

Lysyl oxidase promotes anaplastic thyroid carcinoma cell proliferation and metastasis mediated via BMP1

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Background: Anaplastic thyroid carcinoma (ATC) is an extremely aggressive solid tumor with no effective treatment at present. Because of the rapid growth and aggressiveness, nearly all patients die within six months after developing ATC. Hence, more research regarding novel therapeutic targets for ATC is urgently needed.

Methods: Single-cell RNA sequencing data and microarray data of ATC were retrieved from the Gene Expression Omnibus (GEO) database. Cell clustering was performed using the Seurat package. Then, differential expression and functional enrichment analyses were performed. Gene set enrichment analysis (GSEA) was further used to investigate the functional enrichment of lysyl oxidase (LOX) and bone morphogenetic protein-1 (BMP1). The expression levels of LOX and BMP1 were measured using quantitative real-time PCR and Western blot. LOX and BMP1 were knocked down using si-RNAs. Cell proliferation was evaluated by the CCK-8 and clone formation assays. Cell migration and invasion were assessed by the wound healing assay and Transwell assay, respectively.

Results: LOX was upregulated at the single-cell level, as well as in ATC tissues and cell lines. LOX knockdown significantly inhibited ATC cell proliferation. Furthermore, the migration and invasion of ATC cells were remarkably inhibited after LOX inhibition. In addition, BMP1 regulated LOX expression in 8505C cells, while BMP1 overexpression restored the LOX activity blocked by the LOX inhibitor BAPN. BMP1 could also induce the cell proliferation and metastasis of ATC.

Conclusions: LOX/BMP1 mediates the malignant progression of ATC, highlighting the potential application of LOX/BMP1 in the treatment of ATC. This study provides new insights for efficient therapeutic agents based on the LOX/BMP1 axis.

Keywords: Lysyl oxidase (LOX); bone morphogenetic protein-1 (BMP1); anaplastic thyroid carcinoma (ATC)

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Introduction

Thyroid carcinomas are one of the most common endocrine cancers, including papillary thyroid carcinoma (PTC), follicular carcinoma, anaplastic thyroid carcinoma (ATC), and medullary carcinoma (1). Although most thyroid cancers are localized and are usually associated with good survival, population-based studies have documented a significant increase in the incidence of thyroid cancer worldwide across all pathological types over the past decades (2). ATC was arise from the follicular cells of the thyroid gland and compared to other thyroid cancers, ATC cells do not retain any of the biological features of normal follicular cells, such as thyroglobulin synthesis, iodine uptake, and thyroid stimulating hormone dependence (3). Characterized by aggressive local disease, rapid growth, high rates of metastasis, and fatal clinical outcomes, the median survival of patients with ATC was less than three to six months (4). Therefore, more studies regarding valid and practical therapeutic targets for ATC are urgently needed.

Previous studies have revealed several biological factors that may have predictive or prognostic relevance for ATC by means of immunohistochemistry and bioinformatics. The lysyl oxidase (LOX) family is included among the identified biomarkers. LOX is an enzyme that catalyzes the cross-linking of elastin and collagen in the extracellular matrix, thus controlling the tensile strength of tissues (5,6). Furthermore, LOX family proteins have been shown in multiple studies to drive the processes of carcinoma progression, including gene expression regulation, cell growth, invasion, and metastasis (7,8). LOX has thus been suggested as a promising strategy to prevent metastasis and invasion of ATC. Whether LOX is involved in ATC progression is yet to be established, and the specificity and effects of LOX expression in malignant and non-malignant cells in ATC are unclear. This current study aimed to investigate the functional role of intracellular LOX in ATC.

LOX is a forty-eight kd protein and N-glycosylation and signal peptide cleavage are essential for LOX protein activation. After that, the 50 kd proenzyme is secreted and converted into a mature, active 30 kd form. In addition, the catalytic activity of LOX can be specifically and irreversibly inhibited by beta-aminopropionitrile (BAPN) (9). Procollagen C proteinase/bone morphogenetic protein-1 (BMP1) was originally identified as a secreted metalloprotease of the astacin metalloproteinase family, which plays a role in LOX proteolytic processing (10,11). By inducing the initial cleavage and release of some complexes, such as TGF- β , BMP1 has been regarded to be involved in tumor progression (12). For example, BMP1 overexpression reflects poor prognosis in clear cell renal cell carcinoma (13), whereas the role of BMP1 in thyroid carcinomas, especially in ATC, has never been reported.

In the present study, we accessed both single-cell sequencing data and microarray data from public databases and analyzed them by several computational biology methods. We also examined the expression of LOX in ATC tissues, PTC tissues, and cell lines. Then, we investigated the regulatory effect of LOX and BMP1 on the proliferation, migration, and invasion of ATC cells. Further validation of the interaction between LOX/BMP1 was conducted to determine whether LOX/BMP1 could be identified as a novel target of ATC.

We present the following article in accordance with the MDAR reporting checklist (available at https:// gs.amegroups.com/article/view/10.21037/gs-21-908/rc).

Methods

Data sources

Single-cell sequencing ATC data (GSE148673, number of samples: 5) and normal data (GSE134355, number of samples: 2) were acquired from the Gene Expression Omnibus (GEO) database (www.ncbi.nlm.nih.gov/ geo). Four microarray datasets (GSE29265, GSE33630, GSE65144, and GSE76039) associated with ATC were also extracted from the GEO database. Information about cell types was acquired from annotations of the downloaded data.

Data processing

R software (version: 4.0.4) was used for all the analyses in the study.

Data quality control and data merging

The single-cell sequencing data was read and filtered with Scrublet (14). The cells were screened out according to 200< nFeature_RNA <8,000, nCount_RNA >10, and mitochondria-expressed genes <20%. The batch effect from microarray data was removed by the "limma" package (function: removeBatchEffect()) while preserving differences between groups.

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Table 1 Clinical and pathological characteristics of patients

Patient No.	Histology	Age (years)	Sex
1	PTC	65	Male
2	PTC	68	Female
3	PTC	58	Male
4	PTC	54	Male
5	PTC	54	Male
6	PTC	67	Female
7	PTC	43	Female
8	PTC	42	Male
9	PTC	58	Male
10	PTC	58	Female
11	PTC	58	Male
12	PTC	59	Female
13	PTC	22	Female
14	PTC	66	Male
15	PTC	21	Female
16	PTC	25	Female
17	PTC	47	Female
18	PTC	28	Female
19	PTC	52	Male
20	PTC	45	Female
21	PTC	62	Female
22	PTC	56	Female
23	PTC	50	Female
24	PTC	54	Female
25	PTC	48	Male
26	PTC	48	Male
27	PTC	55	Male
28	PTC	53	Male
29	ATC	75	Male
30	ATC	54	Male
31	ATC	47	Male
32	ATC	72	Female
33	ATC	77	Male

PTC, papillary thyroid cancer; ATC, anaplastic thyroid cancer.

Single-cell copy number variation analysis and cell type annotation

We applied CopyKAT to analyze copy number variations of the single-cell sequencing data (15). Aneuploid copy number profiles which have diploid genomes are common (88%) in tumors but are not found in normal cell types, by means of identifying of aneuploid copy number profiles, It is effective to separate normal cells from tumor cells (16). Aneuploidy cells were defined as malignant ATC cells. In regard of the batch effects in different samples, R package Harmony was used to integrate these samples to eliminate the batch effects. After standardizing and scaling the data, principal component analysis (PCA) was conducted for the purpose of dimensionality reduction (17). UMAP was applied to cluster and visualize data which was then identified and annotated with the SingleR package.

LOX expression analyses

The expression of LOX in single-cell sequencing samples was obtained and visualized by the Seurat package. We defined LOX-positive groups as LOX >0 while the others were negative groups. We screened the top 20 LOX-related marker genes from LOX-positive groups by performing differential gene expression analysis. The expression of these LOX-related marker genes in single-cell sequencing and microarray data were visualized by a heatmap.

Gene set enrichment analysis (GSEA)

GSEA (http://software.broadinstitute.org/gsea/index.jsp) was used to assess LOX- or BMP1-related pathways and molecular mechanisms in ATC samples (18). Enriched gene sets with P values of <0.05 were considered statistically significant.

Expression analyses based on ATC, PTC, and normal tissues

Samples of ATC, PTC, and normal tissues were obtained from ATC patients and PTC patients who were performed surgery in Zhejiang Cancer Hospital, Clinical and pathological characteristics of the patients were shown in *Table 1*. This study was approved by the Ethics Committee of Zhejiang Cancer Hospital (No. 2020QT259) and performed according to the Declaration of Helsinki (as

 Table 2 The primers of the genes

Gene symbol	Forward/reverse primer		
LOX	Forward: 5'-TCCAGGACTGAGAAAGGGGAA-3'		
	Reverse: 5'-GGGTGGGATTCAGACCCTTC-3'		
BMP1	Forward: 5'-CACACCTGTGTCACCTTCCT-3'		
	Reverse: 5'-CCGAGCCGCAAGGTC-3'		
GAPDH	Forward: 5'-GTCATCCATGACAACTTTGG-3'		
	Reverse: 5'-GAGCTTGACAAAGTGGTCGT-3'		

revised in 2013) and Good Clinical Practice guidelines. Patients provided written informed consent for their tissue to be used in this study. Total RNA was extracted using TRIzol[®] reagent (Cat No.15596-018, Life Technologies, Carlsbad, CA, USA) following the manufacturer's instructions and checked to inspect RNA integrity. Qualified total RNA was further purified by the RNeasy mini kit (Cat No.74106, QIAGEN, GmBH, Hilden, Germany) and labeled. Each slide was hybridized with 600 ng Cy3labeled cRNA using the Gene Expression Hybridization Kit (Cat No. 5188-5242, Agilent Technologies, Santa Clara, CA, USA), then slides were scanned with default settings. The original data from the gene microarray was adjusted using the limma package (19) and screened for differentially expressed genes (DEGs) using 2^{Ilog2(fold change)I} >1 with P<0.05.

Cell culture

The PTC cell line BCPAP was obtained from Professor Haixia Guan (Department of Endocrinology and Metabolism, The First Affiliated Hospital of China Medical University, Shenyang, China), and the ATC cell line 8505C was purchased from the Shanghai Cell Bank of the Chinese Academy of Sciences (Shanghai, China). The thyroid epithelial cell line Nthy-ori3-1 was bought from the Shanghai Institute of Cell Biology, Chinese Academy of Sciences (Shanghai, China). Nthy-ori3-1 cells and BCPAP cells were cultured in RPMI 1640 and 8505C cells were cultured in MEM. All the medium was added with streptomycin (100 U/mL), penicillin (100 U/mL) and 10% fetal bovine serum (FBS). Cultures were incubated at 37 °C in a humidified atmosphere of 5% CO₂.

Cell transfection

Cells were plated into 6-well plates one day before

transfection. Transfection was performed with si-LOX, si-BMP1, and the corresponding scramble control (si-NC) (Applied Biological Materials Inc., Vancouver, BC, CANADA) using Lipofectamine[®] 3000 (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA), according to the manufacturer's protocol. After transfection for 6 h, the culture medium was removed and replaced with fresh DMEM containing 10% FBS.

Total RNA extraction and real-time PCR quantification

Total RNA was isolated using the TRIzol[®] reagent (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA), and quantified by absorbance at OD 260 nm. The purity of the RNA was assessed by measuring the ratio of OD 260 nm/OD 280 nm. The total RNA was reversetranscribed into cDNA using the PrimeScript RT Reagent kit (Takara Biotechnology Co., Ltd., Dalian, China). *Table 2* is the primers for related genes. All the PCR reactions were carried out using SYBRs Premix Dimer-Eraser[™] (Takara Biotechnology Co., Ltd., Dalian, China). The real-time PCR assays were conducted by LightCycler 480 (Roche Diagnostics) and the mRNA expression was normalized by *GAPDH*.

Western blot assay

Cells were washed with pre-cooled phosphate-buffered saline (PBS) on ice and then boiled in SDS sample buffer. Proteins samples were electrophoresed on polyacrylamide gel electrophoresis gels to resolve, and then transferred to polyvinylidene fluoride membranes. After blocking with skim milk, the membranes were incubated with primary antibodies against the following proteins: LOX, BMP1, and β -actin (Proteintech, Rosemont, IL, USA). Membranes were then blotted with horseradish peroxidase (HRP)-conjugated secondary antibodies (Jackson Immuno Research Laboratories, West Grove, PA, USA). The images of the gels were scanned using the Bio-Rad Gel Doc XR+ system (Bio-Rad, Hercules, CA, USA).

Co-immunoprecipitation (co-IP) assays

PureProteome Protein G Magnetic Beads (Millipore Corp., Billerica, MA) were washed 3 times with PBS contain 0.01% Tween 20 and centrifuged at 2,500 ×g for 5 minutes. Then the beads were suspended in PBS with 2 µg of the specific antibodies for 2 hours at 4 °C. After incubation,

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the beads were centrifuged at 2,500 ×g and removed the supernatant. After washing them with PBS, the beads were then resuspended in protein extract in immunoprecipitation buffer and PI for 500 μ L total volume, the samples were rotated overnight at 4 °C. The beads were recovered by centrifugation at 2,500 ×g for 5 minutes and washed 3 times in PBS before solubilizing in 20 μ L RIPA buffer. Loading dye (5 μ L of a 5× solution) was added and the beads were resuspended prior to preparation for SDS-PAGE.

Cell counting kit-8 (CCK-8) assay

Cells were seeded into 96-well plates at 5×10^3 cells/well. The CCK-8 assay was performed to detect cell proliferation ability at 72 h following inoculation. In brief, 10 µL of CCK-8 dye (Beyotime Institute of Biotechnology, Shanghai, China) was added into each well and cells were incubated for 1 h at 37 °C under 5% CO₂. Finally, we used microplate reader (Bio-Rad Laboratories, Hercules, CA, USA) to obtain the absorbance of each well at 450 nm.

Clone formation assay

From each group, approximately 1×10^3 cells were plated in each well of a 6-well culture plate. Cells were incubated at 37 °C for 2 weeks with culture media being replaced every 3 days. Then, the cells were washed 3 times with PBS and stained with 0.5% crystal violet. Photomicrographs were taken at a magnification of 10× for all groups.

Transwell assay

The cells invasive ability was determined by the Transwell assay. Cells were cultured in 100 μ L serum-free medium after transfection for 48 hours. Cells were plated into the upper chamber containing a Transwell pre-coated Matrigel membrane filter and 8 μ m pore inserted in the culture plates. The cells remaining on the upper membrane were removed with cotton swabs after 48 h incubation. As for the cells which had invaded through the membrane, they were fixed in 4% paraformaldehyde and stained with 0.1% crystal violet for 20 min at 4 °C. Counts were finally obtained from 5 random fields at 10× magnification.

Scratch wound healing assay

The migratory ability of cells was analyzed using a wound healing assay. Briefly, cells were plated on 6-well plates and scraped by a pipette tip to generate uniform wounds. The initial gap length at 0 h and the residual gap length at 24 h after wounding were observed and photographed at a magnification of $10\times$ for each group.

LOX activity assay

ATC cells conditioned media (50 µL) was added to 150 µL of reaction mixture containing 10.6 mM cadaverine (Sigma-Aldrich), 1 M urea (Sigma-Aldrich), 0.2 U/mL HRP (Sigma-Aldrich), and 20 µM amplex red (Molecular Probes, Invitrogen) in the 96-well clear-bottom black plate (Perkin Elmer, Waltham, MA). Plates were incubated at 37 °C for 1 h. Finally, microplate reader (Bio-Rad Laboratories, Hercules, CA, USA) was used to read the absorbance of each well at 544 nm excitation and 590 nm emission.

Correlation analysis

The matrix of gene expression data [GSE29265 (n=9)] associated with ATC was extracted from the GEO database (http://www.ncbi.nlm.nih.gov/geo/; species: homo). The Pearson method was used to determine the correlation coefficient. The R4.0.1 Ggstatsplot package was used to visualize the correlation analysis results.

Immunofluorescence

Cells were plated onto coverslips and fixed in cold methanol for 10 minutes. Then the cells were blocked with 5% FBS and incubated with the primary antibodies and secondary antibodies. The sections were counterstained with 4,6-diamino-2-phenylindole and observed under a fluorescence microscope. Photomicrographs were taken for all groups.

Statistical analysis

All the experiments were performed 3 times independently,



Figure 1 LOX expression in ATC cells delineated by single-cell sequencing analysis. (A) tSNE of single-cell profiling with each cell colorcoded for sample type and associated cell type (up). Expression of LOX for the cell types defined above each panel (down). (B) Violin plots show the expression levels of LOX in malignant ATC cells and normal cells. (C) Heatmap representing the upregulated and downregulated top 20 LOX-related genes in aneuploid and diploid clusters. t-SNE, t-distributed stochastic neighbour embedding; LOX, Lysyl oxidase; ATC, anaplastic thyroid carcinoma.

and the experiments data were analyzed by means of GraphPad Prism software (La Jolla, CA, USA). Results are presented as mean \pm SD. Differences between groups were tested using paired and paired Student's *t*-tests. P<0.05 was considered statistically significant.

Results

Single-cell profiling and clustering of ATC cells

First, single-cell sequencing analysis was performed to analyze LOX expression. ATC and normal cells were both grouped into 11 clusters. LOX expression in malignant ATC cells was significantly higher than that of normal cells (P<0.05) (*Figure 1A,1B*). By leveraging single-cell melanoma expression data, we identified gene profiles that were significantly co-expressed with LOX in different cell types, and the heatmap shows the top 20 marker genes in aneuploid and diploid subsets (*Figure 1C*). Overall, LOX showed abnormally high expression at the single-cell level.

LOX plays an important role in ATC and is associated with proliferation and migration

We used microarray data from the GEO dataset to verify the expression profiles of the above-mentioned LOXrelated genes in *Figure 1*, and 14 of the top 20 genes were overexpressed in ATC (*Figure 2A*). Consistent with singlecell sequencing analysis, LOX expression in the ATC group (n=52) was markedly higher than in the normal group (n=78) from microarray analysis (median 7.48 *vs.* 4.20, P<0.05) (*Figure 2B*), which proved that LOX are malignant gene expressed in ATC. GSEA showed that the high LOX expression group was mainly enriched in Gene Ontology (GO) biological processes related to cell population proliferation and cell migration (*Figure 2C*).

LOX is significantly upregulated in ATC tissues

We first examined the expression of LOX in ATC, PTC, and normal tissues, and a significantly higher LOX level was observed in ATC tissues (*Figure 3A*, *3B*) (P<0.01). Subsequently, total RNA was extracted from the thyroid epithelial cell line Nthy-ori3-1, the PTC cell line BCPAP, and the ATC cell line 8505C. RT-qPCR analysis revealed that *LOX* expression was significantly increased in 8505C cells compared to BCPAP and Nthy-ori3-1 cells (*Figure 3C*, P<0.05). These results indicated that LOX was upregulated in ATC.

Knockdown of LOX suppresses the proliferation, migration, and invasion of ATC cells

LOX was significantly knocked down in 8505C cells after transfection with si-RNA (*Figure 4A*, P<0.01). LOX activity was also decreased in si-LOX 8505C cells



Figure 2 The expression of LOX was upregulated in ATC tissues and cell lines. (A) Heatmaps representing the upregulated and downregulated top 20 LOX-related genes in ATC and normal subsets using microarray analysis. (B) The expression level of LOX in the ATC and normal groups (7.48 *vs.* 4.20). (C) GSEA showing high-LOX groups are associated with cell proliferation- and migration-related GO pathways. LOX, Lysyl oxidase; ATC, anaplastic thyroid carcinoma; GSEA, gene set enrichment analysis; GO, Gene Ontology.

(*Figure 4B*, P<0.001). Both growth curves and colony formation efficiency showed that LOX downregulation impeded the proliferation of 8505C cells compared with cells transfected with control (*Figure 4C*,4D, P<0.01). In addition, the migratory and invasive abilities of 8505C cells were examined using the Transwell migration and scratch wound healing assays. The numbers of invaded (*Figure 4E*) and migrated (*Figure 4F*) 8505C cells were much lower in the in si-LOX group than in the control group. Similar results were observed in 8505C cells treated with the LOX inhibitor BAPN (*Figure 4G*,4H). All these results illustrated the effects of LOX in the malignant progression of ATC.

The effect of BMP1 in the proliferation and metastasis of ATC cells

A previous study revealed that LOX can be activated by

BMP1 in other tumors. In this study, BMP1 expression in the ATC group was markedly higher than in the normal group as determined by microarray analysis (median 6.02 vs. 5.39, P<0.05) (Figure 5A). We conducted GSEA to determine the molecular function of BMP1 in ATC, and the high-BMP1 group was also enriched in cell proliferationand migration-related GO pathways (Figure 5B). To confirm whether BMP1 plays a role in promoting the proliferation and metastasis of 8505C cells, we knocked down BMP1 by si-RNA. The CCK-8 and clone formation assays both demonstrated that BMP1 knockdown could reduce ATC cell viability (*Figure* 5C, 5D). The migratory and invasive abilities of 8505C cells were also impaired by BMP1 knockdown (Figure 5E, 5F). Moreover, si-BMP1 could enhance the inhibition effect of BAPN. Taken together, BMP1 protein exerted a promotive effect on ATC malignant progression, and therefore may be regarded as a



Figure 3 The expression of LOX was upregulated in ATC tissues and cell lines. (A) The protein expression level of *LOX* in ATC, PTC, and normal tissues was measured using protein mass spectrometry analysis. (B) The mRNA level of LOX in ATC, PTC, and normal tissues was determined by quantitative real-time PCR. (C) The mRNA level of *LOX* in Nthy-ori3-1, BCPAP, and 8505C cells was detected using quantitative real-time PCR. *, P<0.05. LOX, Lysyl oxidase; ATC, anaplastic thyroid carcinoma; PTC, papillary thyroid carcinoma; mRNA, messenger RNA.



Figure 4 LOX knockdown inhibited the proliferation, migration, and invasion of ATC cells. (A) The expression of LOX was knocked down using si-RNAs in 8505C cells. (B) The activity of LOX was downregulated after knocking down LOX in 8505C cells. (C) The proliferation of 8505C cells after LOX knockdown was assessed using the cell counting kit-8 assay. (D) The proliferative ability of 8505C cells after LOX silencing was evaluated using the clone formation assay (crystal violet staining at 10× magnification). (E,G) Cell migration of 8505C cells was evaluated by the wound healing assay (at 10× magnification). (F,H) Cell invasion of 8505C cells was evaluated by the Transwell assay (crystal violet staining at 10× magnification). **, P<0.01; ***, P<0.001. LOX, Lysyl oxidase; si-RNAs, short interfering RNAs.

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Figure 5 The effect of BMP1 on the proliferation and metastasis of ATC cells. (A) The expression level of BMP1 in the ATC and normal groups (6.02 *vs.* 5.39). (B) GSEA showing that the high-BMP1 group was associated with cell proliferation- and migration-related GO pathways. (C,D) The proliferative ability of 8505C cells was evaluated by the cell counting kit-8 assay (C) and clone formation assay (D) (crystal violet staining at 10× magnification). (E) Cell invasion of 8505C cells was evaluated by the Transwell assay (crystal violet staining at 10× magnification). (F) Cell migration of 8505C cells was evaluated by the wound healing assay (at 10× magnification). ***, P<0.001. BMP-1, bone morphogenetic protein-1; ATC, anaplastic thyroid carcinoma; GSEA, gene set enrichment analysis; GO, Gene Ontology; PTC, papillary thyroid carcinoma.

new therapeutic target of ATC.

Association between LOX and BMP1 in ATC cells

The correlation between BMP1 and LOX, as determined by the Person method based on the GEO database (*Figure 6A*), was R=0.52, which indicates a significant positive correlation. In order to investigate the association between LOX and BMP1 in ATC cells, we conducted an immunofluorescence assay to analyze the co-localization of BMP1 and LOX proteins in 8505C cells (*Figure 6B*). Co-IP revealed that BMP1 might bind with LOX in 8505C cells (*Figure 6C*).

BMP1 could mediate the activity and expression of LOX in ATC cells

In order to further investigate the effect of BMP1 on the function of LOX in ATC, we knocked down BMP1 by si-RNA (*Figure 7A*). Compared with the control group, RT-PCR and Western blot assays revealed that LOX was significantly downregulated in si-BMP1 8505C cells (*Figure 7B*, P<0.01). Furthermore, the activity of LOX was also decreased when BMP1 was knocked down (*Figure 7C*, P<0.05). When BMP1 was overexpressed, the activity of LOX inhibited by BAPN could be rescued, as shown in *Figure 7D*. These results showed that BMP1 could mediate



Figure 6 Association between LOX and BMP1 in ATC cells. (A) The correlation between BMP1 and LOX using the Pearson method based on the GEO database. (B) The co-localization of BMP1 and LOX proteins in 8505C cells was shown by immunofluorescence (at 10x magnification). (C) The interaction between BMP1 and LOX was detected using co-IP. BMP-1, bone morphogenetic protein-1; LOX, Lysyl oxidase; ATC, anaplastic thyroid carcinoma; co-IP, co-immunoprecipitation.



Figure 7 BMP1 could mediate the activity and expression of LOX in ATC cells. (A) The expression of BMP1 was knocked down using si-RNAs in 8505C cells. (B) The expression level of LOX in 8505C cells after si-BMP1 transfection was measured by Western blot and quantitative real-time PCR. (C) The activity of LOX was downregulated after knocking down BMP1 in 8505C cells. (D) The activity of LOX was determined after BMP1 overexpression and treatment with BAPN in 8505C cells. *, P<0.05; **, P<0.01. BMP-1, bone morphogenetic protein-1; si-RNAs, short interfering RNAs; LOX, Lysyl oxidase; BAPN, beta-aminopropionitrile (LOX inhibitor).

the activity and expression of LOX in ATC cells.

Discussion

In this study, we found abnormal expression of LOX and

BMP1 in ATC from the single-cell and gene expression level up to cell lines or tumor tissue level.

LOX is a secreted amine oxidase that causes stiffening of the matrix and plays a crucial role in mediating tumor microenvironment, which enhances the invasive and metastatic properties of the tumor (20). Previous study showed that LOX was significantly associated with breast cancer bone metastasis. High expression of LOX in primary breast tumors leads to osteolytic lesion formation (5). Therefore, LOX can prompt more aggressive tumor behavior.

Using the retrieved single-cell sequencing data, the molecular signature of ATC tumor tissues was analyzed. In tumor cells, a high expression of LOX was found at the single-cell level, and a subset of LOX-related genes was identified. In support of this, we examined LOX expression in thyroid tumor tissues. LOX expression was undetectable in normal tissue, thyroid epithelial cell line, and PTC cell line, whereas LOX had high abnormal expression in ATC tissues samples and the ATC cell line (8505C). Furthermore, functional analysis demonstrated that knocking down LOX with siRNA or inhibiting LOX with BAPN led to a significant reduction in the migration and invasion of 8505C cells. BAPN was obtained originally from sweet peas, Lathyrus odoratus L and was the first chemical compound described as LOX potent irreversible inhibitor. BAPN has shown anticancer properties in several different cancer types. Such findings are in agreement with a previous study that LOX exacerbated the malignant and invasive properties of cancer.

BMP1 was reported that was responsible for processing LOX into the active form (21), and an association between LOX activation and BMP1 has been previously reported in other cancers (22). Similar results were confirmed in this study by knocking down BMP1 in ATC. To further understand the interaction between BMP1 and LOX expression in ATC, we used co-IP and immunofluorescence co-localization analysis. Our results identified relevant protein-protein interactions of LOX and BMP1. Importantly, GSEA was performed and demonstrated that BMP1 was related to cell proliferation and migration. For validation, the in vitro assays demonstrated that si-BMP1 could inhibit ATC cell proliferation and invasion. Moreover, BMP1 overexpression restored the activation of LOX repressed by BAPN in 8505C cells. Although there are no current agents that are known to target LOX/BMP1, our results suggest that LOX/BMP1 depletion or LOX inhibition could be an effective treatment strategy in ATC.

Functional enrichment analysis of the LOX-related genes from ATC cells may help to define the specific roles of LOX expression in malignant tumor development and progression. However, we do not have further information on the involvement of LOX in influencing PTC cell proliferation and migration through these genes. In future work, we will further explore the relationship between them.

However, our study had limitations. Our study provides evidence that LOX and BMP1 promote the proliferation and migration of ATC cells. But the analysis was only conducted *in vitro* study, further *in vivo* experiments of ATC need to be explored in future studies.

Overall, our results confirmed that LOX plays an important role in ATC malignant progression. BMP1 can activate LOX and promote the proliferation and migration of ATC cells. The inhibition of LOX and BMP1 was suggested as a promising therapeutic strategy for oncological diseases, including ATC. In the upcoming years, we expect to obtain more detailed information on the research of LOX and BMP1. The combination of LOX enzyme inhibitors with standard anticancer treatments is another approach that should be further explored in future studies. Our data will be contributed to the progress of therapeutic development against ATC.

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Footnote

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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. This study was approved by the Ethics Committee of Zhejiang Cancer Hospital (No. 2020QT259), and performed according to the Declaration of Helsinki (as revised in 2013) and Good Clinical Practice guidelines. Patients provided written

informed consent for their tissue to be used in this study.

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