Angiomotin-p130 inhibits β-catenin stability by competing with Axin for binding to tankyrase in breast cancer

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Abstract
Growing evidence indicates that Angiomotin (Amot)-p130 and Amot-p80 have different physiological functions. We hypothesized that Amot-p130 is a tumor suppressor gene in breast cancer, in contrast with the canonical oncogenicity of Amot-p80 or total Amot. To clarify the role of Amot-p130 in breast cancer, we performed real-time quantitative PCR, western blotting, flow cytometry, microarray, immunofluorescence, immunoprecipitation, and tumor sphere-formation assays in vitro, as well as tumorigenesis and limited-dilution analysis in vivo. In this study, we showed that Amot-p130 inhibited the proliferation, migration, and invasion of breast cancer cells. Interestingly, transcriptional profiles indicated that genes differentially expressed in response to Amot-p130 knockdown were mostly related to β-catenin signaling in MCF7 cells. More importantly, most of the downstream partners of β-catenin were associated with stemness. In a further validation, Amot-p130 inhibited the cancer stem cell potential of breast cancer cells both in vitro and in vivo. Mechanistically, Amot-p130 decreased β-catenin stability by competing with Axin for binding to tankyrase, leading to a further inhibition of the WNT pathway. In conclusions, Amot-p130 functions as a tumor suppressor gene in breast cancer, disrupting β-catenin stability by competing with Axin for binding to tankyrase. Amot-p130 was identified as a potential target for WNT pathway-targeted therapies in breast cancer.

Introduction
Breast cancer (BCa) is the most common cancer in the female population, showing the highest incidence and prevalence among female cancers1. Although precision therapy has improved BCa survival, most patients inevitably suffer from disease recurrence or metastasis. It is, therefore, important to explore the potential mechanism underlying breast carcinogenesis.

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in BCa cells, and DNA vaccines targeting Amot-p80 inhibit tumor growth and metastasis in vivo. However, Amot-p130 has been shown to inhibit the proliferation of non-cancerous breast epithelial cells.

Amot isoforms have distinct physiological functions. During embryonic development, Amot-p80 is expressed early, whereas Amot-p130 is expressed later. In endothelial cells, Amot-p80 is found at the leading edge of migrating cells and diffuses throughout the cytoplasm when not migrating, whereas Amot-p130 is primarily located at cell junctions. The difference between the isoforms is also apparent in the regulation of endothelial cell migration, in which Amot-p80 and Amot-p130 play promotive and inhibitive roles, respectively. The Amot-p80/Amot-p130 ratio is used as an indicator of migration activity. We hypothesized that Amot-p80 and Amot-p130 have different functions in breast carcinogenesis. In a previous work from our group, we have shown that Amot-p130 decreases the motility of BCa cells. Here, we have investigated the link between the inhibition of metastasis and Amot-p130 in BCa.

Amot-p130 shows a high structural homology with AmotL2. AmotL2 inhibits WNT signaling by trapping β-catenin in recycling endosomes. However, it is unclear whether Amot-p130 regulates the WNT/β-catenin pathway. In the present study, the modulation of Amot-p130 expression revealed that Amot-p130 inhibited the cancer stem cell (CSC) potential of BCa, disrupting β-catenin stability by competing with Axin for binding to tankyrase (TNKS), leading to a further inhibition of cell proliferation and epithelial–mesenchymal transition (EMT) in BCa.

Results

Amot-p130 inhibits the proliferation of BCa cells

The basal expression of Amot-p130 varied significantly among the different BCa cell lines (Fig. 1a), showing lower expression levels in basal-like cell lines than in luminal cell lines. MCF7 with Amot-p130 knockdown (MCF7KD) and MM231 with Amot-p130 overexpression (MM231OE) cells were established using Amot-p130-targeted lentivirus (Fig. 1b) to determine the role of Amot-p130 in cell proliferation. The results of the cell count assay showed that MCF7KD cells grew faster, whereas MM231OE cells grew at a slower rate than control cells (Fig. 1c). Consistently, the extent of colony formation was higher in MCF7KD (42% vs 25%) and lower in MM231OE cells than in control cells (19% vs 37%) (Fig. 1d). An increase in the percentage of MCF7KD cells in S and G2/M phases occurred concomitantly with a decrease in MM231OE S- and G2/M-phase cells (Fig. 1e). Apoptosis was consistently decreased in MCF7KD cells (10.4% vs 7.1%) and increased in MM231OE cells (4.9% vs 11.6%) (Fig. 1f).

Amot-p130 inhibits the EMT of BCa cells

The activation of EMT allows cancer cells to acquire migratory, invasive, and CSC potential. In wound-healing assays, the recovery of confluency was faster in MCF7KD cells and markedly slower in MM231OE cells than in control cells at 24 h post-scratch (Fig. 2a). The number of cells migrating across the Matrigel was higher in MCF7KD (113) than in MCF7 control cells (67), whereas in MM231OE cells was less than one-third of those in the MM231 controls (21 vs 75) (Fig. 2b). The results of invasion assays showed that the number of invading cells was more than six folds higher in MCF7KD than in MCF7 cells (24 vs 150) and less than one-tenth in MM231OE than in MM231 cells (118 vs 6) (Fig. 2c). The EMT phenotype of cancer cells is marked by the balance between mesenchymal and epithelial markers. In MCF7KD cells, mesenchymal markers, including vimentin and Snail, were upregulated, whereas epithelial markers, including E-cadherin and ZO-1, were downregulated compared with those in MCF7 cells (Fig. 2d). By contrast, mesenchymal marker expression was lower, whereas epithelial marker expression was higher in MM231OE than in MM231 cells. Specifically, E-cadherin disappeared from the membrane and vimentin was transferred from the cytoplasm to the nucleus in MCF7KD cells (Fig. 2e). Consistently, E-cadherin appeared in the membrane and vimentin transitioned from a diffuse pattern to a dense dot in the cytoplasm of MM231OE cells. Collectively, these results suggested that Amot-p130 inhibits proliferation and EMT in BCa cells.

Identification of β-catenin as an effector of Amot-p130 in BCa

The molecular mechanisms underlying the function of Amot-p130 in BCa were examined using transcriptome microarrays. A total of 255 genes were upregulated and 251 genes were downregulated in MCF7KD cells compared with MCF7 cells (Fig. 3a, Supplementary Table S1). The annotations of these differentially expressed genes were significantly related to Wnt/β-catenin signaling in breast carcinogenesis (Fig. 3b). β-Catenin was identified as the pivotal link among all deregulated genes (Fig. 3c). All β-catenin downstream partners, including ID1, ID4, TCF4, CCND1, CTGF, ETS1, and AXN, were associated with CSC potential. Using real-time quantitative PCR, we confirmed that the expression of these genes was higher in MCF7KD than in MCF7 cells and lower in MM231OE than in MM231 cells (Fig. 3d). Taken together, these results indicate that Amot-p130 may affect the CSC potential of BCa by modulating β-catenin signaling.

Amot-p130 inhibits the CSC potential of BCa

Based on the microarray results, we assessed the effect of Amot-p130 on the CSC potential of BCa.
ALDH-positive populations or CD24-low/CD44-high populations are believed to contain breast CSC subsets\textsuperscript{25,26}. In vitro, the percentages of ALDH-positive cells increased from 2.79% to 22.8% in MCF7KD compared with MCF7 cells, whereas it decreased from 13.6% to 7.32% in MM231OE compared with MM231 cells (Fig. 4a, b). Similarly, the proportion of CD44-high/CD24-low cells was higher in MCF7KD (1.48% vs 0.70%) and lower in MM231OE (77.1% vs 88.2%) cells than in the controls. CSCs play an important role in drug resistance. Amot-p130 knockdown increased the resistance of MCF7 cells to tamoxifen, whereas Amot-p130 overexpression induced susceptibility to cisplatin in MM231 cells (Fig. 4c). In addition, the stem cell markers Sox2, Nanog, OCT4, YAP, and TAZ were upregulated in MCF7KD and downregulated in MM231OE cells (Fig. 4d). MCF7KD cells formed a two folds higher number of tumor spheres and with a larger size than the control cells, whereas MM231OE cells formed fewer (33 vs 76) and smaller spheres than control cells (Fig. 4e).

Next, we examined the ability of Amot-p130 to suppress tumorigenesis in vivo using a nude mouse xenograft model. MCF7KD cell injection induced the formation of larger tumors, whereas MM231OE cell injection induced smaller tumors than those in the respective control cells (Fig. 4f). The limited dilution-condition assay revealed that the rate of tumor formation was higher in MCF7KD and lower in MM231OE cells than in the respective controls (Fig. 4g). Immunohistochemistry (IHC) staining showed that E-cadherin was downregulated and vimentin was upregulated in MCF7KD cell-derived xenograft tissues (Fig. 4h). Conversely, MM231OE-derived xenograft tissues showed E-cadherin upregulation and vimentin downregulation. Taken together, these results indicated that Amot-p130 suppressed the CSC potential of BCa.
Amot-p130 inhibits the WNT pathway without direct interaction with β-catenin

To elucidate the role of Amot-p130 in the WNT/β-catenin pathway, the protein levels of β-catenin were assessed by western blotting. Both cytoplasmic and nuclear β-catenin levels were increased by Amot-p130 knockdown in MCF7 cells (Fig. 5a). TOP/FOP luciferase reporter assays confirmed the higher level of β-catenin-driven transcription in MCF7KD than in MCF7 cells (Fig. 5b). Opposite results were observed in MM231OE cells. The WNT pathway controls the transcription of many important genes, including Cyclin D1, TCF4, LEF, Met, and C-Myc. The protein levels of these downstream targets were upregulated in MCF7KD cells and downregulated in MM231OE cells (Fig. 5c). IHC analysis showed that β-catenin levels were upregulated in MCF7KD-derived xenografts and downregulated in MM231OE xenografts (Fig. 5d). These results indicate that Amot-p130 inhibits the WNT pathway. Immunofluorescence staining showed the co-localization of Amot-p130 with β-catenin at cell–cell junctions in MCF7 control and MM231OE cells (Fig. 5e). The close relationship between Amot-p130 and β-catenin being revealed in the present study, we investigated whether Amot-p130 directly interacts with
β-catenin. Co-immunoprecipitation assays showed that the anti-Amot-p130 antibody did not pull down β-catenin (Fig. 5f), suggesting that Amot-p130 inhibits WNT pathway activity without directly interacting with β-catenin.

Amot-p130 regulates β-catenin stability by competing with Axin for binding to TNKS

Next, we explored the mechanism by which Amot-p130 affects β-catenin. Firstly, cells were treated with cycloheximide (CHX) to inhibit protein synthesis, alone or in combination with MG132. MG132 was added to prevent the ubiquitination-mediated degradation of β-catenin. The results showed that β-catenin protein levels decreased with time after a single CHX treatment in all the cell lines. The addition of MG132 to CHX-treated cells prolonged the half-life of β-catenin in MCF7 and MM231OE groups, but not in MCF7KD and MM231 groups (Fig. 6a and Supplementary Fig. S1a). This suggested that Amot-p130 was involved in the degradation of β-catenin. Next, the cells were treated with XAV939 to block the acetylation of Axin by TNKS and then induce the phosphorylation of β-catenin by the Axin-containing destruction complex. β-catenin levels only decreased significantly in MCF7KD and MM231 cells (Fig. 6b and Supplementary Fig. S1b). Lastly, the cells were treated with SKL2001, a β-catenin stabilizer, which resulted in β-catenin levels increasing significantly only in MCF7 and MM231OE cells (Fig. 6c and Supplementary Fig. S1c). Specifically, XAV939 treatment reduced the levels of nuclear β-catenin in MCF7KD and MM231 cells, whereas
Fig. 4 (See legend on next page.)
Fig. 4 Amot-p130 inhibits the CSC potential of breast cancer cells. a ALDH-positive population was detected by flow cytometry (left). The proportions were calculated from three independent experiments (right). b CD24-low/CD44-high populations were detected by flow cytometry (left). The proportions were calculated from three independent experiments (right). c The tolerance of MCF7 cells to tamoxifen and MM231 cells to cisplatin was determined by MTT assay. d Stemness markers were assessed by western blotting. GAPDH was used as the loading control. e The formation of tumor spheres was recorded under microscopy (left). Scale bar = 50 µm. The proportions were calculated from three independent experiments (right). f Xenograft formation in NOD-SCID mice evaluated at 6 weeks after injection. Tumor volumes were determined using the formula volume = 1/2 × length × width². g The number of outgrowths was recorded at the end of the 6th week using limited-dilution assays. h IHC analysis of protein expression of Amot-p130, E-cadherin, and vimentin in xenograft tissues. Scale bar = 200 µm. All P values were calculated by paired Student’s t test. *P < 0.05, **P < 0.01. IHC, immunohistochemistry; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide

Fig. 5 Amot-p130 inhibits WNT pathway activation in breast cancer cells. a Protein levels of total β-catenin, cytoplasmic β-catenin, and nuclear β-catenin as determined by western blotting. GAPDH was used as the loading control for the total and cytoplasmic protein. Lamin A was used as the loading control for nuclear protein. b β-Catenin-driven transcription activity was determined by TOP/FOP luciferase reporter assays. Normalization was based on internal Renilla luciferase activity. The final reporter activity was measured as TOP/FOP ratio and was expressed as the mean ± SD of three independent experiments. c Protein levels of WNT downstream targets were determined by western blotting. d β-Catenin expression was determined by immunohistochemistry staining in xenograft tissues. Scale bar = 200 µm. e Co-localization of Amot-p130 (red) and β-catenin (green) in cell–cell contacts was indicated by immunofluorescence confocal microscopy. DAPI was used for nuclear staining (blue). f The interaction between Amot-p130 and β-catenin was evaluated in MCF7 cells by co-immunoprecipitation assay. IgG was used as the negative control. ***P < 0.001. DAPI, 4′,6-diamidino-2-phenylindole
SKL2001 upregulated nuclear β-catenin in MCF7 and MM231OE cells (Fig. 6d and Supplementary Fig. S1d). The growth rates of MCF7 and MM231OE cells were slower in response to XAV939, whereas MCF7KD and MM231 cells grew faster upon SKL2001 treatment (Fig. 6e and Supplementary Fig. S1e).

TNKS plays an important role in the WNT pathway by directly binding to Axin. Recently, TNKS was also found to bind to the N-terminal domain of Amot-p130.27,28 Hence, we speculated a competitive relationship between Amot-p130 and Axin in binding to TNKS. Our co-immunoprecipitation assays showed that less Axin was pulled down by the TNKS antibody in MCF7 and MM231OE cells (Amot-p130 positive cells) than in MCF7KD and MM231 cells (Amot-p130 negative cells) (Fig. 6f and Supplementary Fig. S1f). In summary, a higher
proportion of TNKS bound to Amot-p130 than to Axin in Amot-p130 positive cells. Collectively, the data suggested that Amot-p130 regulates β-catenin stability by competing with Axin for binding to TNKS.

Discussion

The present study established the tumor suppressive role of Amot-p130 in BCa, in contrast with the canonical oncogenicity of Amot-p80 or total Amot. The results indicated that Amot-p130 inhibits WNT/β-catenin signaling by competing with Axin for binding to TNKS. As summarized in Fig. 7, Axin/TNKS binding induces the nuclear translocation of β-catenin and the activation of the WNT pathway (left panel). Amot-p130/TNKS binding allowed Axin to form β-catenin-destruction complexes, thereby inactivating WNT/β-catenin signaling (right panel) and inhibiting the CSC potential in BCa.

Unlike Amot-p130, the Amot-p80 isoform is oncogenic in BCa, as is total Amot8,29. Total Amot was expressed at higher levels in BCa tissues than in matched normal tissues, and BCa patients with high Amot expression were associated with a poorly differentiated disease, multiple metastases, and poor survival10. Additionally, in vitro studies show that total Amot promotes the proliferation and invasion of BCa cells3. The opposite roles of Amot isoforms in BCa are consistent with their roles in the regulation of endothelial cell migration15, as Amot-p130 stabilizes migration whereas Amot-p80 promotes migration. Migration is repressed in endothelial cells expressing only Amot-p130, whereas a relatively small amount of Amot-p80 compared with Amot-p130 is sufficient to induce a migratory phenotype20. Amot-p80 and Amot-p130 expression undergoes dynamic changes, and a balance between the expression of the two isoforms controls BCa phenotype. Amot-p80 expression is responsible for the advantageous oncogenic role of total Amot in BCa cell lines. We found that the ratio of Amot-p130 to Amot-p80 expression was high in luminal BCa cells, whereas it was low in triple-negative BCa cells. The tumor suppressive role of Amot-p130 was demonstrated in several cancers4,5. Clinically, low Amot-p130 expression is an independent indicator of poor survival in lung cancer patients30.

Amot-p130 is a key regulator of embryonic development and is involved in regulating visceral endoderm movement during mouse embryonic stem cell differentiation31,32. Amot-p130 loss leads to the differentiation of the inner cell mass33. Amot is one of the three genes showing expression alterations at the mRNA and protein levels in mesenchymal stem cells in the dermis of patients with psoriasis compared with healthy controls34.

However, little attention has been paid to the function of Amot-p130 in CSCs. Our work began by investigating
the involvement of Amot-p130 in cell growth and EMT in BCa cells. We showed that the differentially expressed genes induced by endogenous Amot-p130 knockdown were mostly associated with breast CSCs, such as ID4 and CTGF. We further demonstrated that Amot-p130 inhibits breast CSC potential, including the regulation of CSC-enriched populations, drug resistance, the expression of stemness markers, tumor-sphere formation, and xenograft outgrowth.

The opposite functions of Amot-p130 and Amot-p80 are mostly mediated by the N-terminal domain. This extra domain is involved in several protein interactions, such as the classical interaction between the PPXY and the WW domains of YAP; therefore, Amot-p130 is an important effector in the HIPPO pathway. In the present study, the WNT signaling pathway was ranked highest among the breast carcinoma-related pathways affected by Amot-p130 knockdown. Amot-p130 downregulated cytosolic and nuclear β-catenin, suppressed β-catenin-driven transcription, and decreased the protein levels of WNT downstream partners. The WNT pathway is crucial for regulating the self-renewal, migration, and tumorigenicity of breast CSCs. Growing evidence indicates that Motin family members play important roles in the WNT pathway. AmotL2 reduces cytosolic and nuclear β-catenin levels by redirecting β-catenin into recycling endosomes, decreasing the tissue size during zebrafish development. Mechanistically, the N-terminal glutamine-rich domain is essential for the interaction between AmotL2 and β-catenin. Moreover, the exogenous expression of Motin family members inhibits β-catenin-driven transcription in HEK293T cells.

Amot-p130 is a classical component of cell-junction complexes, and β-catenin plays a central role in adherens junctions. In the present study, Amot-p130 and β-catenin co-localized at the cell–cell junction, suggesting that the two molecules function together in junction formation. Given the similarity in structure between Amot-p130 and AmotL2, we hypothesized that Amot-p130 retains β-catenin at the cell–cell junction, leading to a further inhibition of WNT signaling. Consistent with the role of Amot-p130, the formation of cell junctions negatively affects cell growth, migration, and invasion. In this case, the level of total β-catenin would decrease upon Amot-p130 silencing, because most of the free β-catenin released from the junction complex would be degraded in the cytoplasm. This is not the case, however. No direct interaction between Amot-p130 and β-catenin was detected. Instead, Amot-p130 may indirectly regulate β-catenin stability, as suggested by the present results showing the effect of Amot-p130 on (1) the protein levels of β-catenin and p-β-catenin and (2) the reversal of the decreasing trend of β-catenin levels induced by a cotreatment with MG132 and CHX.

TNKS regulates the degradation of Axin through binding and acetylation. The specific PRPPVGEE sequence in the N-terminal region of Axin and the ankyrin repeat domain of TNKS are required and sufficient for their physical interaction. Similarly, Amot-p130 physically interacts with TNKS through the highly conserved sequence QEPQGQE at its N terminus. Compared with Amot-p130 negative cells, Amot-p130 positive cells showed (1) equal levels of the TNKS protein, (2) lower levels of Axin/TNKS binding, (3) decreased β-catenin degradation upon XAV939 treatment, and (4) decreased β-catenin phosphorylation upon SKL2001 treatment, leading to the nuclear translocation of β-catenin. We demonstrated that Amot-p130 competes with Axin for binding to TNKS in BCa. Hence, β-catenin-driven transcription and cell growth were significantly affected by SKL2001 treatment in Amot-p130 positive cells and by XAV939 in Amot-p130-negative cells. XAV939 treatment inhibits clone formation in MCF10A cells, but not in HEK293T cells. To some extent, these results support our hypothesis, because HEK293T has high levels of Amot-p130.

Future studies need to be focussed on the effect of WNT inhibitor and/or agonist treatment on CSC potential. Whether Amo-p130 expression in BCa tissues is associated with clinicopathological features, metastasis, and prognosis needs to be evaluated. Future bioinformatics analyses should consider the different Amot isoform types.

In conclusion, we showed that Amot-p130 functions as a tumor suppressor in BCa, targeting CSC potential and leading to the inhibition of cell growth and EMT. Mechanistically, Amot-p130 inhibited the WNT/β-catenin pathway, modulating β-catenin stability by competing with Axin for binding to TNKS. The present study provides a new mechanistic insight into the regulation of Axin protein homeostasis and suggests new avenues for WNT pathway-targeted therapies in BCa.

Methods
Cell culture and reagents
MCF7 and MM231 cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with 10% fetal bovine serum (HyClone) and maintained in a humidified incubator at 5% CO2 and 37°C. Lentivirus targeting human Amot-p130 (NM_00113490.1) were constructed by GeneChem (sequences, Table 1). Cells were infected using polybrene (Santa Cruz) and treated with puromycin for 3 weeks to generate stable Amot-p130-knockdown (MCF7KD) or Amot-p130-overexpression (MM231OE) cells. The antibody against Amot was synthesized by Genemed Synthesis Inc. The specific antibody against Amot-p130 was purchased from Santa Cruz Bio-Technology (sc-166924). Antibodies against β-catenin (#8084P), p-β-catenin (#9561T), YAP (#14074S), TAZ (#4883), vimentin (#5741P), E-cadherin (#3195P), OCT-4 (#8084P), p-β-catenin (#9561T), YAP (#14074S), TAZ (#4883), vimentin (#5741P), E-cadherin (#3195P), OCT-4
Table 1 Sequences of shRNA and primers used

| Name                  | Sequence                                      |
|-----------------------|-----------------------------------------------|
| Amot-p130 shRNA #1    | CATACACCAGCAAGCCACAGGGAAT                     |
| Amot-p130 shRNA #2    | CAAGAATCCCACAAGTTCCAGTTGAA                   |
| Amot-p130-F           | AGCCCTGGCTGACATTACACC                        |
| Amot-p130-R           | CTTACCATAGGTCGGAGTCTT                        |
| GAPDH-F               | CTCCCTCCACCTTTGACGCTG                       |
| GAPDH-R               | TCCTCTGTCTGTCGTTG                            |
| ETS1-F                | TGATTTTGCATCCCGTT                            |
| ETS1-R                | ACCGGCATGATCCGGACT                           |
| TCF4-F                | GAGCCAGCCATTCTCTTG                            |
| TCF4-R                | CAGGTCTTCACTACCCCTG                         |
| ID1-F                 | GGAATCCGGAAGTGGAAAC                         |
| ID1-R                 | GCCTCAGGACACACAGAT                          |
| IDA-F                 | CCACATCAGCGCAACAGAG                         |
| IDA-R                 | CTCCCCCTCTAGTGTCTCTG                         |
| CTNN1-F               | ACCTAGGAGCCCCAACAACAA                       |
| CTNN1-R               | TTCTCTGCGTGGACGGCC                         |
| ANXA6-F               | CTGACATAATACACCTCACGT                      |
| ANXA6-R               | TTGGCATCAAATACGGAGGAG                      |

(#2750), Nanog (#4903P), SOX2 (#3579P), C-myc (#5605), Cyclin D (#2978), LEF1 (#2230P), TCF4 (#2565), MET (#8198), Gsk3β (#12456T), and β-trcp (#4394S) were from Cell Signaling Technology. HRP-labeled GAPDH (HRP-60004) and antibody against lamin A (10298-1-AP) were from Proteintech Group. XAV939 and SKL2001 were from Selleck Chemicals.

**Cell count assay**

Cells were seeded in 24-well plates at a density of 4 × 10⁴ cells/well for MCF7 group and 2 × 10⁴ cells/well for MM231 group. After culturing for the indicated time points, cells were washed with phosphate-buffered saline (PBS), detached with 0.25% trypsin for 10 min at 37 °C, and resuspended with an equal volume of the growth medium. Viable cells were counted with a hemocytometer under an inverted microscope system. The experiment was performed in triplicate.

**Wound-healing assay**

Cells were seeded in six-well plates at a density of 1 × 10⁶ cells/well for MCF7 group and 5 × 10⁵ cells/well for MM231 group, and cultured in complete medium for 24 h. After the removal of the culture medium, a monolayer of the sub-confluent cells was scratched with a 200 μl pipette tip to create a wound area. The wounded monolayer was washed with PBS twice and cultured for 24 h in 2% fetal bovine serum (FBS) medium for MCF7 group or FBS-free medium for MM231 group. Cell migration into the wound area was monitored by inverted microscopy and photographed at the indicated time points. The experiment was performed in triplicate.

**Cell cycle and apoptosis assay**

For the cell cycle assay, 1 × 10⁶ cells were fixed overnight in 70% ethanol and stained using 10 μg/ml propidium iodide (Sigma-Aldrich) and 50 μg/ml RNase A (Sigma-Aldrich). For apoptosis assay, 1 × 10⁵ cells were collected and stained with 7AAD and Annexin V (#559763, Apoptosis Detection Kit; BD Bioscience). Both data were analyzed by flow cytometry (BD Biosciences).

**Migration and invasion assays**

Migration and invasion assays were performed using the BioCoat cell migration chamber (Corning), which consists of a 24-well companion plate with cell culture inserts containing a filter with 8 µm-diameter pores. The transwell for the invasion assay was coated with matrigel (1:4 dilution with FBS-free DMEM; Corning). The cells were trypsinized, suspended with FBS-free DMEM, and seeded at 2 × 10⁵/ml for the migration assay and at 2 × 10⁵/ml for the invasion assay. Hundred microliters of the cell suspension was added to the upper well, and 600 µl of DMEM containing 10% FBS was added to the lower well. The plates were placed in 5% CO₂ at 37 °C and observed every 4 h. When the appearance of cells was observed in the lower well, the cells in the upper wells were gently removed by scrubbing, fixed in 95% ethanol for 15 min, and stained with 0.4% crystal violet for 30 min. The migrated or invaded cells were subsequently photographed with a microscope. The experiment was performed in triplicate.

**Microarray**

Total RNA was purified from cells using Trizol and examined using Thermo NanoDrop 2000. After purification and amplification (GeneChip), the RNA was hybridized onto Human Gene 1.0 ST arrays (Affymetrix). Raw data were normalized using Robust Multichip Array and log2 transformed using BRB-ArrayTools v4.3.0. Differentially expressed genes between MCF7 and MCF7KD cells were defined by P < 0.05 and by a fold change > 1.5 (Supplementary Table S1). Gene Ontology and Ingenuity Pathway Analysis were performed to identify enriched functionally associated pathways.

**Tumor-sphere formation**

Cells were plated at a density of 10000 cells/well for MCF7 and 5000 cells/well for MM231 in ultra-low attachment six-well plates and cultured in DMEM/F12 medium containing 20 μg/ml epidermal growth factor and...
2% B27. Cells were maintained in 5% CO₂ at 37 °C, and
the medium was replaced every 3 days. The total number
of spheres greater than 80 μm in diameter was counted at
14 days under an inverted microscope.

**Luciferase reporter assay**

Cells were plated at a concentration of 5000 cells/well in
96-well plates. After overnight serum starvation, the cells were co-transfected with 0.2 μg TOPflash or FOPflash
plasmids (Addgene) and 0.1 μg Renilla TK-luciferase vector (Promega) as a control using Lipofectamine 2000 (Invitrogen). Luciferase activity was measured by the
Dual-Luciferase Reporter Assay (Promega) using a Glo-
max 96 Microplate Luminometer 48 h later. The firefly
luciferase activity was normalized to the internal Renilla
luciferase activity, and the β-catenin-driven transcription
was measured as TOP/FOP ratio.

**Surface staining by flow cytometry**

Trypsinized cells were washed twice with PBS, blocked
with 2% bovine serum albumin, and stained with anti-
CD44-APC conjugate (Biolegend) and anti-CD24-PE
conjugate (Biolegend) in dark at 4 °C for 30 min. After
washing twice, the cells were analyzed by flow cytometry
(Becton Dickinson) using Cell Quest Pro software. The
ALDEFLUOR kit (Stem Cell) was used to detect intra-
cellular aldehyde dehydrogenase (ALDH) enzyme activity.
One million cells were suspended in ALDEFLUOR assay
buffer containing the ALDH substrate and incubated for
60 min at 37 °C. As a negative control, an aliquot of each
cell sample was treated with diethylaminobenzaldehyde, a
specific ALDH inhibitor.

**MTT assay**

Cells were seeded in 96-well plates at a density of 5 ×
10⁵ cells/well for MCF7 group and 2 × 10⁵ cells/well
for MM231 group. Twenty microliters of 5 mg/ml 3-(4,5-
dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide
(MTT) solution (Sigma) was added to each well and
incubated for 4 h. After the removal of the supernatant,
150 μl of dimethyl sulfoxide was added to each well.
The plates were kept in the dark and slightly shaken for
30 min at room temperature. The OD values of each well
were measured by a microplate reader at a wavelength of
490 nm. The experiment was performed in triplicate.

**Xenograft and limited-dilution assays**

NOD-SCID female mice aged 4 weeks were bred at a
specific pathogen-free animal facility. For xenograft
assays, 5 × 10⁶ cells diluted in 100 μl PBS were injected
into the mammary fat pad of mice on both the flanks. For
limited-dilution assays, cells were injected at concentra-
tions of 5 × 10³, 5 × 10⁴, 5 × 10⁵, and 2 × 10⁶ cells into the
fat pads (n = 6). Mice were monitored for tumor
development twice a week using a caliper. All the mice
were euthanized at the end of the 6th week. Xenograft
tumors were measured, weighed, and analyzed by IHC as
previously described36. All animal procedures complied
with the guidelines of the Institutional Animal Use and
Care Committee and were approved by the Animal Ethics
Committee of Xi’an Jiaotong University.

**Other assays**

Real-time quantitative PCR (primers, Table 1) was
performed as previously described10. Assays including
western blotting, immunofluorescence, plate clone for-
mation, and immunoprecipitation were also done as
previously described36.

**Statistical analysis**

Statistical analyses were performed using GraphPad
Prism software (version 5.0). Data were analyzed by paired
Student’s t test and presented as mean ± SEM. P < 0.05 at
two sides was considered statistically significant. The
asterisks *, **, and *** stand for P less than 0.05, 0.01, and
0.001, respectively. Statistical calculations were derived
from at least three independent replicates.

**Data availability**

All the data generated or analyzed during this study are included in this
published article (and its supplementary information files).

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**Conflict of interest**

The authors declare that they have no conflict of interest.

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