LIM protein JUB promotes epithelial–mesenchymal transition in colorectal cancer

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Metastasis is the leading cause of cancer-related death in almost all types of cancers, including colorectal cancer (CRC). Metastasis is a complex, multistep, dynamic biological event, and epithelial–mesenchymal transition (EMT) is a critical process during the cascade. Ajuba family proteins are LIM domain-containing proteins and are reported to be transcription repressors regulating different kinds of physiological processes. However, the expression and pathological roles of Ajuba family proteins in tumors, especial in tumor metastasis, remain poorly studied. Here, we found that JUB, but not the other Ajuba family proteins, was highly upregulated in clinical specimens and CRC cell lines. Ectopic expression of JUB induced EMT and enhanced motility and invasiveness in CRC, and vice versa. Mechanistic study revealed that JUB induces EMT via Snail and JUB is also required for Snail-induced EMT. The expression of JUB shows an inverse correlation with E-cadherin expression in clinical specimens. Taken together, these findings revealed that the LIM protein JUB serves as a tumor-promoting gene in CRC by promoting EMT, a critical process of metastasis. Thus, the LIM protein JUB may provide a novel target for therapy of metastatic CRC.

Metastasis accounts for approximately 90% of cancer-associated deaths.1–3 An estimated 1.2 million cases of colorectal cancer (CRC) were recorded worldwide in 2008. Approximately 35% of patients present with stage-IV metastatic disease when diagnosed with primary disease, and 20–50% of subjects with stage-II or -III disease will progress to stage-IV disease.4,5 Mortality has declined by approximately 1.8% per year for stage-IV disease6,7 but the prognosis of metastatic CRC remains extremely poor (overall survival at 5 years <10%).8,9 Metastasis remains the most poorly understood component of the pathogenesis of cancer.

Epithelial–mesenchymal transition (EMT) is known to be fundamental for embryonic development, wound healing and fibrosis, but also for tumor invasion and metastasis.5,6 EMT is the initial process of tumor metastasis. Cells undergoing EMT usually acquire a mesenchymal phenotype, spindle-like morphology, high motility and invasiveness. Downregulation of E-cadherin is a hallmark of EMT. Control of transcription of the E-cadherin gene is the main mechanism accounting for downregulation of this protein.10 Several transcription factors have been reported to repress expression of E-cadherin: Snail, Slug, Twist and SIP,8,11 among which Snail was the first to be discovered. Snail has been reported to be a major transcription repressor of E-cadherin, frequently upregulated in breast cancer,8,11 esophageal squamous cell carcinoma12 and CRC.13

The LIM protein Ajuba family contains three members, JUB, WTIP and LIMD1, and is characterized by a unique N-terminal region, the preLIM region, and three tandem C-terminal LIM domains.14–16 This family of proteins function as scaffolds and have been reported to modulate many events in cells, such as cell proliferation and tissue size,17,18 DNA damage response,19 meiotic maturation of oocytes14 and...
embryonal cell proliferation and differentiation.(20) However, the pathologic roles of Ajuba family proteins in diseases remain unexplored. Langer et al.(21) find that LIM protein Ajuba family proteins interact with the SNAG domain of the Snail family. They use Xenopus neural crest as a model of in vivo Snail-induced EMT and demonstrate that Ajuba LIM proteins contribute to neural crest development as Snail/Slug corepressors and are required for in vivo Snail/Slug function.(21) Thus, we propose that Ajuba family proteins may also regulate EMT in cancer and promote metastasis.

In the present study, surprisingly, we find that the LIM protein JUB, but not the other two members of the Ajuba family, WTIP and LIMD1, was highly upregulated in CRC specimens and cell lines. Ectopic overexpression of JUB induces EMT and promotes migration and invasion in CRC cells. Silencing of JUB impairs EMT and inhibits migration and invasion in CRC cells. Further mechanistic study revealed that JUB promoting EMT depends on Snail, and JUB is also required for Snail to induce EMT, which is consistent with previous reports that JUB serves as a corepressor of Snail. The present study uncovered an important role of the LIM protein JUB in the pathological progress of tumorigenesis in CRC. Hereafter, JUB could be a novel therapy target for metastatic CRC.

Materials and Methods

Plasmids and antibodies. For overexpression of JUB, human JUB (538aa) was amplified by PCR from cDNA of SW620 cells and subcloned into a pSin-EF2-puro retrovector (Addgene). For depletion of JUB, two human shRNA sequences were cloned into the pSuper-retro-puro vector to generate pSuper-retro-JUB-RNAi(s). The target sequences were: RNAi#1, GGACCGGGATTATCACTTT, and RNAi#2, CCAAGTATACTGTGTCACC, as previously reported.(22) Human Snail gene was amplified from cDNA and inserted into the EcoRI and XhoI sites of the pCDNA3.1 vector. For silencing of Snail, oligonucleotides were purchased from RiboBio (RiboBio, Guangzhou, Guangdong) and the target sequence is 5’-gctgcagaccttctggcga-3’.

Antibodies, including anti-JUB (Cell Signaling Technology, Danvers, MA, USA), anti-WTIP (Sigma Aldrich, Santa Cruz, CA, USA), anti-LIMD1 (Sigma, Saint Louis, MO, USA), anti-β-actin (Sigma Aldrich), anti-E-cadherin (BD Biosciences, Bedford, MA, USA), anti-Vimentin (BD Biosciences) and anti-Snail (Cell Signaling Technology) were used for western blot analysis.

Cell culture and stable cell line establishment. Colo rectal cancer cell lines, including SW480, SW620, KM12, HCT15, HCT116, Caco-2 and LoVo, were purchased from ATCC (Manassas, VA, USA) and maintained in RPMI-1640 medium (Invitrogen, Carlsbad, CA, USA) supplemented with 10% FBS (HyClone, Logan, Utah, USA), 1% penicillin/streptomycin and non-essential amino acids (HyClone) at 37° in 5% humidified CO2 atmosphere. SW620 and SW480 cells were selected for stable overexpression or knockdown of JUB.

For stable cell line establishment, the indicated plasmids (pSin-EF2-puro-retro-JUB and pSuper-puro-retro-JUB-RNAi(s)) were packed into retrovirus in 293T cells. Then, the viruses were harvested and SW620 and SW480 cells were infected. After infection for 48 h, the cells were selected with medium containing puromycin (0.5 ug/mL) over 1 week.

RNA extraction, RT-PCR and real-time RT-PCR. Total RNA from cultured cells was extracted using Trizol reagent (Invitrogen) following the manufacturer’s instructions. The cDNA was amplified and quantified using an ABI Prism 7500 Sequence Detection System (Applied Biosystems, Lab India, Haryana, India), with SYBR Green I dye (Molec-well Probes, Invitrogen). β-actin was used as an internal control. Expression data were normalized to β-actin, and calculated as 2(-[(Ct of gene) – (Ct of beta-actin)]), where Ct represents the threshold cycle for each transcript. The primers for qRT-PCR were:

| Gene    | Forward Primer            | Reverse Primer            |
|---------|---------------------------|---------------------------|
| JUB     | AGAGGGCCAGGGAGGACTACT     | GAGGACAGAACAAACAGCTG     |
|         | TGGTGGCTTGGCATACG         | TGGCTGAAACCGGAGTACAG     |
|         | TCACCGGAGGCTGATTACT       | AGGGTAAAGCATGTCATGG      |
|         | GCACAGAGGCCCCTGGCCTT      | CTTTGCACATGCCCAGG        |

Clinical tissue specimens and ethics statement. Tissue specimens were freshly collected from Zengcheng People’s Hospital (Boji-Affiliated Hospital of Sun Yat-Sen University), so that patients could be histopathologically and clinically diagnosed. All samples were obtained with prior written informed consents from the patients and approval from the Institutional Research Ethics Committees of Zengcheng People’s Hospital (Boji-Affiliated Hospital of Sun Yat-Sen University) ethics Committee.

Transwell assays. A total of 4 × 10^4 of each the indicated cells were suspended in 200 µL basic 1640 medium and seeded into the upper trans-well cell culture chambers (BD Biosciences) coated with Matrigel (for migration assay) or without Matrigel (for migration assay). The lower chamber was loaded with 500 µL of 1640 medium with 10% FBS. After incubation for 48 h at 37°C in 5% CO2, cells were fixed with methanol and stained with 1% crystal violate. Cells presented on the lower surface of the membrane were quantified and photographed under a microscope. The average number of five randomly selected microscopic fields was presented.

3-D spheroid invasion assay. First, we coated the 24-well plates with 100% Matrigel (BD Biosciences) for the bottom layer. We incubated the coated plates at 37° for at least 30 min and then seeded 1 × 10^4 of the indicated cells mixed with 10% Matrigel on the top layer. The medium was refreshed every other day. Cells forming a 3-D spherical structure (spheres) were photographed at 2-day intervals for 2 weeks. Spheroids with outgrowth were considered to be invasive spheroids and the quantification of invasive spheroids was carried out as follows: high contrast images of spheroids were captured using AxioVision Rel.4.6 software (Carl Zeiss, Oberkochen, Germany) and 10 randomly selected fields of each well were analysed. The final data was presented as the percentage of invasive spheroids.

Dual-luciferase reporter assay. A total of 4 × 10^4 cells were seeded in triplicate in 24-well plates. Twenty-four hours later, 500 ng of vector, pSin-EF2-puro-JUB or pSuper-puro-JUB-RNAi plasmids plus 150 ng E-cadherin-luciferase reporter construct and 5 ng pRL-TK Renilla plasmids were co-transfected into cells using Lipofectamine 2000 Reagent (Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s instructions. Forty-eight hours after transfection, reporter luciferase activity was measured using the Dual-Luciferase Reporter Assay (Promega), according to the manufacturer’s instructions.

ChIP assay. Cells were grown to 70–90% confluence, fixed in 1% formaldehyde, and harvested by scratching. ChIP assays
were performed using the EZ-CHIP kit (Upstate, Temecula, CA, USA) according to the manufacturer’s instructions. Quantitative real-time PCR was performed on immunoprecipitated DNA. The primer sequences for detecting E-cadherin promoter are: forward, 5’-AGGTGAACCCCTCAGCCAATC-3’; reverse, 5’-ACGGGCTGGAGTCTGAA-3’.

**Immunofluorescence staining.** Cells were seeded on cover-slips in 24-well plates (Costar; Corning Incorporated, Corning, NY, USA). After 24 h, the cells were washed with 1 × PBS and fixed with ice-cold methanol for 10 min at room temperature. Cell membrane was permeabilized with 0.2% Triton X-100 in PBS (PBS-T), blocked with 10% BSA in PBS-T for 30 min, then incubated with primary antibodies, E-cadherin, Vimentin (1:500, BD Biosciences), in 10% BSA for 1 h at room temperature. After three washes with PBS, the slides were incubated for 1 h in darkness with Rhodamine-conjugated secondary goat anti-mouse antibody (The Jackson Laboratory, Bar Harvot, Maine USA). After washing, the slides were counterstained with DAPI (Sigma Aldrich). Images were captured using the AxioVision Rel.4.6 computerized image analysis system.

**Statistical analysis.** Student’s two-tailed t-test was used for the statistical analysis. Statistical analyses were performed using the SPSS 13.0 statistical software package. Data represented mean ± SD. Pearson correlation statistical analysis was used to analyze the correlation between JUB and E-cadherin expression. P-values less than 0.05 were considered statistically significant.

**Results**

**JUB, but not the other members of the Ajuba family proteins, is upregulated in CRC.** Until now, the expression of Ajuba family proteins had not been PCR. First, we analyzed their expression, including JUB, WTIP and LIMD1, in a publicly-available dataset (http://www.ncbi.nlm.nih.gov/gds/). As shown in Figure 1a, we were surprised to find that JUB, detected by different probes, but not the other two members of the Ajuba family proteins, was highly upregulated in tumor tissues compared to the matched adjacent normal tissues (GSE32323, n = 34). Furthermore, we detected the expression of Ajuba family proteins, E-cadherin and Vimentin (Fig. S1), in normal colorectal epithelial cells and CRC cell lines. Compared to the normal colorectal epithelial cells, JUB was significantly overexpressed in both protein and mRNA levels; however, the expression of WTIP and LIMD1 showed no obvious changes (Fig. 1b,c).

**Ectopic overexpression of JUB induces epithelial–mesenchymal transition in colorectal cancer cells.** Snail was proved to repress E-cadherin expression to induce EMT. Previous reports have shown that JUB functions as a corepressor of Snail, but whether it could also induce EMT has not been considered. Thus, we investigated the role JUB in EMT. As shown in Figure 2(a), western blot analysis revealed that ectopic expression of JUB in SW480 cells significantly reduced the expression of the epithelial marker, indicated by E-cadherin, and enhanced the expression of the mesenchymal marker, indicated by Vimentin. This was further confirmed by immunofluorescence staining (Fig. 2b).

It is well known that EMT aids metastasis by augmenting the motility and invasiveness of tumor cells. Indeed, overexpression of JUB in SW480 cells led to more cells invading and migrating through the Matrigel-coated or non-coated membrane of the chamber compared to vector cells (Fig. 2c). Strikingly, in the 3-D spheroid invasion assay, SW480-JUB cells displayed morphologies that were much more invasive, presenting more outward projections compared with vector cells (Fig. 2d and Fig. S2a). These data demonstrate that JUB does promote EMT and invasiveness in CRC.

**Silencing of endogenous JUB represses epithelial–mesenchymal transition in colorectal cancer cells.** To further confirm the role of JUB in promoting EMT in the opposite way, SW620, a derivative of the SW480 cell line and a highly metastatic CRC cell line, which showed a high expression of JUB (Fig. 1b), was selected for depletion of the endogenous expression of JUB. Indeed, silencing of JUB repressed EMT in SW620 cells. Knockdown of JUB reduced expression of E-cadherin and enhanced expression of Vimentin, indicated by both western blotting and immunofluorescence staining assays (Fig. 3a,b). The motility and invasive ability were also impaired by silencing of JUB as fewer cells were presented in the under-surface of the transwell chamber membrane (Fig. 3c). Consistent with
the data detailed above, 3-D culture showed that silencing JUB reduced the invasiveness of SW620 cells, exhibiting fewer or no outward projections (Fig. 3d and Fig. S2b). Taken together, the abovementioned data strongly suggest that JUB promotes EMT in CRC.

Snail is essential for JUB-induced epithelial–mesenchymal transition and JUB is required for Snail-induced epithelial–mesenchymal transition. Langer et al. (2008) demonstrate that JUB acts as a corepressor of Snail and targets the promoter region of E-cadherin to repress its expression.21 We tested whether this occurred in our current CRC model. First, we confirmed the interaction of JUB and Snail in CRC cells (Fig. S3a). Then, we measured the activity of an E-cadherin promoter containing reporter,21 which contains three Snail-binding E-boxes driving luciferase expression. Indeed, upregulation of JUB decreased while downregulation of JUB increased the reporter activity in
both SW480 and SW620 cells (Fig. 4a). We noticed that overexpression of JUB increased while silencing JUB decreased the repression effect of Snail on E-cadherin promoter activity (Fig. S3b). In addition, JUB presented on the promoter of E-cadherin as shown by JUB ChIP (Fig. S3c). Furthermore, we silenced Snail in SW480-JUB cells or reintroduced Snail-ORF into SW620-JUB-Ri cells. As shown in Figure 4(b) and (c), silencing of Snail derepressed the expression of E-cadherin and repressed the motility and invasion abilities of SW480-JUB cells, which indicates that Snail is essential for JUB-induced EMT. However, overexpression of Snail in SW620-JUB-Ri cells only partially depressed E-cadherin expression and partially enhanced motility and invasion abilities (Fig. 4b,c), which means that JUB is also required for Snail-induced EMT.

Expression of E-cadherin inversely correlated with JUB expression in specimens. Finally, we examined whether JUB repressing E-cadherin expression to induce EMT identified in CRC cells is also evident in clinical CRC specimens. By analyzing 10 cases of freshly collected CRC tissue specimens, we found that specimens with higher expression of JUB showed lower expression of E-cadherin (Fig. 5a). Quantification and correlation analysis proved that expression of E-cadherin and JUB significantly correlated, inversely ($r = -0.649$, $P = 0.042$, Fig. 5b).

Discussion

In the current study, we investigated the role of Ajuba family proteins in CRC cells. First, we found, surprisingly, that only JUB but not WTIP or LIMD1 of the Ajuba family was highly upregulated in CRC in a publicly available dataset, and validated this in cell lines. Further investigation proved that JUB could modulate EMT program of CRC cells. SW480-JUB cells

![Fig. 4. Snail is required for JUB-induced epithelial-mesenchymal transition. (a) Transient luciferase reporter assay using luciferase driven by the E-cadherin promoter in SW480 and SW620 cells. (b) Western blot of the indicated proteins in SW480 and SW620 cells transfected with indicated plasmids or siRNA. β-actin was used as a loading control. (c) Quantification of migrated and invaded cells of Transwell assay in the indicated cells. Data are the mean ± standard deviation (SD) from three independent experiments. *P < 0.05 based on Student’s t-test.

![Fig. 5. Reverse correlated expression between JUB and E-cadherin in clinical specimens. (a) Western blot analysis of JUB and E-cadherin in clinical specimens. (b) Quantification and Pearson correlation statistical analyzing of JUB and E-cadherin expression.]
showed enhanced EMT, indicated by decreased expression of E-cadherin, increased expression of vimentin, and enhanced motility and invasiveness compared to vector cells. Silencing of JUB in SW620 cells dramatically impaired EMT, leading to increased expression of E-cadherin, decreased expression of vimentin, and impaired motility and invasiveness compared with control cells. Further investigation revealed that JUB requires Snail to induce EMT and is required for Snail-induced EMT. Correlation analysis between the expression of JUB and E-cadherin in clinical specimens validated the conclusion that JUB represses E-cadherin and promotes EMT. EMT is closely associated with stemness in several cancer models. Our preliminary experiment found that, in SW480 cells, overexpression JUB forms more and larger spheres as indicated by tumor sphere formation assay (Fig. S4). However, further confirmation and investigation is needed.

There are three members of Ajuba family proteins, JUB, LIMD1, and WTIP. Various reports have shown that Ajuba family proteins share a lot in common. They have been found to bind the same partners or the same complexes and to regulate the same cellular events, especially JUB and LIMD1. Ajuba family proteins share three tandem, homologous LIM domains at the COOH terminus, but contain divergent proline rich NH2 termini (preLIM region). Thus, it is possible and reasonable that Ajuba proteins possess specific roles for each other. For example, Witzel et al. report that JUB is the prominent Isl1-binging partner while LIMD1 shows the weakest interaction with Isl1. Using LAST2 as a bait, only JUB of the family was identified as a binding partner. Here, we also found, surprisingly, that only JUB, but not LIMD1 or WTIP, was highly upregulated in the samples we obtained and in the publicly-available dataset, which indicates a specific role of JUB in CRC progression. However, it is still possible that WTIP and LIMD1 might also regulate EMT in CRC, but, absolutely, they do not play the prominent role.

According to various studies, JUB may regulate EMT via contrary mechanisms. Loss of cell–cell adhesion is a crucial aspect of EMT. JUB was originally found as a cytosolic protein. It was reported that JUB was recruited to cadherin-depen-
dent cell junctions and was required for maintenance of E-cadherin adhesion. Thus, JUB seems to repress EMT rather than promote EMT. However, inhibition of Pak1, which directly phosphorylated Ajuba at Thr172, phenocopies depletion of Ajuba, which means that it is the phosphorylated JUB but not the un-phosphorylated JUB involving in E-cadherin adhesion maintenance. Recently it was proved that JUB contains a functional nuclear export signal and shuttles between cytoplasm and nucleus. Therefore, it is possible that phosphorylation of JUB modulated its subcellular localization, leading it to be recruited to cell junctions. JUB was proved to be a corepressor of Snail family in nuclear to induce EMT. JUB can recruit the effector PRMT5 to Snail to repress Snail-target genes, such as E-cadherin. Our data show that, in CRC, JUB was dramatically upregulated and it repressed E-cadherin expression to induced EMT as a corepressor of Snail, which is consistent with previous studies. However, we cannot rule out the possibility that it is a consequence of the balance between nuclear distribution of cytoplasmic and nuclear JUB, which requires further investigation.

In sum, in the present study we found an elevated expression of JUB in CRC. We proved that upregulated JUB induces EMT, leading to increased motility and invasiveness, by acting as a corepressor of Snail. Our current study indicates a novel and potential target for metastatic CRC.

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**Disclosure Statement**

The authors have no conflict of interest.

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Supporting Information

Additional supporting information may be found in the online version of this article:

Fig. S1. Western blotting analysis of E-cadherin and Vimentin in colorectal cancer (CRC) cell lines.

Fig. S2. Quantification of invasive spheroids. (a) Quantification of invasive spheroids in SW480-JUB and vector cells. (b) Quantification of invasive spheroids in SW620 JUB-RNA is and vector cells. Data presented as mean ± SD. *P < 0.05 based on Student’s t-test.

Fig. S3. JUB interacts with and functions as a corepressor of Snail. (a) Co-immunoprecipitation between JUB and Snail. IgG was used as a negative control. (b) Transient luciferase reporter assay using luciferase driven by the E-cadherin promoter in SW480. (c) ChIP analysis of JUB on the promoter of E-cadherin. Snail was used as a positive control. Data presented as mean ± SD. *P < 0.05 based on Student’s t-test.

Fig. S4. Tumor sphere formation assay analysis in SW480 cells.