genes, indicating that low expression of BUBs plays a role in the mechanism of drug resistance in BrCa (Figure 10(d)). To avoid cell death, cancer cells have been found to evade drug-induced mitotic arrest (mitotic slip). This mitotic slip is assumed to be the key mechanism for this resistance to drugs [11]. For aneuploid cells to survive, mitotic slippage is required. Changes in the genome composition of aneuploid cells may cause these cells to acquire drug resistance. Chromosomal instability (CIN) is common in solid cancers, mostly due to weakened or overactive mitotic checkpoints. The deteriorated function of the spindle examination point caused by downregulated BUBs expression may lead to mitotic slippage of BrCa cells and then cause acquired drug resistance [12, 13].

4. Discussion

SAC is a monitoring system that assists in the appropriate separation of chromosomes during mitosis. Because of less time during mitosis, the chromosomes do not join and separate in the normal manner, resulting in daughter cells with an abnormal number of chromosomes, a state called aneuploidy. Aneuploidy can result in failure to survive, developmental abnormalities, or the initiation and development of diseases such as cancer, depending on its sternness [14–16]. Some studies have shown that aneuploidy induced by SAC damage in the midgut of drosophila will lead to abnormal intestinal development or tumorgenesis [17]. The SAC mechanism is used by cells during mitosis to guarantee that all chromosomes have enough time to attach to spindle microtubules before the cell divides [5, 18]. SAC mainly consists of mitotic arrest defect (MAD) and BUB genes. As a critical regulator of mitosis, BUBs play a critical function in mitotic spindle inspection [19]. BUB1 phosphorylates BUB3 during mitosis, and the BUB1-BUB3 complex acts on APC/C inhibition when the SAC is activated and inhibits APC/C ubiquitin ligase activity by phosphorylating its activator CDC20. Additionally, this compound phosphorylates MAD1L1. In contrast, to its requirement for suppressing APC/C CDC20 and chromosome alignment, kinase activity in SAC activity is only a bit important [20]. Even though BUB1’s kinase activity is not essential for SAC, it appears that BUB1’s primary job is to recruit other SAC components such as MAD1, MAD2, CDC20, and BUB1 to the kinase complex. Additionally, it catalyses the production of the MAD2-CDC20 complex at the centromeres, which eventually forms the mitotic checkpoint complex (MCC), a powerful inhibitor of APC/C CDC20. It is essential for SAC signaling and correct chromosome alignment [3]. BUB1B, a paralogue of BUB1, is likewise necessary for normal mitotic progression. One of its checkpoint functions is to delay the late onset of disease by inhibiting APC/C and ensuring the correct separation of chromosomes. In addition, it is necessary to monitor the activity of kinetochore...
2.6 t–tests $p = 0.00076$

BUB1 expression

2.8

2.4

G1
($n = 3$)

G2
($n = 3$)

(a)

2.50

2.75

2.25

2.50

G1
($n = 3$)

G2
($n = 3$)

(b)

Figure 10: Continued.
motors that are dependent on the CENPE kinetochore motor. This protein is required for the localization of the CENPE kinetochore. CENPE interacts with BUB1B, resulting in the activation of its kinase activity, which is important for the activation of SAC. Kinetochore localization of BUB1B is not critical to checkpoint signaling, but it is important to regulate motile microtubule connections.

Additionally, BUB1B is important in initiating apoptosis in polyploid cells that retreat abnormally from mitotic stasis, a process that contributes to tumor suppression [2, 21]. Like BUB1, BUB3 locates at the centromere before chromosomal alignment, facilitating kinetochore localization of BUB1, thereby activating checkpoints in response to the unconnected kinetochore. It has dual functions in SAC signal...
transduction and in establishing proper kinetochore-microtubule attachments [22, 23].

There are reports of BUB gene imbalance in many cancers. For example, BUB1 can enhance the proliferation of hepatocellular carcinoma cells by activating the phosphorylation of SMAD2 [24], and BUB1B promotes liver cancer progression by activating the mechanistic target of rapamycin complex 1 (mTORC1) signaling pathway [25]. Overexpression of BUB1 accelerates the growth of prostatic cancer and predicts a negative prognosis in patients, according to the National Cancer Institute [26]. Overexpression of BUB1 and BUB1B in tumor tissues is related to a poor prognosis in pancreatic ductal adenocarcinoma, and it is also associated with advanced tumor stage and tumor development [27, 28]. Highly expressed BUB3 may weaken the SAC signal by changing the stoichiometric balance of SAC proteins required for MCC formation and checkpoint activity, which may lead to chromosomal missegregation, aneuploidy, and reduced residence time in mitosis [29]. It has also been reported that SAC is essential for the survival of BrCa cells. Inhibition of BUB1B leads to a decrease in the viability and clonogenicity of BrCa cell lines and significantly increases cell apoptosis and cell death. BUB1B gene knockdown can also cause acute chromosomal abnormalities. Knockout of BUB1B on the mouse MDA-MB-468 cell line can reduce tumor growth [2]. Although studies have reported the prognostic effect of BUBs on low-grade BrCa at the protein level [30], BUBs as a gene family interact and are inseparable in the carcinogenesis process. This study analyzed the expression of BUB1, BUB1B, and BUB3 in BrCa from various angles and further explained that the specific BUBs play a role in the incidence and progression of BrCa. We used an online database to compare the differential expression of BUBs in BrCa and normal breast tissues and found that the expression of BUBs in BrCa was higher than that in normal tissues. In addition, Kaplan-Meier Plotter was used to determine a significantly reduced RFS in BrCa patients who overexpressed BUBs, while high expression of BUB1 and BUB1B exhibited a strong correlation with the poorer OS. From the Cox regression analysis on a univariate approach, it is concluded that BUB1 and BUB1B can be used as biomarkers for the prognosis of BrCa. Further analysis of multivariate Cox regression shows that BUB1B and BUB3 may be independent prognostic indicators of BrCa. Immune cells in the body are constantly watching the cells around them, and when they spot alien cells, they activate their immune mechanisms to get rid of the “bad cells” and keep the body safe. It can be seen from the above studies that the high expression of B lymphocytes is beneficial to improving the prognosis of patients, but BUB3 negatively regulates B lymphocytes, making it easy for tumor cells to escape immune surveillance. BUB1 and BUB1B suppress the immune response by increasing the expression of immune checkpoint molecules, causing tumor cells to form immune escape. Studies have found that cells with faulty mitotic checkpoints are more resistant to a variety of anticancer treatments, including microtubule disruptors and DNA disruptors, than cells with normal checkpoints. Mitotic checkpoints are also related to the response to DNA damage. As a result, cancer cells with malfunctioning mitotic checkpoints become resistant to certain anticancer treatments that employ DNA damage as a mechanism of action [31]. We found MCF-7 and MCF-7/ADR in the GEO database and found that BUBs and their coexpression genes were downregulated in drug-resistant cell lines. Anti-microtubular drugs are commonly used as first-line agents for the treatment of a variety of malignancies. In one study, it was suggested that the basis of antiproliferative effects is the result of continuous activation of SAC, prolonged mitotic block, and mitotic cell death caused by microtubule dynamic disturbances. However, cells may also adopt the process of mitotic slippage in addition to mitotic cell death, thereby promoting cell survival and drug resistance [32]. In our study, BUBs are highly expressed in BrCa. Although it cannot be ruled out that the increase in gene expression may lead to instability of the genome, it is more likely to be cell compensation for other molecular component defects in the mitotic spindle damage checkpoint. Increased expression of these genes may be a BrCa marker for chromosomal instability [33, 34].

5. Conclusion

Through the above analysis, BUBs as biomarkers of BrCa may be potential treatment targets.

Abbreviations

BUB1: Budding uninhibited by benzimidazoles 1
BUB1B: BUB1 mitotic checkpoint serine/threonine kinase B
BUB3: Budding uninhibited by benzimidazoles 3
BUBs: BUB1, BUB1B, and BUB3
BrCa: Breast cancer
SAC: Spindle assembly checkpoint
APC/C: Anaphase-promoting complex or cyclosome
CDC20: Cell division cycle 20
CENPE: Centromere protein E
OS: Overall survival
RFS: Relapse-free survival
CI: Confidence interval
HR: Hazard ratios
PPI network: Protein-protein interaction network
GO: Genetic Ontology
KEGG: Kyoto Encyclopedia of Genes and Genomes
MF: Molecular function
CC: Cellular component
BP: Biological process
TNBC: Three negative breast cancer
PFS: Progression-free survival
TIME: Tumor-immune microenvironment
TIL: Tumor infiltration lymphocyte
ADR: Adriamycin
CIN: Chromosomal instability
MAD: Mitotic arrest defect
MCC: Mitotic checkpoint complex
mTORC1: Mechanistic target of rapamycin complex 1.

Data Availability

The datasets of bioinformatics used in this study are available from the corresponding author upon reasonable request.

Conflicts of Interest

The authors declared that no competing interests exist in this study.

Authors’ Contributions

XCW is responsible for conceptualization and methodology; SNW is responsible for software, data management, data analysis, and writing—original draft preparation; XYL is responsible for visualization, reviewing, and verification; MY is responsible for quality assessment; DQY, KY, and XQ are responsible for resources and supervision. All authors read and approved the final manuscript. Shunan Wang and Xinyu Liu contributed equally to this work.

Acknowledgments

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Supplementary Materials

Supplementary 1. Figure S1 The expression of BUBs in different T/N stages of BrCa. ns, \( p \geq 0.05; ^* p < 0.05; ^{*}{*} p < 0.001 \).

Supplementary 2. Figure S2 BUBs and their coexpressed genes protein interaction network.

Supplementary 3. Figure S3 Analysis of BUB gene family in GeneMANIA database.

Supplementary 4. Supplementary Table 1 Differential expression of BUBs and their coexpressed genes in the GSE155478 dataset.

References


