INTRODUCTION

Tumor metastasis, the movement of tumor cells from a primary site to progressively colonize distant organs, is responsible for more than 90% of deaths of cancer patients. The metastasis of cancerous tumors relies on the release of circulating tumor cells (CTC) that migrate to distant sites and form secondary tumors. Compared with single CTC, CTC clusters, arising from the fragmenting of primary tumor cell clumps into the vasculature in the circulation, have 23 - 50-fold increased metastatic potential, and dissociating CTC clusters into single cells can suppress metastasis. The fundamental process of CTC cluster formation is termed "collective invasion": cancer cells invade the surrounding stroma as cohesive clusters. "collective invasion" is still not well understood.

One way for cancer cells to collectively invade is to rely upon the motility of migratory stromal cells, such as fibroblasts and macrophages. In fact, tumors also contain multiple subpopulations of tumor cells with distinct genotypic and phenotypic characteristics. Importantly, this cellular heterogeneity is associated with differences in metastatic potential and therapeutic response.

Collective invasion of cancer cells is the key process of circulating tumor cell (CTC) cluster formation, and greatly contributes to metastasis. Cancer stem-like cells (CSC) have a distinct advantage of motility for metastatic dissemination. To verify the role of CSC in the collective invasion, we performed 3D assays to investigate the collective invasion from cancer cell spheroids. The results demonstrated that CSC can significantly promote both collective and single-cell invasion. Further study showed that CSC prefer to move outside and lead the collective invasion. More interestingly, approximately 60% of the leader CSC in collective invasion co-expressed both epithelial and mesenchymal genes, while only 4% co-expressed in single invasive CSC, indicating that CSC with hybrid epithelial/mesenchymal phenotype play a key role in cancer cell collective invasion.

KEYWORDS
- cancer stem-like cells
- collective invasion
- hybrid epithelial/mesenchymal
- leader
- single invasion
Cancer stem-like cells (CSC) are subpopulations in the tumor that are endowed with the ability to self-renew and differentiate into non-stem cancer cells (NSCC) that comprise the bulk of the tumor. Studies suggest that CSC not only have stem-like cell characters but also demonstrate the potential of a distinct advantage for metastatic dissemination. An increased proportion of CSC in tumors correlated with the occurrence of metastasis and a reduced survival rate in patients. Comparison of gene signatures in CSC in tumors correlated with the occurrence of metastasis and a re-tumor cells and 16.7% of CTC possessed stem-like signatures, suggesting that CSC may participate in the initiation of metastasis. However, to our knowledge, there are no reports that shed light on the function of CSC in collective invasion.

Transitions between epithelial and mesenchymal phenotypes (the epithelial to mesenchymal transition [EMT] and its reverse, the mesenchymal to epithelial transition [MET]), are hallmarks of cancer metastasis. The connection among EMT, MET and stemness indicates that CSC come in two distinct states: "epithelial-like" and "mesenchymal-like." Recent studies have attempted to resolve this contradiction by suggesting that instead of the cells in pure epithelial (E) or pure mesenchymal (M) states, cells in hybrid E/M or partial EMT state (which express epithelial and mesenchymal markers concurrently) are most likely to gain stemness. In fact, many CTC that survive in blood exhibit hybrid E/M phenotypes, become resistant to drugs and exit the bloodstream more efficiently. Cells in the hybrid E/M phenotype have both epithelial (cell-cell adhesion) and mesenchymal (migration) traits, and, hence, may avail collective cell invasion and migration.

Here we hypothesized that CSC may play a key role in collective invasion. To test this hypothesis, we used 3D invasion assays with spheroids to identify the most invasive cancer cells in an unbiased fashion. The results demonstrated that the CSC promote collective invasion significantly by moving outside of the spheroids to lead the collective invasion. To further explore the properties of the leader CSC, cell lines with E-cadherin, N-cadherin and Nanog promoter labeling were generated. Co-expression of these genes in the leading CSC suggested that CSC is the leader, with hybrid E/M phenotype playing a key role in collective invasion.

## 2 | MATERIALS AND METHODS

### 2.1 | Cell culture

Human breast cancer cell lines MCF7 and MDA-MB-231 were purchased from ATCC. Cells were cultured at 37°C and 5% CO₂ in DMEM high glucose medium (HyClone) with 10% FCS (HyClone) and 1% penicillin-streptomycin (HyClone).

### 2.2 | Cells sorting

Cells were trypsinized into single cell suspension and were counted. For staining, samples were incubated with antibodies for 30 minutes at 4°C. Unbound antibody was washed off and cells were sorted by flow cytometry for no longer than 30 minutes post-staining on a BD AriaIIl. The antibodies used to obtain breast CSC were anti-CD44-PE, anti-Nanog-APC and anti-CD24-FITC (BD Pharmingen). The purity of isolated breast CSC was determined by standard flow cytometry analysis. The purity of isolated CD44⁺CD24⁻ CSC regularly exceeded 98%.

### 2.3 | Cells spheroids formation and invasion assay

Cells were removed from the cell culture dishes with trypsin and resuspended in sterile 0.25% methylcellulose solution in DMEM. The cellulose solution contained cells with different ratios needed at a concentration of 10⁴ cells per mL. Next, 20-μL droplets were plated onto the underside of 10-cm culture dishes or 24-well plates (Corning) and inverted culture plates to form cell spheroids by gravity in a 37°C incubator in 4 days. Compacted spheroids were collected, then embedded in extracellular matrix (medium/ Geltrex gel [BD Bioscience] mix at a concentration of 1.6 mg/mL) in 24-well plates (Corning) at 37°C and 5% CO₂. The gel was incubated for at least 30 min at 37°C before the spheroids were embedded. The spheroids were then imaged with Zeiss LSM710 at a magnification of ×10 and ×20 from 0-96 h post-embedding. Z-stack images spanning 100-150 μm were collected and image stacks were processed by ZEN software (Carl Zeiss) to yield maximum-intensity projections.

### 2.4 | Cell tracker

Cells were harvested by removing culture media from cells after centrifugation, and then suspended gently in pre-warmed CellTracker (Thermo Fisher)[1:1000 dilution] staining solution with culture medium, and incubated at 37°C for 20 minutes. After that, cells were centrifuged to remove the CellTracker solution and then added with culture medium for use.

### 2.5 | Immunofluorescence Staining

MCF7 cells were plated onto glass coverslips for 4 hours, fixed in 4% paraformaldehyde for 15 minutes at room temperature and then washed with PBS. Then, cells were incubated with E-cadherin-FITC-antibody and N-cadherin Alexa Fluor 350 Antibody (1:200, BioLegend) overnight at 4°C. After washing with PBS, cells were monitored using a confocal microscope (Zeiss, LSM710).

### 2.6 | Western blot

Cells were washed with PBS three times and scraped into a lysis buffer (Beyotime, Shanghai, China) with proteinase inhibitor
After centrifugation, the supernatant was collected. The concentration was determined using a bicinchoninic acid protein assay (Beyotime). Equal amounts of proteins were electrophoresed on a 6% or 8% Bis-Tris gel and transferred to nitrocellulose membranes (Beyotime). After being blocked by 5% skim milk in Tris-buffered saline, membranes were incubated with primary antibodies of rabbit anti-Zeb1 (1:200, Abcam, USA), mouse anti-twist (1:500, Abcam, USA), mouse anti-vimentin (1:500, Abcam, USA), mouse anti-Nanog (1:500, Abcam, USA), rabbit anti-Snail (1:500, Abcam, USA) or polyclonal anti–β-actin (1:1000, Amersham, USA) overnight at 4°C, followed by incubation with corresponding secondary antibodies at 37°C for 1 hour. The blot signal was detected using an ECL detection system (Amersham, USA).

2.7 | Spheroid image analysis

The invasive distance was defined as the mean distance of all points on the cells to the centroid of spheroid minus the radius. Distribution of different cells in spheroids was quantified by measuring both the total spheroid area around the outer perimeter and the inner spheroid core in ImageJ.

2.8 | Stable cell lines labeling

pE-cad-Ctrine, pN-cad-mCherry and pNanog-CFP expressing plasmids which comprises specific gene promoter (promoter information in supporting information) and a fluorescent label were transfected into MCF7 cells using Lipofectamine 2000 according to the manufacturer's instructions. Forty-eight hours after transfection, cells started to be selected by G418. After 2 weeks of selection, cells with signals were sorted and collected by FACS. Insertion of the target sites was analyzed by sequencing of genomic DNA using the following primers: E-cadherin sequence 1: AGACATTCTCTGATTATCCC, E-cadherin sequence 2: ATGGTGAGCAAGGGCGAGGAG; N-cadherin sequence 1: CCCGCCCT-CATTCCACA, N-cadherin sequence 2: ATGGTGAGCAAGGGCGAGGAG. Nanog sequence 1: AAAAAAGAAATGGCTGGTTAATTAT, Nanog sequence 2: ATGGTGAGCAAGGGCGAGGAG.

2.9 | Time-lapse image

To observe the E-cadherin/N-cadherin dynamics at the spheroid edges, spheroids cells expressing E-cad-Ctrine and pN-cad-mCherry, pNanog-CFP were prepared as described above. Leader and follower spheroids were imaged using a confocal microscope. Images were collected every 4 hours beginning at 20 hours post–embedding for 40 hours. To observe the cell spheroid assay with salinomycin added, spheroids were imaged using a Nikon inverted microscope with a live cell chamber at ×10 objective; images were captured every hour from beginning of embedding for 72 hours.

2.10 | In vivo experiments

Mice were purchased from the Beijing Laboratory Animal Center of China and maintained in the Animal Resources Center of the China Agriculture University. The protocol was approved by the Committee on the Ethics of Animal Experiments of the China Agriculture University (permit number: 2019-009-012). Nude mice were maintained in a specific pathologic-free environment. A specified number of NSCC or hybrid epithelial/mesenchymal CSC suspended in total 100-μL serum-free medium and Matrigel (1:1) were inoculated subcutaneously on the left and right side separately (5 animals per experimental group). For the in vivo tumorigenic assay, tumor incidence was monitored after cells were inoculated for 4 weeks. In addition, 1 × 10⁶ cells were intravenously injected into the tail vein of BALB/c mice (n = 5 for each group) to analyze the tumor cell metastasis ability. Mice were imaged by bioluminescence imaging (BLI).

2.11 | Statistics and reproducibility

The experimental data were analyzed using OriginLab. To exclude the influence of cell spheroid size on invasion, the cell spheroids with radius of 10-15 µm were counted and normalized. The results were shown as mean ± SD. Statistical differences were performed using the t test. When representative images are shown, the representatives are from at least three independent samples.

3 | RESULTS

3.1 | Different invasion types of cells on spheroid

CD44⁺CD24⁻ and Nanog⁺ cell subpopulations are two kinds of commonly studied CSC. Many studies have shown that the subpopulations have stronger abilities in spheroid formation and tumorigenesis. In MCF7 cells, the average proportion of CD44⁺CD24⁻ CSC was 1.49% (Figure S1); meanwhile, our results showed that 95.2% of the CD44⁺CD24⁻ CSC were Nanog⁺ cells (Figure 1A). Within this proportion, cells can invade the surrounding environment (Figure S2C). To probe the different types of invasion present in the total invasive population over time, we performed an invasion assay of tumor cells as reported previously. As illustrated in Figure S2A, tumor spheroids were embedded in a 3D matrix and imaged over time. There were 2 types of cell invasion observed. As shown in Figure S1B and S1C, cells in spheroids showed phenotypically heterogeneous invasions including single-cell invasion and collective invasion. We analyzed the cell number in different invasive
types, and the results showed that the cell number for collective invasion was more than that of single invasion (Figure S2D).

3.2 | The different collective invasive ability of cancer stem-like cells and non–stem cancer cells

To study the difference between CSC and NSCC in the collective invasion, spheroids made by pure CSC, pure NSCC, as well as a 50% mixture of CSC and NSCC at 1:1 were used for invasion assay experiments (Figure 1B). In the pure NSCC group, there is scarcely any invasion post–72 h (Figure 1B-D). While in the 50% CSC group, the probability of collective invasion is 49.4%, which is 19 times higher than that of the pure NSCC group (Figure 1C). The number of collective invasion cells per spheroid is 6.1, approximately 21 times more than that of the pure NSCC group (Figure 1D), and the number of single invasive cells is also more than that of the NSCC group (Figure 1E). In addition to breast cancer cell line MCF7, conducted the same research with triple-negative breast cancer cell line MDA-MB-231 (Figure S3A-C). In the pure CSC group, the probability and the number of collective invasion cells were both higher than those in the other two groups (Figure 1C,D). The results were similar for both Nanog+ and CD44+CD24− CSC for collective invasion (Figure 1C,D). With the CSC proportion increased, there were more collective invasive cells than single invasion cells exhibited (Figure 1E). In addition, the average invasion distance of cells increased in direct proportion to the CSC ratio (Figure S3D).

3.3 | More cancer stem-like cells distributed in the outer layer of cell spheroids

To explore how CSC enhanced collective invasion of the cells, and to determine the difference between CSC and NSCC in invasive behavior, we labeled CSC and NSCC in red and green with a cell tracker, respectively. In spheroids with mixed CSC and NSCC at 1:1, the red CSC gradually diffused to the outer layer of the spheroids during the invasion process (Figure 2A). The distributions of CSC and NSCC on the cell spheroids were counted by using circles with a radius which increased by 10 μm in turn from the center of spheroids at different
time points (Figure 2B,D). The results showed that CSC were significantly more distributed in the outer layer of the spheroids than NSCC 72 h post-invasion, while there was no difference in the distribution at the beginning (Figure 2C,E). The same phenomenon was also found in the MDA-MB-231 cell spheroids (Figure S4D-F). Meanwhile, there was no difference between the size of single CSC and NSCC, eliminating the disturbance of the size of CSC and NSCC on the distribution (Figure S4A-C).

3.4 Cancer stem-like cells acted as leaders to drive collective invasion

Invading cells displayed phenotypically heterogeneous, collective chain invasion with leader cells defined as the first cell of a chain with trailing follower cells. Moreover, from the invaded cells, the CSC, either Nanog+ or CD44+CD24−, were mostly located in the leader position of the invading chain, leading the collective invasions (Figure 3A,B and Figure S4G,H). By counting all invasive chains, the results showed that 81% and 85% of the leader cells were CD44+CD24− CSC or Nanog+, respectively (Figure 3C and Figure S4I), suggesting that CSC led the collective invasion.

To verify the impact of CSC proportion on collective invasion, we increased the proportion of CSC in cell spheroids, from 10% to 50% to 90% in total cell content (Figure S5A), respectively. In all groups, CSC were found at the leading tip of invasive chains, followed by other CSC and/or NSCC, to invade surrounding extracellular matrix collectively (Figure S5B). With the increase in the proportion of CSC, the relative number of NSCC in collective invasion also increased (Figure S5D). At the same time, we found that adding salinomycin could reduce the outer distribution of CSC on the cell spheroids and also reduce the collective invasion of cells (Figure S6).
The results showed that CSC and NSCC can have both collective and single-cell invasions (Figure S5C,D). From spheroids with 50% CSCs group, CSCs account for 57% in single invasive cells and 74% in collective invasive cells (Figure 3D). As CSC are involved in both single and collective invasion, we further explored the difference in the CSC contributing to the two distinct invasion patterns.

### 3.5 Hybrid E/M phenotype in invasive leaders

Previous studies have shown that there are epithelial and mesenchymal cells in metastatic cancer cells.\(^{36,37}\) We labeled epithelial marker E-cadherin promoter in pE-cad-Ctine and mesenchymal marker N-cadherin promoter in pN-cad-mCherry. In addition, pNanog-CFP was used for labeling the Nanog promoter (Figure S7). To verify the consistency of fluorescence and native expression, Native E-cadherin and N-cadherin protein expression in cells were confirmed by immunofluorescence using their antibodies (Figure S8). Nanog was confirmed by western blot using their antibodies (Figure S9). In the collective invasion, we found that most leaders expressed all of the three markers simultaneously (Figure 4A-C). Western blot results using their antibodies show that the hybrid epithelial/mesenchymal CSC also express ZEB1, Snail, Twist and vimentin (Figure S9). In the follower cells, it was rare for all three markers to be expressed at the same time (Figure 4D). Most follower cells only expressed E-cadherin alone. For single cell invasion, most single invasion CSC only expressed N-cadherin alone (Figure 4E); few cells expressed all three markers at the same time (Figure 4F).

Based on the time-lapse results, we further recalled the differences in the starting cells of different cell invasion patterns (Figure 5A,B). Results showed that 59.6% of starting collective invasion cells expressed all of the E-cadherin, N-cadherin and Nanog simultaneously (Figure 5D), and evolved into the leader cells. After 40 h of continuous observation of the invasion process, we found that the leader cells continued to express three genes from the beginning, driving follower cells to invade from cells spheroids. While in single cell invasions, few cells expressed all three markers at the same time, with approximately 23% of cells expressing Nanog alone and approximately 26% of cells expressing both Nanog and N-cadherin (Figure 5C).

Since finding the collective invasion led by CSC with hybrid E/M phenotype, we have been exploring the tumorigenesis of hybrid epithelial/mesenchymal CSC. Cultured NSCC and hybrid epithelial/mesenchymal CSC were serially transplanted into the mammary fat pad (MFP) of immunocompromised mice. The enhanced tumorigenic capacity with hybrid epithelial/mesenchymal CSC was assessed by tumorigenesis rate. The tumorigenesis rate of the side inoculated with hybrid epithelial/mesenchymal CSC was higher than that of the side inoculated with NSCC (Figure S10A). In addition, 1 × 10^6 cells were intravenously injected into the tail vein of BALB/c mice to analyze the tumor cell metastasis ability. Tumor growth and metastatic burden are the BLI value of signals from metastatic tissues; CSC also resulted in improved tumor growth and metastatic burden (Figure S10B).

### 4 DISCUSSION

Circulating tumor cell cluster formation involves collective cancer cell invasion, which has been the focus of many studies.\(^{5,11,12,38,39}\) Why do cancer cells collectively invade as a strand? One possible reason that has been proposed is that cells cooperate to promote
Multicellular packages may provide survival or invasion advantages to escape; studies have shown that metastasis is supported by the polyclonal metastasis of tumor clusters rather than single cell seeds. Compared with single CTC, CTC clusters, as a group of invasive cells, often show a greater survival rate and are accompanied by poor clinical prognosis. Previous studies have shown that the keratin-14 positive subpopulation plays a pioneering role in the collective invasion of breast cancer cells. Our results showed that breast CSC can augment the probability of collective invasions occurring and make the invasion distance longer, which may increase the probability of metastasis of a tumor.

As the exact markers could be cell line-dependent, numerous studies have shown that CSC have stronger migration and invasion ability, as well as the ability to differentiate into many different subpopulation of tumor cells. In the mammary epithelial cell (hMEC)-derived HMLER cells, CD44+CD24− cells with hybrid E/M cells have mammosphere-forming capacity and exhibit stem-like characteristics. In human breast carcinoma tissue, CD44+CD24− cells (mesenchymal CSC) were present at the tumor invasive edge, while an ALDH1+ epithelial-like and hybrid E/M BCSC population was localized in the interior close to the tumor stroma. In MCF7 and MDA-MB-231, CD44+CD24− has classically been defined as belonging to the cancer stem cell population that has the ability of metastasis initiation and tumor formation, as well as, according to many studies, higher drug and radiation resistance. Presently, the experimental data showed that CD44+CD24− CSC are gradually distributed in the outer layer of the MCF7 and MDA-MB-231 cell spheroids during the process, which makes it easier for CSC to become leaders. An increase in the CD44+CD24− population was also observed.
in taxanem, and anthracycline induced a chemotherapy-tolerant state, which explains a new mechanism of CSC against specific drugs.54 Salinomycin can selectively reduce the CD44+CD24− CSC population.32,55,56 We used Salinomycin to treat cell spheroids and found that the outer distribution of CSC could be reduced. This result suggested that eliminating CSC can block the collective invasion, suggesting a new idea for suppressing cancer metastasis. Our results also provide a basis for the study of CSC in collective invasion and new evidence for the role of CSC in tumor metastasis.

The most significant differences between single and clustered CTC involve the expression of cell-cell junction components, and stemness-related transcription factor networks being accessible in CTC clusters.3 We found that this difference may be due to the differential expression of invasive cells. The heterophilic junction between E-cadherin expressed by cancer cells and N-cadherin expressed by CAF enables cancer cell adhesion, migration and collective invasion.9 Recent studies have found that the specific epigenetic phenotype of the breast cancer cell subpopulation promotes the transition to a more aggressive cell state, in which typical mesenchymal markers are insufficient to mark leading cells.57 Similarly, the leading cells here lack complete EMT characteristics, supporting the notion that invasive cancer cells have partial EMT phenotypes.58 In the present study, more than half of the leader cells expressed both E-cadherin and N-cadherin; they also have the characteristics of stem cells with Nanog expression.

The connection of hybrid E/M cells to collective motility has been shown previously in 2D tissue culture motility.59 We further verified that CSC can, indeed, act as leaders to promote collective invasion without complete loss of epithelial morphology and complete acquisition of mesenchymal morphology caused effective metastasis (Figure 4); it is hard to see three positive co–expressions simultaneously in followers and in single invasion cells. In Ewald’s group findings, E-cad functions as an invasion suppressor, survival factor and promoter of metastasis, but cells can continue to express E-Cadherin in a hybrid state to invade the ECM, so the balance E-cad protein level dynamics or a partial-or-hybrid EMT state may be particularly important during systemic dissemination, invasion and early seeding.60 By acquiring partial EMT, cancer cells with hybrid E/M phenotypes can undergo collective invasion through their remaining epithelial character4,11 and enhance motility by achieving mesenchymal character.61-63 In addition, with stem cell characteristics, they have a stronger viability;43,64 this makes the collective invasion led by CSC with hybrid E/M phenotype more aggressive.

Together, our study provides key insights into the biology of cancer cell invasion and metastasis. CSC are actively involved in the progression of cancer and provide a new concept for the collective cancer invasion model. Through co–expression of E-cadherin and N-cadherin, CSC promoted collective cancer invasion and metastasis as leaders. Our present observations may predict a new strategy for cancer metastasis, including interventions targeting CSC.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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