Cancer-associated fibroblast heterogeneity in axillary lymph nodes drives metastases in breast cancer through complementary mechanisms

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Although fibroblast heterogeneity is recognized in primary tumors, both its characterization in and its impact on metastases remain unknown. Here, combining flow cytometry, immunohistochemistry and RNA-sequencing on breast cancer samples, we identify four Cancer-Associated Fibroblast (CAF) subpopulations in metastatic lymph nodes (LN). Two myofibroblastic subsets, CAF-S1 and CAF-S4, accumulate in LN and correlate with cancer cell invasion. By developing functional assays on primary cultures, we demonstrate that these subsets promote metastasis through distinct functions. While CAF-S1 stimulate cancer cell migration and initiate an epithelial-to-mesenchymal transition through CXCL12 and TGFβ pathways, highly contractile CAF-S4 induce cancer cell invasion in 3-dimensions via NOTCH signaling. Patients with high levels of CAFs, particularly CAF-S4, in LN at diagnosis are prone to develop late distant metastases. Our findings suggest that CAF subset accumulation in LN is a prognostic marker, suggesting that CAF subsets could be examined in axillary LN at diagnosis.
Breast cancers (BCs) are the most frequent cancers in women worldwide. Diagnosis is mainly based on molecular subtype (Luminal A/B, HER2 and triple-negative (TN) defined according to cancer cell expression of progesterone, estrogen and HER2 receptors), tumor size and grade, as well as axillary lymph node (LN) metastasis. Most BC deaths arise from distant metastases in bone, liver and lung1,2. LNs are first reached by tumor cells evading from primary tumors (PTs) and number of invaded LNs is a strong prognostic factor3. Tumor metastasis is a multi-step process, including local invasion, intravasation, migration in blood or lymph stream, extravasation and distant organ colonization. Among others, one mechanism involves cancer cell epithelial-to-mesenchymal transition (EMT), which involves TGFβ but also NOTCH and WNT signaling pathways4-7.

Tumors are complex ecologies composed of numerous cell types, which participate in tumorigenesis and modulate cancer cell invasiveness8-11. Cancer-associated fibroblasts (CAFs) are abundant and involved in many tumor hallmarks such as angiogenesis, tumor cell proliferation, treatment resistance, immunomodulation and metastases9,12-15. In particular, CAFs are well-described to enhance tumor invasion and metastases, especially in BC. Indeed, CAFs are able to secrete factors, such as TGFβ and CXCL12, that directly stimulate cancer cell proliferation, EMT and migration16-19. Moreover, CAFs indirectly promote tumor spread via angiogenesis induction through VEGF and IL69,20. Besides, CAFs can directly interact with cancer cells via heterotypic E-cadherin–N-cadherin (CDH1–CDH2) adhesion and drive their invasion21. They remodel the extracellular matrix (ECM)22,23 and produce spaces or even tracks followed by cancer cells24-26. Recently, lung epithelial cells were shown to acquire cancer-associated parenchymal-like feature in metastatic niche27.

CAFs are poorly described in metastases, including axillary LN metastases, with only few studies assessing some markers, such as podoplanin (PDPN) or α-smooth muscle actin (αSMA)27-30. CAFs are known to be heterogeneous in PTs31-34, but characterization of this heterogeneity and its link with CAF functions is still far from being understood. If the role of CAFs in tumor invasion is well established, here we investigated if and how CAF subsets act on metastatic spread. As we recently identified four CAF subsets (named CAF-S1 to -S4) in BC32, we investigated the analogy between normal stromal cells and CAF subsets in LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e). CAF-S2 and CAF-S3 subpopulations are thus detected in non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e). CAF-S2 and CAF-S3 subpopulations are thus detected in non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e). CAF-S2 and CAF-S3 subpopulations are thus detected in non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e). CAF-S2 and CAF-S3 subpopulations are thus detected in non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e).

Results
Metastatic breast cancer axillary LNs exhibit four CAF subsets.
To investigate CAF heterogeneity in BC metastatic axillary LNs, we first performed multicolor flow cytometry (fluorescence-activated cell sorting (FACS)) on freshly resected invaded LN, with their matched PTs as controls (Supplementary Table 1). We applied the same gating strategy on both tissues by first excluding debris, dead cells, doublets, hematopoietic (CD45+), epithelial (EPCAM+), endothelial (CD31+) and red blood cells (CD235a+) (Supplementary Fig. 1a). We then characterized the CD45−EPCAM−CD31−CD235a− fraction enriched in fibroblasts by studying five CAF markers: FAP (fibroblast activation protein α1), CD29 (Integrin β1), αSMA, PDGFRβ (platelet-derived growth factor receptor-β) and PDPN (Fig. 1a). On FACS data, we applied FlowSOM13, an unsupervised automated algorithm which orders cells according to their phenotypic similarities (Fig. 1b, c). FlowSOM clustered CAFs into four main branches and thus distinguished four CAF subsets (CAF-S1 to -S4) in invaded LNs (Fig. 1b and Supplementary Fig. 1b). Four branches were also detected in corresponding PT (Fig. 1c and Supplementary Fig. 1c), confirming data in breast and ovarian PT12,32,33. Although expression of CD29 in CAF-S3 was slightly lower and PDGFRβ in CAF-S4 slightly higher in LNs than in PTs, systematic pairwise comparison of each CAF marker protein level between LNs and PTs showed the identification of CAF-S1, -S2, -S3 and -S4 in metastatic LNs (Fig. 1d, e). Hence, these data reveal that four CAF subsets are detected in metastatic BC LNs.

As normal LN structure relies on a fibroblastic network constituted by fibroblastic reticular cells (FRCs) described as PDPN+ cells35, we investigated the analogy between normal stromal cells and CAF subsets in LNs. Even though non-invaded LNs were hardly accessible because almost fully used for diagnosis, we had access to two non-invaded specimens (Supplementary Fig. 1d), along with their matched invaded LNs. Non-invaded axillary LNs were clearly enriched in CAF-S2- and CAF-S3-like cells, while the matched invaded LNs showed a much higher proportion of CAF-S1 and CAF-S4 (Fig. 1f and Supplementary Fig. 1e). CAF-S2 and CAF-S3 subpopulations are thus detected in metastatic LNs, but also in non-invaded LNs. These results corroborated our previous data showing that CAF-S2- and CAF-S3-like cells are detected in healthy breast tissue32, suggesting that these CAFs might derive from normal resident fibroblasts. In that sense, the pattern of CAF-S3 in LNs was slightly different than the one detected in PTs, as observed with CD29 staining (Fig. 1b–d), suggesting that normal-like CAF-S2/S3 could be more affected by their tissue of origin than CAF-S1 and CAF-S4. In contrast to CAF-S2 and CAF-S3, CAF-S1 and CAF-S4 were strictly observed in invaded LNs and positively correlated with tumor cell invasion (Fig. 1g). Thus, these data highlight a potential link between both CAF-S1 and CAF-S4 and tumor cell invasion in LNs. In conclusion, we identified four CAF subsets in metastatic LNs defined as: CAF-S1: FAPHigh CD29Med-High αSMAHigh PDPNHigh PDGFRβHigh; CAF-S2: FAPneg CD29Low αSMANeg-Low PDPNβneg PDGFRβLow; CAF-S3: FAPNeg-Low CD29Med αSMANeg-Low PDPNβLow PDGFRβMed-Low; CAF-S4: FAPNeg-Low CD29High αSMAβHigh PDPNβLow PDGFRβMed. Besides, the amounts of both CAF-S1 and CAF-S4 subsets in LNs are linked to BC cell metastatic spread.

CAF-S1 and CAF-S4 are the most abundant subsets in metastatic LN. To decipher the link between CAF subsets and metastatic spread, we studied metastatic LN sections from a retrospective cohort of 124 BC patients (Supplementary Table 2). We analyzed invaded zones of metastatic LN, identified using EPCAM marker (Supplementary Fig. 2a). We first observed that LN stroma represented around 25–30% of invaded areas, independently of BC subtypes (Fig. 2a). We performed immunohistochemistry (IHC) of five CAF markers (FAP, CD29, PDGFRβ, αSMA) on serial LN sections (Fig. 2b, c). Here, we replaced PDPN by FSP1 because we could not find a PDPN-specific antibody for IHC, but we verified that PDPN and FSP1 markers recognized the same cells by FACS (Supplementary Fig. 2b). Histological scoring of each CAF marker demonstrated that invaded LNs from Luminal (Lum A and B) cases exhibited the lowest histological scores (H-scores) except for PDGFRβ, whereas both HER2 and TN LNs showed the highest H-scores.
(Fig. 2b, c and Supplementary Fig. 2c). When applying a decision tree algorithm to determine CAF subset enrichment (Fig. 2d), we found that 96% of metastatic LNs showed accumulation of CAF-S1 and CAF-S4 (Fig. 2e). Luminal LNs were mainly enriched in CAF-S4, while HER2 and TN cases displayed both CAF-S1 and CAF-S4 predominance. We observed that the median percentage of fibroblasts positive for FAP, SMA and CD29 (reflecting CAF-S1 identity) reached 75% of total CAFs in CAF-S1-enriched LNs, and that fibroblasts negative for FAP but positive for SMA and CD29 (reflecting CAF-S4 identity) reached 60% of total fibroblasts in CAF-S4-enriched LNs. We also developed an image analysis tool that combined spatial...
Fig. 1 Metastatic BC LNs exhibit four CAF subsets. a Representative FACS plots showing FAP, CD29, PDPN, αSMA and PDGFRβ protein levels in DAPI+ EPCAM− CD45− CD31− CD235a− cells from a primary breast tumor (PT, left) and its corresponding metastatic axillary lymph node (LN, right). b, c FlowSort trees built on CAF from LNs (b, n = 20) and PTs (c, n = 16) and annotated for each CAF marker expression. Colors show CAF marker intensities. Node size depends on the number of phenotypically close cells. d Specific mean fluorescence intensity (speMFI) for each marker per CAF subset in PT and LN. Values are in log2 adjusted with offsets per marker. Each dot represents one sample (n ≥ 13 PT/LN pairs). Boxplots are median ± 25%-75% quantiles, whisker values range 1.5 × IQR above 75th or below 25th percentiles. p Values from Wilcoxon signed rank test. e Correlation plots between each marker speMFI in PT and LN, matched by patient and CAF subset (n ≥ 13 PT/LN pairs). p Values from Spearman’s test. f Same as in a for an invaded axillary LN (left) and its corresponding non-invaded LN (right). g Correlation plots between the percentage (%) of each CAF subset among total CAF and EPCAM+ cells among live cells, in invaded axillary LN (n = 19). p Values from Spearman’s test. Source data provided in Source Data file, with R scripts used.

registration and computational analysis to generate maps of CAF subsets at cellular level in invaded LNs (Fig. 2f and Supplementary Fig. 2e). Representative pictures of CAF subset maps confirmed the enrichment in CAF-S1 or CAF-S4 in invaded LNs. Interestingly, among the 124 LNs analyzed, we had access to 41 corresponding PTs (Lum A, HER2, TN) for matched analyses (Fig. 2g and Supplementary Fig. 2e). We found that αSMA H-scores were similar between PTs and LNs, whereas CD29 and PDGFRβ were higher and FSP1 lower in LNs (Supplementary Fig. 2e). When we combined H-scores using the decision tree algorithm, we found that CAF enrichments in metastatic LNs were significantly different from those in matched PTs, with a strong increase in CAF-S4 in LNs (Fig. 2g). Similarly, when comparing invading LNs to PTs (unmatched samples) from a larger BC cohort with at least one invading LN at diagnosis (N+) (Supplementary Table 2), we confirmed the accumulation of CAF-S4, the increase in CAF-S1 and the decrease in CAF-S2 and CAF-S3 in invaded LNs compared to PTs (Fig. 2h). Metastatic LNs were particularly enriched in CAF-S4 compared to PTs in Luminal A and TN subtypes (Supplementary Fig. 2i), possibly because HER2 PTs LNs were particularly enriched in CAF-S4 compared to PTs in metastatic LN stroma is enriched in myofibroblastic CAF-S1 and CAF-S4 (Fig. 2f), thereby confirming the existence of pericyte-like CAF and their potential pericyte origin32,38–41. Moreover, there was a strong overlap between PT and LN signatures for each CAF subset (Fig. 3d), suggesting that CAF-S1 and CAF-S4 from metastatic LN are molecularly close to CAF-S1 and CAF-S4 from PTs (Supplementary Tables 4 and 5).

CAF-S1 and CAF-S4 exhibit distinct migration and invasion properties. As invaded LNs show predominantly CAF-S1 and CAF-S4 fibroblasts, we hypothesized they could be involved in metastatic spread and we analyzed their properties by functional assays. We established CAF-S1 and CAF-S4 primary cell cultures from BC patients. We first confirmed by RNAseq on four CAF-S1 and CAF-S4 pairs (three couples from PTs and one from LN) that, after several passages in culture, these cells kept similar molecular identities as those detected without culture (Supplementary Fig. 3a–c). Moreover, we confirmed by FACS that CAF markers at protein levels corroborated data from fresh samples (Supplementary Fig. 3d). We next deciphered CAF-S1 and CAF-S4 intrinsic characteristics. We first observed that CAF-S1 proliferated slightly faster than CAF-S4 (median doubling time = 2.1 days for CAF-S1 and 3.0 days for CAF-S4) (Fig. 4a). Moreover, CAF-S1 displayed higher migration skills compared to CAF-S4, as assessed by Transwell assays (Fig. 4b). We validated this result using cell exclusion zone assays by manually tracking cells and calculated their velocity, persistence and directionality. CAF-S1 were faster and more persistent than CAF-S4, CAF-S1 MCAM-positive and CAF-S4 MCAM-negative fibroblasts in invaded LNs (Supplementary Fig. 2f), thereby confirming the existence of pericyte-like CAF and their potential pericyte origin32,38–41. Moreover, there was a strong overlap between PT and LN signatures for each CAF subset (Fig. 3d), suggesting that CAF-S1 and CAF-S4 from metastatic LN are molecularly close to CAF-S1 and CAF-S4 from PTs (Supplementary Tables 4 and 5).

CAF-S1 and CAF-S4 subsets show similar signatures between PTs and metastatic LNs. As CAF-S1 and CAF-S4 subsets were largely predominant in LNs, we next aimed at characterizing these LN CAF subsets on a molecular basis and compared them to PT CAF subsets. CAF-S1 and CAF-S4 from both PTs and LNs (n = 5 pairs) were FACS-sorted by excluding dead, hematopoietic, epithelial, endothelial cells and erythrocytes and analyzing CAF markers (using the same gating strategy as in Supplementary Fig. 1). We performed RNA sequencing (RNAseq) on CAF-S1 and CAF-S4 fibroblasts, and EPCAM+ cancer cells sorted from PTs and matched invaded LNs (Supplementary Table 1). Unsupervised principal component analysis (PCA) and hierarchical clustering (HC) built on the 500 most variant genes distinguished tumor cells, CAF-S1 and CAF-S4 fibroblasts (Fig. 3a, b). The first PCA component (51% variance) differentiated EPCAM+ cells from CAF, and the second (23%) variance CAF-S1 from CAF-S4 (Fig. 3a). HC enabled us to visualize specific transcriptomic profiles (Fig. 3b). PCA and HC showed that samples from PTs and LNs were mixed within each cellular population, highlighting that differences between CAF subsets (CAF-S1/CAF-S4) were higher than between tissue of origin (PT/LN). We performed PCA on each population on their respective 500 most variant genes and still noted that PT and LN samples did not segregate (Fig. 3c). In line with published data3,41, tumor cells clustered with their matched EPCAM+ LN cells, suggesting that tumor cell transcriptomes vary more between patients than between PTs and LNs. In contrast, patient effect was less detectable in CAF-S1 or CAF-S4 (Fig. 3c). We next performed paired differential analyses of CAF-S1 and CAF-S4 considering either PT or LN samples. CAF-S1 upregulated genes were mainly involved in ECM organization and CAF-S4-specific genes in muscle contraction (Supplementary Table 3 and Fig. 3d). As CAF-S4 exhibited pericyte-like signature, we confirmed that we could detect MCAM-positive fibroblasts in invaded LNs (Supplementary Fig. 2f), thereby confirming the existence of pericyte-like CAF and their potential pericyte origin32,38–41. Moreover, there was a strong overlap between PT and LN signatures for each CAF subset (Fig. 3d), suggesting that CAF-S1 and CAF-S4 from metastatic LN are molecularly close to CAF-S1 and CAF-S4 from PTs (Supplementary Tables 4 and 5).
Spheroids made by CAF-S4 pulled more efficiently on collagen fibers than CAF-S1 (Fig. 4e). To further quantify the traction forces exerted by CAF on matrix, we performed traction force microscopy (TFM). We observed that CAF-S1 were more elongated compared to CAF-S4 that were larger (Fig. 4f and Supplementary Fig. 3f, h). Moreover, the strain energy (that measures the energy spent to deform the substrate) developed by CAF-S4 was more than twofold higher than that of CAF-S1 (Supplementary Fig. 3g). To take into account differences in surface area between CAF-S1 and CAF-S4 (Supplementary Fig. 3i).
Fig. 2 Metastatic LNs are enriched in CAF-S1 and CAF-S4. a Percentage (%) of stroma in invaded LN according to BC subtypes (n = 124), LumA (violet), LumB (blue), HER2 (light gray), Lum B/HER2 (dark gray), TN (black). b Histological scores (H-Scores) for each CAF marker in invaded LN according to BC subtypes (n = 124). c Representative images of CAF marker staining on serial LN sections (LN#1 HER2; LN#2 TN). Scale bar, 50 μm. d Decision tree defining CAF subset identity based on quartile (Q) and median (Mdn) distribution of CAF markers. Thresholds and decision rules were first established on FACS data and next applied to IHC data. e Repartition of CAF subset enrichments (CAF-S1 (red), CAF-S2 (orange), CAF-S3 (green) and CAF-S4 (blue)) in LN according to BC subtypes (n = 124). f Representative views of CAF marker immunostaining on serial LN sections used for building maps of CAF subsets at cellular scale using the decision tree algorithm, shown in d. CAF-S1 are in red, CAF-S4 in blue and epithelial cells in black. CAF-S1- and CAF-S4-enriched LNs are shown. Scale bar, 200 μm. g Repartition of CAF subset enrichments in PTs and matched LNs (n = 41 pairs). h Same as in g with unmatched samples (N − cases, n = 75 PT and 84 LN). i Same as in h according to BC subtypes (Lum A: n = 38 PTs, 30 LNs; HER2: n = 16 PTs, 26 LNs; TN: n = 21 PTs, 28 LNs). j Contingency table showing repartition of CAF subset enrichments in PTs and corresponding invaded LNs (n = 41 pairs). In all panels, boxplots are median ± 25%–75% quantiles, whisker values range 1.5 × IQR above 75th or below 25th percentiles. b p Values from Mann–Whitney test. e, g, h p Values from Fisher’s exact test. Non-significant p values are not mentioned. Source data provided in Source Data file, with R scripts used.

Fig. 3 CAF subsets show same identity in PTs and metastatic LNs. a PCA based on the 500 most variant genes from RNAseq data of CAF-S1, CAF-S4 and EPCAM* cells sorted from paired PT and LN (n = 5 pairs for CAF-S1 and CAF-S4; n = 4 pairs for EPCAM* cells). b Hierarchical clustering on the same samples and same genes as in a using Ward’s method with Pearson distances. Rows represent samples and columns genes. Color saturation shows gene expression deviation from the mean (above in red, below in blue). c PCA based on the 500 most variant genes from RNAseq data of EPCAM* cells (n = 4 PT/LN pairs, top left), CAF-S1 (n = 5 PT/LN pairs, top right) and CAF-S4 (bottom left: n = 5 PT/LN pairs; bottom right: PCA restricted to n = 8 samples, as indicated). d Venn diagram showing overlaps between the transcriptomic signatures of CAF-S1 from PT, CAF-S1 from LN, CAF-S4 from PT and CAF-S4 from LN. Circled numbers show common genes between tissue localization for each cell type. p Values from hypergeometric test indicate significance of overlaps between subgroups. Source data provided in Source Data file, with R scripts used.
Fig. 3h), we compared the strain energy densities (strain energy normalized to the cell surface) and confirmed that CAF-S4 deployed more contractile energy per surface unit than CAF-S1 (Fig. 4g). Similar results were obtained using mean traction stress amplitude (mechanical forces cell develop per unit area to deform the substrate; arrows Fig. 4f, h). Collagen fiber density was also increased around CAF-S4 compared to CAF-S1 cells (Fig. 4i), which is consistent with CAF-S4 capacity to exert higher traction forces on their environment, to pull and contract collagen fibers. Overall, these data demonstrate that CAF-S1 are motile, while CAF-S4 are contractile with strong capacity to pull on collagen fibers.

**CAF-S1 and CAF-S4 promote tumor cell spread by distinct mechanisms.** While CAF function in metastases was investigated in the past, the specific role of CAF subsets on metastatic spread...
remains unknown. We assessed CAF-S1 and CAF-S4 effects on BC cell properties by using MCF7 and T47D luminal cell lines, as CAF switch between PTs and LNPs was prominent in luminal cases. Co-culture of BC cells with CAF-S1 or CAF-S4 increased the number of tumor cells, with a stronger impact of CAF-S1 compared to CAF-S4 (Fig. 5a). This effect was also detected with CAF-S1- or CAF-S4-conditioned medium (CM), although at a lower extent (Fig. 5b). CAF subset proliferation increased at long time points of co-culture with cancer cells (Fig. 5c), but not at short term (Supplementary Fig. 4a). Transwell assays showed that BC cell migration was strongly increased toward CAF-S1, but at a much lower extent toward CAF-S4 (Fig. 5d). The CAF-S1-pro-migratory effect was independent of its pro-proliferative effect, while the slight impact of CAF-S4 was proliferation-driven (Supplementary Fig. 4b). In agreement with their intrinsic features (Fig. 4), CAF-S4 fibroblasts (easily distinguished from cancer cells by F-actin staining) exhibited more F-actin stress fibers and Vinculin-stained focal adhesions than CAF-S1 (Fig. 5e and Supplementary Fig. 4c for quantification). Interestingly, in presence of CAF-S1 or CAF-S4, cancer cells were more scattered (Fig. 5e and Supplementary Fig. 4c). Indeed, in absence of CAF-S1, BC cells were less cohesive and more dispersed. With CAF-S4, cancer cells were also less packed but with numerous remaining F-actin interconnections. In agreement with these observations, E-cadherin/CDH1 protein levels decreased at cancer cell membrane in presence of CAF-S1, effect which was also visible but less pronounced with CAF-S4 (Fig. 5f and Supplementary Fig. 4d). Thus, these results suggest that CAF-S1—and to a lower extent CAF-S4—promote cancer cell spreading by initiating first steps of EMT.

As CAF-S4 properties were linked to 3D environments, we moved to such systems to analyze their impact on BC cell invasiveness in surrounding matrix. We first used an inverted Transwell assay to test CAF subset-mediated matrix remodeling on BC cell invasion (Fig. 6a–f). In this assay, cancer cells need to cross the membrane to invade the matrix. MCF7 and T47D cells are not invasive enough to obtain quantifiable results, we thus chose to work with MDA-MB-231, a TN invasive cell line. The highest z-distance covered by cancer cells in the matrix was significantly increased in CAF-S1- and even more in CAF-S4-embedded collagen (Fig. 6a, b). If cancer cells browsed in average 72 μm above the membrane in a CAF-free matrix, they reached 109 μm in presence of CAF-S1 and 129 μm with CAF-S4 (Fig. 6b). We next looked at the average distribution of cancer cell frequency along z-axis, every 10 μm from the membrane up to 200 μm. Major part of cancer cells was below 30 μm, distance from which a major drop was observed (Fig. 6c). Interestingly, beyond 30 μm, CAF-S4 significantly increased the proportion of cancer cells compared to control (Supplementary Fig. 4e, left).

Similar results were obtained when increasing the invasion threshold, with an even greater impact of CAF-S4 on tumor cell invasion (Fig. 6d and Supplementary Fig. 4e, right). To go deeper in the characterization of BC cell motility in 3D, we took advantage of the tumor-on-chip system, a microfluidic device where cancer cells and CAF subsets were co-cultured in 3D and followed by time-lapse video microscopy. In this system, we reproduced the intrinsic CAF subset motility, CAF-S1 being more motile than CAF-S4 (Supplementary Fig. 4f). In addition, velocity of GFP-labeled BC cells increased in presence of CAF-S4, and at a lower extent CAF-S1 (Fig. 6e, f). In tumor-on-chip, CAF subset-mediated effect was more important on T47D (Fig. 6f) than MCF7 (Fig. 6e), probably because T47D cells are less motile at basal levels than MCF7 (around 0.1 μm/min for MCF7 and 0.05 μm/min for T47D). Taken together, these results show that both CAF-S1 and CAF-S4 stimulate BC cell motility, via different effects: CAF-S1 initiate first steps of EMT and secrete factors that attract cancer cells, while CAF-S4 remodel the matrix and promote cancer cell invasion in 3D.

CAFS1 and CAF-S4 induce cancer cell invasion by TGFβ/CXCL12 and NOTCH. We next sought to define the molecular actors involved in CAF-S1 and CAF-S4 pro-invasive effects. As CAF-S1 attract cancer cell at distance and secrete high levels of CXCL12 (Supplementary Fig. 5a), we first analyzed the impact of CXCL12 silencing in CAF-S1 and CAF-S4 (Supplementary Fig. 5b) on BC cell proliferation and migration. Interestingly, CAF-S1 silenced for CXCL12 partly lost their ability to initiate EMT in cancer cells, while CXCL12-silencing had no impact in CAF-S4 (Fig. 7a). Indeed, BC cells were more clustered and E-cadherin protein levels in tumor cells increased after CXCL12 silencing in CAF-S1 but not in CAF-S4 (Fig. 7a and Supplementary Fig. 5c). We also observed that CXCL12-silencing in CAF-S1 decreased BC cell chemo-attraction without affecting their proliferation rate (Supplementary Fig. 5d, e). In agreement with these observations, even partial inhibition of CXCR4 in cancer cells (Supplementary Fig. 5f) affected their scattering and EMT initiation by CAF-S1 (Fig. 7b). This suggests that both CXCL12 secretion by CAF-S1 and CXCR4 expression in cancer cells are involved in CAF-S1-mediated EMT initiation in BC cells. As we observed an upregulation of the TGFβ-signaling pathway in CAF-S1 (Supplementary Fig. 5g), we wondered whether this pathway, strong EMT inducer, could also be involved in CAF-S1-mediated EMT initiation. As the expression of TGFβ receptors (TGFBR1, 2 and 3) was upregulated in CAF-S1 cells compared to CAF-S4 (Supplementary Fig. 5h), we compared the impact of TGFBR inhibitor on CAF-S1- and CAF-S4-induced EMT initiation on cancer cells (Fig. 7c and Supplementary Fig. 5i). TGFBR inhibition affected initiation of EMT in cancer cells in presence of...
CAF-S1 cells, while it had no impact on cancer cells alone and a weak effect with CAF-S4 (Fig. 7c). Taken together, these results show the role of TGFβ and CXCL12 signaling pathways in CAF-S1-dependent pro-tumorigenic phenotype.

Concerning CAF-S4, NOTCH pathway and three out of the four NOTCH receptors (NOTCH 1–3) were significantly upregulated in CAF-S4 (Supplementary Fig. 6a, b). We observed that DAPT, a γ-secretase inhibitor and pan-NOTCH inhibitor, severely reduced CAF-S4 contractile capacity assessed in collagen gel assays, while DAPT had no impact on CAF-S1 fibroblasts (Fig. 7d). Due to this strong effect of NOTCH inhibition on CAF-S4 contractility, we next analyzed in-depth the effect of NOTCH on CAF-S4. We verified that DAPT had no effect on CAF-S4 viability (Supplementary Fig. 6c). As expected, Blebbistatin and Y27632, potent myosin and ROCK inhibitors used as positive controls, also abolished CAF-S4 contraction (Supplementary Fig. 6d). Using TFM, we confirmed that DAPT treatment also diminished the Strain Energy Density globally exerted by CAF-S4 cells (Fig. 7e), although it did not significantly affect the magnitude of the stress which averages the local traction forces (Fig. 5e). Moreover, the inverted Transwell invasion assay confirmed the role of NOTCH in CAF-S4-mediated BC cell invasion. Indeed, both the maximal distance reached and the magnitude of the stress which averages the local traction forces (Fig. 5f).

**Fig. 5 CAF-S1 promote proliferation and initiate cancer cell epithelial-to-mesenchymal transition.**

- **a** Total number of viable BC cells (DAPI− cells by FAC) in co-culture with CAF-S1 (red) or CAF-S4 (blue relative to control (− gray, without CAF) (n = 6 per BC cell type). **b** Total number of viable BC cells (Resazurin staining) with CAF-S1- or CAF-S4-conditioned medium (CM) relative to control (−, without CM) (n = 9 per BC cell type). **c** Total number of viable CAF-S1 (red) and CAF-S4 (blue) (DAPI− cells by FAC) in co-culture with BC cells relative to control (−, without BC cells) (n ≥ 6 per BC cell type). **d** CAF-S1 and CAF-S4 capacities to attract BC cells (n ≥ 8 per BC cell type). **e** Left, Images of co-staining of Vinculin (red), F-actin (green) and DAPI (blue) in MCF7 (left) or T47D (right) cultured alone, or in presence of CAF-S1 or CAF-S4. Scale bars, 20 μm; inset, 10 μm. Arrows show reduced BC cell cohesion in presence of CAF-S1/S4; asterisks show F-actin interconnections between cells in presence of CAF-S4. Vinculin and F-actin individual staining in Supplementary Fig. 4c. Right, Number of BC cells per tumor area (at least 13 images per condition, n = 3 per BC cell type). **f** Left, Images of E-cadherin (red), F-actin (green) and DAPI (blue) co-staining (top) or of E-cadherin (red) and DAPI (blue) staining (bottom) in BC cells alone or in presence of CAF-S1/S4. Scale bars, 20 μm; inset, 10 μm. Right, Quantification of E-cadherin staining per BC cell area (at least eight images per condition) (n = 2 per BC cell type). In all panels, barplots are mean ± SEM and n number of independent experiments; a. u., arbitrary units. **a**, **b**, **c** Values from Student’s t-test. **b**, **c** Values from paired t-test. **e**, **f** Right: p Values from Student’s t-test (MCF7) and Mann–Whitney test (T47D). At least two CAF-S1 and CAF-S4 pairs tested, except in **e**, **f** for T47D, where one CAF-S1/S4 pair is used. Source data provided in Source Data file, with R scripts used.
CAF-S4 (Supplementary Fig. 6i, j). In conclusion, CAF-S1 fibroblasts promote BC cell migration and EMT initiation in cancer cells in a CXCL12- and TGFβ-dependent manner. In addition, CAF-S4 fibroblasts stimulate cancer cell invasion and motility in 3D by increasing contractility through the NOTCH signaling pathway.

CAF-S4-enriched LNs are associated with poor patient survival. We finally investigate the clinical relevance of our findings. We analyzed the impact of global stroma content and CAF subset enrichment in LN on disease-free survival, i.e. survival without local, regional or distant relapse (Fig. 8a and Table 1). As CAF-S1 are able to initiate EMT, we quantified the percentage of stromal and epithelial compartments in LNs. High content in stroma in LNs was associated with a shorter disease-free survival, with a similar tendency for overall survival ($p = 0.06$ by Log Rank test and $p = 0.07$ by Cox regression model) (Fig. 8a, b and Table 1). Using multivariate Cox regression analysis with additive model, we observed that the stromal content was independent of BC subtype and LN status at diagnosis, two well-established independent prognostic parameters (Table 1). As invaded LNs are mainly composed of CAF-S1 or CAF-S4 (Fig. 2), these results suggested that both CAF subtypes, when present in high quantity in invaded LNs at diagnosis, might be deleterious for patient survival. We found that CAF subsets on their own exhibited a faint but significant prognostic value, in particular when patients were stratified according to their stromal content in LNs (Fig. 8c and Supplementary Table 6). Indeed, among patients who displayed high stromal content in LNs at diagnosis, those enriched in CAF-S4 showed the poorest overall survival, while no difference according to CAF subset was observed in the context of low stromal content (Fig. 8c). Of note, the prognostic value of CAF subset content in LNs is interesting—although faint—as it is not observed in PT3,33. Considering that CAF-S1 and CAF-S4 enhance BC cell invasive capacities and that most BC patients decease from metastases, we next looked at distant metastases developed after diagnosis. In line with our results, patients who developed such metastases were mainly patients with CAF-S4-enriched LNs at diagnosis, with a similar repartition in BC subtype (Fig. 8d). The impact on distant metastatic spread was even stronger in CAF-S4- than in CAF-S1-enriched LN patients, effect detected in all BC subtypes (Fig. 8e). Overall, patients who showed LNs with high stromal content and specific enrichment in CAF-S4 were more prone to develop late distant metastases than any other patients, in particular in liver (Fig. 8f, g). Although metastatic sites vary according to BC subtypes with bone as the...
most common site except in TN BC associated with other niches, such as liver,\(^1,2\), CAF-S4 impact was not associated to a specific BC subtype. Indeed, among all patients displaying late liver metastasis, only one patient (6%) presented HER2 overexpression at diagnosis. Moreover, CAF-S4-enriched metastatic patients were numerous both within Lum and TN BC subtypes (Fig. 8g). We even detected a higher proportion of patients categorized as CAF-S4-enriched with high stromal content in LNs in the Lum BC subtype compared to TN BC patients (60% Lum versus 33% TN). Thus, patients with high-CAF-S4 content in LNs, who

\[\text{Velocity (μm/min)}\]

\[
\begin{array}{cccc}
0.00 & 0.05 & 0.10 & 0.15 & 0.20 \\
\end{array}
\]

\[\text{Relative velocity}\]

\[
\begin{array}{cccc}
\text{CAF-S4} & \text{CAF-S4} & \text{DAPT} & \text{DAPT} \\
\end{array}
\]

\[p = 7 \times 10^{-13}\]

\[p = 0.5\]

\[p = 0.5\]

\[p = 0.001\]

\[p = 0.01\]

\[p = 0.003\]
tended to metastasize into liver, were not enriched in TN BC subtype in the cohort analyzed. The impact of CAF subset on metastatic spread was thus not related to a bias linked to BC subtype. On the whole, these data highlight the role of CAF-S1 and particularly CAF-S4 sub-sets under BC node status and metastases (see Model Fig. 8h).

Discussion

Although the role of CAF in metastatic spread is well established, when analyzed as a global population, we address here the link between CAF subsets and BC cell spread. Axillary metastatic LN exhibit four CAF subsets (CAF-S1 to S4) to be highly enriched in αSMA+ CAF-S1 and CAF-S4 subsets. The abundance of αSMA+: CAF at PT has been associated with poor prognosis in BC. In agreement with our findings, a recent study reported that the global stromal content in metastatic LN provides a prognostic stratification of BC patients. We go a step further by differentiating CAF-S1 and CAF-S4 and by showing that the CAF-S1/CAF-S4 status in LNs exhibits a prognostic value, while this is not the case when considering CAF subset enrichment in PT. This underlines the relevance of our study in BC pathology and indicates that assessing the content in CAF-S1 and CAF-S4 sub-sets in LNs might be helpful for clinical diagnosis.

Both CAF-S1 and CAF-S4 sub-sets exhibit pro-invasive properties yet with distinct modes of action. CAF-S1 enhance tumor cell migration and EMT initiation, while CAF-S4 favor cancer cell invasion and motility in 3D. Although the impact of CAF in metastatic spread has been previously demonstrated, we show here the complementary role of two distinct CAF sub-populations in metastases. We identify a crosstalk between CAF-S1 and cancer cells involving both CXCL12 and TGFβ, consistent with data showing that CAFs secrete CXCL12 and promote cancer cell migration. In addition, we highlight the role of TGFβ, a well-known EMT inducer, and an important player in fibroblast activation. Both CXCL12 and TGFβ expression are found in BC metastatic sites, in particular lung and bone, supporting they are particularly relevant in BC. In contrast to CAF-S1, CAF-S4 are highly contractile and exert a pro-invasive effect on cancer cells. Although mechanisms linking CAF contractility to cancer cell invasion have been described, we unveil here the NOTCH pathway as a key player. NOTCH pathway has been studied in cancer with a strong focus on cancer cells. Our data show that NOTCH exerts a key role in CAF-S4 contractility and its pro-invasive action in BC. Thus, we report specific CAF-S1 and CAF-S4-mediated processes and mechanisms that promote metastatic spread in BC.

The four CAF sub-sets we identified in LN mirror CAF sub-populations in PTs. This is consistent with studies showing that LN stroma mimics PT microenvironment. But, our results provide new insights. Indeed, if the four CAF sub-sets are detected in both PT and LN, their abundance strikingly differs. Compared to PT, LNs are almost exclusively enriched in activated CAF-S1 and CAF-S4. The transcriptomic signatures of CAF subsets from LN validated molecular similarities between CAF-S1 (or CAF-S4) from PT and LN. If few studies addressed CAF transcriptomic profiles in both PTs and LNs from BC patients, a short gene list identified in ref. allows to discriminate LN from PT bulk samples. Interestingly, most of these genes are highly detected in CAF-S4, consistent with their abundance in metastatic LNs. Although CAF origin remains an open question, CAF-S1 and CAF-S4 transcriptomic profiles generated from PT and LN indicate that subset identity is similar between the two tissues. A recent study identified nine stromal populations by single-cell RNAseq in mouse LN, including the established FRCs, FDC and perivascular cells. Although these cells were identified in non-cancerous mouse lymphoid tissues, CAF-S1 are close to CD34+ stromal cells and CAF-S4 to perivascular cells. Our data on CAF-S4 are consistent with previous studies showing the existence of pericyte-like CAF and their potential pericytes-derived origin. Thus, CAF subsets in invaded LNs could derive from different resident mesenchymal cells. In conclusion, by combining the study of BC patient samples and performing functional assays on human CAF subset cultures, we demonstrate that CAF subsets exhibit distinct functions in metastatic spread. Considering the deleterious effect of metastases on BC patients' survival, our data might strengthen the interest in assessing CAF subsets content in LN for clinical diagnosis and in using anti-TGFβ and/or anti-NOTCH therapies in BC.

Methods

Cohorts of BC patients. The projects developed here are based on surgical residues, available after histopathological analyses, and not required for diagnosis. There is no interference with clinical practice. Analysis of PT and metastatic LN samples was performed according to the relevant national law on the protection of people taking part in biomedical research. All patients included in our study were informed by their referring oncologist that their biological samples could be used for research purposes and they gave their verbal informed consent. In case of patient refusal, that could be either orally expressed or written, residual tumor samples were not included in our study. Human experimental procedures were approved by the Institutional Review Board and Ethics committee (CRI) of the Institut Curie Hospital group (approval 12 February 2014) and CNIL (Commission Nationale de l’informatique et des Libertés) (No. approval: 1674356 delivered 30 March 2013). HER2-amplified carcinomas have been defined according to ERBB2 immunostaining using ASCO’s guideline. Luminal (Lum) tumors were defined by positive immunostaining for ER (estrogen receptor) and/or PR (progesterone receptor). The cut-off used to define hormone receptor positivity was 10% of stained cells. Ki67 (proliferation) score further distinguishes Lum A from Lum B tumors (below 15%: Lum A, above: Lum B). TN immunophenotype was defined as...
follows: ER−PR− ERBB2− with the expression of at least one of the following markers: KRT5/6+ or EGF-R+.

PT (up to 75) and LN samples (up to 124) from retrospective cohorts of BC patients suffering from invasive BC cancers with at least one metastatic LN at diagnosis, have been collected as part of routine standard of care and analyzed by IHC. Retrospective cohorts include Lum, HER2 and TN subtypes. IHC were performed on residual surgery samples prior to any treatment (i.e. prior to radiation, hormonal or chemo-therapy). Overall survival and disease-free survival have been defined as followed: overall survival is defined as the period of time going...
from diagnosis to death of the patient: disease-free survival is the period of survival without local, regional or distant relapse, meaning the period of time going from diagnosis to the first event occurring among local, regional or distant relapse. Clinical features of prospective and retrospective cohorts are listed in Tables S1 and S2.

Flow cytometry on BC PT and LN samples. Fresh human BC PT and LN samples were collected directly from the operating room, after surgical specimen’s macroscopic examination and selection of areas of interest by a pathologist. Samples were cut into small pieces (around 1 mm³) and digested in CO₂-independent medium (Gibco #18045-054) supplemented with 150 μg/ml liberase (Roche #05401020001) and DNase I (Roche #11284932001) during 40 min at 37°C with shaking (180 rpm). Cells were then filtered through a 40-μm cell strainer (Fisher Scientific #223635447) and resuspended in PBS solution (PBS, Gibco #14190; EDTA 2 mM, Gibco #15575; Human Serum 1%, BioWest #S1490-100) at a final concentration between 5 × 10⁵ and 10⁶ cells in 50 μl. Cells in suspension were then stained with an antibody mix containing anti-EPCAM-BV605 (1:50; Biolegend, #324224), anti-CD146-PECy7 (1:100; Biolegend, #301318), anti-CD45–APC-Cy7 (1:20, BD Biosciences, #BD-557833), anti-CD235a-PerCP/Cy5.5 (1:50, Biolegend, #349109), anti-CD29-Alexa Fluor 700 (1:100, Biolegend, #303020), anti-FAP-AF488 (1:50, Biolegend, #357006) for surface staining and anti-SMA-Alexa Fluor 594 (1:25, R&D Systems, #IC14207-025) for intracellular staining. All antibodies except FAP were purchased already conjugated with fluorescent dyes. Anti-FAP antibody was conjugated with fluorescent dye Zenon APC Mouse IgG1 labeling kit (1:100, Thermo Fisher Scientific, #Z25051). Isotype control antibodies for each marker used were: iso-anti-CD29 (1:100, Biolegend, #400144), iso-anti-FAP (primary antibody, 1:200, R&D Systems, #MA0002), iso-anti-PDGFRβ (1:20, Biolegend, #000114), iso-anti-PDPN (1:125, Biolegend, #400525) and iso-anti-SMA (1:125, R&D Systems, #IC0037). For intracellular staining, cells were incubated with violet LIVE/DEAD dye (1:1000, Thermo Fisher Scientific, #L34955) for 20 min at 37°C (in PBS, Gibco #14190) to dead cells, and then fixed in 4% paraformaldehyde (PFA, Electron Microscopy Sciences, #15710) overnight at 4°C. After a rapid washing step in PBS solution, cells were incubated for 45 min at RT with the antibody mix in PBS supplemented with 0.1% saponin (Sigma-Aldrich, #S9090). For surface staining, cell suspensions were stained immediately after dissociation of BC PT and LN samples during 15 min at RT with the antibody mix in PBS solution. 2.5 μg/ml DAPI (Thermo Fisher scientific, #D13063) added just before flow cytometry analysis. For primary CAF-S1 and CAF-S4 culture characterization, cells at 80% confluence were collected after trypsin treatment, resuspended in PBS + 5% FCS, and centrifugation of 10⁵ cells in 50 μl and intracellularly stained with the above-described protocol.

In both conditions (surface and intracellular staining), signals were acquired on the LSRFortessa® analyzer (BD biosciences) for flow cytometry analysis. At least 5 × 10⁶ events were recorded. Compensations were performed using single staining on anti-mouse IgG and negative control beads (BD biosciences, #525843) for each antibody except anti-PDPN, on ArC reactive beads (Molecular probes #A10346) for live/dead staining and on AbC Total Antibody Compensation Beads (Thermo Fisher scientific, #A10513) for anti-PDPN.

Data analysis was performed using FlowJo version 10.7.0r2. Cells were first gated based on their viability using live/dead or side scatter (SSC-A) and side (SSC-B) scatter (measuring cell size and granularity, respectively) to exclude debris. Dead cells were excluded based on their positive staining for Live/Dead (fixed conditions) or DAPI (surface staining). Single cells were next selected based on SSC-A versus SSC-W parameters. Gating included EPCAM−, CD45−, CD31−, CD235a− cells, to remove epithelial (EPCAM−, CD45−, CD31−, CD235a−) cells from the negative fraction were next examined using the 5 CAF markers FAP, CD29, aSMA, PDGFRβ and PDPN.

ICH analyses. BC PT and LN cases were selected by Pathologists (see Supplementary Table 2) and serial sections of paraffin-embedded tissues (PT: Tissue-Micro-Arrays (TMA) using two cores (1 mm diameter, selected as representative from whole-tumor tissue sections by Pathologists) of tumor per case and cut in slices of 3 μm; LN: whole sections of 3-μm thickness) were stained on the Lab Vision Autostainer (Thermo Fisher Scientific). Dewaxing and antigen retrieval on slides prone to immunohistochemical staining were performed with Envision Flex Target Retrieval Solution (high- or low-pH, as required—see below—Dako, #K800441 or #K800521) on the Lab Vision PT Module (Thermo Fisher Scientific). Antigen detection was done using the Envision Flex/XRP (Dako, #K8006) for CD29, aSMA and PDGFRβ or the streptavidin-biotin peroxidase protocol (Vectastain ABC kit; Vector Labs #PK-6101/6102/6104) plus detection with 3,3’-diaminobenzidine for 5 min (DAB, Dako, #K3468) for FSP1, EPCAM (on LN) and FAP. The following antibodies and respective conditions were used: anti-FAP (pH = 6, 1 h at 1:200, Vitaxins #MAB601001), anti-CD29 (pH = 6, 1 h at 1:100, Abcam #ab16176), anti-FSP1 (pH = 6, 1 h at 1:50, Abcam Ab27997), anti-aSMA (pH = 6, 30 min at 1:200, DAKO, #M0851), anti-PDGFRβ (pH = 9, 1 h at 1:100, Abcam #ab23570), anti-EPCAM (pH = 6, 1 h at 1:200, DAKO #M0804), anti-E-cadherin (pH = 9, 1 h at 1:100, Invitrogen #180223). Counterstaining was performed with Mayer hematoxylin freshly prepared (Dako, #S3309), Tissue sections were then submitted to serial gradients of xylene and mounted with coverslip in an automatic deparaffinising unit (Tissue-Tek DriCore). In each sample, CAF marker staining was evaluated as a histological score (H-score), defined by staining intensity (ranging from 0 to 4) multiplied by the percentage of stained fibroblasts. For LNs, only fibroblasts in metastatic zones—assessed by EPCAM staining—were evaluated. For each PT and LN sample, CAF enrichment in specific compartments was evaluated using an algorithm developed by the team, which takes as input CAF markers H-scores, as described in refs. 32,33. In brief, the thresholds were first defined in a learning dataset on the distribution (1st quartile, median and 3rd quartile) of each marker using FACS data. Thresholds were then transposed to IHC data. For each section, the percentages of stroma and epithelial morphology assessment and EPCAM staining performed by an independent researcher (EPCAM/E-cadherin staining of epithelial cells). Quantifications gave very consistent results whatever the person (pathologist versus researcher), the method (manually versus automated) or the staining (EPCAM/E-cadherin), thereby arguing that evaluation of stromal and epithelial content was accurate. In each section, percentage of stroma was evaluated as the fibroblastic area multiplied by 100 divided by the total sample area.

Design of a decision tree for CAF subset prediction. CAF identity was determined by using an algorithm developed by the team (shown in Fig. 2d), which takes as input histological scores of CAF markers. The thresholds were first defined, in a learning dataset, on the distribution (1st quartile, median and 3rd quartile) of each marker using FACS data. Thresholds were then transposed to IHC data.

CAF subsets and epithelial cells at cellular level in situ in invaded LNs. IHC staining from consecutive sections were scanned on Philips Ultra Fast Scanner; 5x images of CAF markers including FAP, CD29, FSP1 and SMA (markers used in the decision tree algorithm, see also Fig. 3d) as well as EPCAM staining from the same
Table 1 Univariate and multivariate Cox regression analyses for progression-free survival and overall survival in N+ breast cancer patients-199 cases (LN cohort).

| Variable | LN | N | Multivariate | Hazard ratio | 95% CI | p Value |
|----------|----|---|-------------|--------------|--------|---------|
| LN fraction | 1 | 2.4 | 1 | 0.02 | 2.3 | 0.9 |
| CAF-S1 | 1 | 1.1 | 1 | 0.5 | 2.1 | 0.8 |
| CAF-S4 | 1 | 1.1 | 1 | 0.5 | 2.1 | 0.8 |

Gene expression profiling by RNAseq. CAF-S1, CAF-S4 and EPCAM+ cells (from PTs and LNs of five patients, n = 28 samples in total, see Supplementary Table 1) were sorted after surface staining (see 'Flow Cytometry on BC PT and LN samples') on FACSAria (BD biosciences), directly into RNase-free tubes (Thermo Fisher Scientific #AM12450). At least 100 cells were collected per cell population. Total RNA was extracted with Single Cell RNA Purification kit (Norgen Biotech #51800) following manufacturer's instructions. RNA integrity and quality were checked with Agilent RNA 6000 Pico Kit (Agilent, #5067-1513) and appara tus. cDNA synthesis and amplification were synthesized using SMART-seq2 Ultra Low Input RNA Kit for Sequencing (Clontech #634892). cDNA quality was checked on Agilent 2100 Bioanalyzer using Agilent High Sensitivity DNA kit (Agilent #5067-4626) and quantified using the Qubit dsDNA HS Assay kit (Life Technologies, #Q32854). cDNA libraries were prepared using Nextera XT preparation kit (Illumina #FC131-10). Samples were sequenced on HiSeq 2500 (Illumina) rapid run flow cells with an average sequencing depth of 34.5 million paired-end reads. Read length was 100 bp. Reads were mapped on reference human genome (hg19/GRCh37 from UCSC genome release) using TopHat 2.0.6 algorithm with the following parameters: global alignment, no mismatch in seed alignment (of size 22), three mismatches in read length. Quality control was performed using FastQC software and duplicates were removed using Samtools rmdup. Gene expression quantification was performed using HTSeq-count and featureCounts (implemented in Bioconductor R package Rsubread #1.18.0). Only genes with at least one read in 5% of all samples were kept for further analyses. Normalization was performed with DESeq2 R package (1.8.2) method. Analysis also included unsupervised analyses such as PCAs and HC, as well as paired differential expression analyses (done with DESeq2 biocomparator package). Biological interpretation of the identified genes was done by computational functional analyses on Metascape (http://metascape.org), which is based on several bioinformatics resources (Gene Ontology, KEGG, Reactome). RNAseq data have been deposited on EGA website (EGA number: EGAS00001003238).

Cell culture. The human BC cell lines MCF-7(-GFP), T47D(-GFP) and MDA-MB-231-GFP were propagated in DMEM (GE, #SH30243.01) supplemented with 10% fetal bovine serum (FBS) (Biosera, #1003/500), penicillin (100 U/ml) and streptomycin (100 µg/ml) (Gibco #15140122). Each cell line identity was verified by Short Tandem Repeat (STR) DNA profiling (Promega #B9510).

CAF-S1 and CAF-S4 primary cultures were established from PT or LN human samples. All along the duration of the project, we isolated 22 CAF-S1 and 22 CAF-S4 paired primary cell lines (referred to as CAF pairs because isolated from the same patient) using FACS-sorted cells full-filling marker criteria shown in Fig. 1. Among them, 64% were from PTs and 36% from LNs. In agreement with the strong transcriptomic analogy of CAF subsets from PT or LN (shown in Fig. 3), we did not observe any difference in functional assays when we performed experiments with CAF-S1 or CAF-S4 isolated either from PT or from LN. Thus, we observed no effect of the site of isolation (PT or LN) on CAF-S1/CAF-S4 properties. Transcriptomic signatures and in vitro properties were thus equivalent whatever the site of CAF initial localization (PT or LN). Although CAF-S1 were quite easily expanded in vitro, CAF-S4 were difficult to maintain. As they exhibit a pericyte-like signature, we adapted the culture conditions to expand and maintain both CAF-S1 and CAF-S4 in similar and comparable conditions. BC samples were processed and sorted after surface staining as described above (see 'Flow Cytometry on BC PT and LN samples') by cell sorting and gating strategy using FACS shown in Fig. 1. Purified CAF-S1 and CAF-S4 were then plated in 96-well plates and expanded in a pericyte medium (Sciencell, #1201) in a humidified, 1.5% O2 and 5% CO2 incubator. Doubling time was assessed during this expansion phase for six CAF pairs over five passages. All cultures were systematically validated by qRT-PCR between passages 3 and 6 using genes upregulated in either CAF-S1 or CAF-S4 (including SFRP2, HMCN1, TINAGLI, PDE3A). Other types of validation were done, such as RNAseq on four CAF-S1 and CAF-S4 pairs and FACS analyses (see 'Flow Cytometry on BC PT and LN samples') to assess CAF marker protein levels (as shown in Supplementary Fig. 3). For RNAseq experiment, cells at 80% confluency were collected after trypan treatment, total RNA was extracted (Qiagen #217004) and DNase-treated (Life technologies #EN0525). mRNAs were retrotranscribed, amplified and cDNA libraries were obtained with TruSeq Stranded.
mRNA library prep (Illumina #RS-122-2101). Samples were then sequenced and analyzed as PT and LN fresh CAF samples (see Gene expression profiling by RNAseq). We defined as the antibody that the position corresponds to the cells were taken from both sides of the cell-free zone border — absolute because cells were taken from both sides of the cell-free zone border — calculated for each successive pair of time points for each cell. Means over all the global tracking are shown. Persistence of cell trajectory, defined as the ratio p where l is the contour length of the total trajectory and d the direct end-to-end distance was also calculated per cell over the same range. Two different CAF-S1 and CAF-S4 pairs were tested in two independent experiments.

CAF spheroids and invasion assay. For spheroid invasion assays, CAFs were trypsinized and resuspended at a concentration of 4 × 10^5 cells/ml. A total of 20 μl drops were pipetted down on a 10-cm dish lid. Lids were then gently put back on the plates filled with medium to humidify the system. After 4 days, CAF spheroids were detached from the hanging drops in a gently transferred to 24-well plates containing collagen solution drops (Conring, #354249) in the center of 6-well plates. At least six spheroids per CAF were embedded in collagen and further analyzed. After polymerization, 0.5% FBS medium was added to the wells. Spheroids were imaged 3 days after inoculation on a Nikon Eclipse TE300 microscope with a ×4 objective mounted on Nikon DS-Fi1c and imaged with Fiji software as an invasion index defined as the total area covered by CAFs (encompassing single cells escaped from the spheroid and the spheroid) divided by the area of the spheroid core. Three CAF-S1 and CAF-S4 pairs were tested in three independent experiments. For time-lapse imaging, time-lapse videos were immediately acquired after collagen polymerization using inversion microscopy and Leica video microscopes, equipped with motorized stage, 37 °C incubator and CO2 controller (Leica DMi8 with a Retiga R6 camera and illumination by Lumencor SOLA SE 365). The automated imaging system was controlled by the software Metamorph (Universal Imaging). Transmission images were acquired every 15 min until 42 h with an objective to encompass the whole spheroid. Videos were mounted with eight frames per second.

Collagen contraction assay. To assess force-mediated collagen contraction, 3 × 10^4 CAFs were embedded in 100 μl of a 2.5-mg/ml collagen solution (Conring, #354249) and seeded in triplicates in 96-well plates. After polymerization, culture medium was added on top of the gels and in the surrounding empty wells to limit evaporation. To assess DAPT (Sigma, #D9942), BLEBBISTATIN (Sigma, #B9650) and Y27632 (STEMCELL Technologies, #72302) impact on collagen contraction, 10 μM of each drug was added to the medium, DMSO as control. Gel contraction was assessed 4 days after plating: plates were scanned (EPSON Perfection V700 Photo scanner) and quantification was performed on Fiji software. Percentage of contraction was calculated using the formula:

\[
\text{Percentage of contraction} = \frac{A_{\text{t=0}} - A_{\text{t=end}}}{A_{\text{t=0}}} \times 100
\]

where

\[
A_{\text{t=0}} = \text{the traction stress at time } t = 0
\]

\[
A_{\text{t=end}} = \text{the traction stress at the end of the experiment}
\]

\[
\text{SE} = \frac{\text{the average stress per cell}}{\text{the area}}
\]

\[
\text{the mean traction stress amplitude which corresponds to the average stress per cell. SE}
\]

\[
= \frac{1}{n} \sum_{i=1}^{n} T_i
\]

\[
\text{where } T_i = \text{the traction stress field and } d_i = \text{the displacement field.}
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Collagen evaluation by second harmonic generation. CAFs were seeded in home-made holes in 30 mm^2 tissue culture plates as described in ref. 22. Three holes of 3-4 mm diameter were drilled in a plate and widened around the edges using a scalpel. The bottom of the dish was covered with epoxy (Loctite), and 20 × 20 mm squares were glued to the dish overnight at RT. The day before the experiment, dishes were silanized with 3-aminopropyl-trimethoxysilane (Sigma-
and immediately used for experiments. For CAF viability assessment, 7 × 10³ CAFs were seeded in triplicates in 96-well plate wells with the different drugs at 10⁻⁶ M DAPT or DMSO was added on both sides of the insert. For control, inserts were first labeled with fluorescent dye Zenon APC Mouse #C34573, for CAF-S4: CellTrace Violet, #C34571) according to manufacturer’s protocol; 2 × 10⁵ cells were then embedded in 100 µl of a 2.5-mg/ml collagen solution (Corning, #354249) and seeded inside 24-well cell culture inserts (Corning, 9 µm pore size, #3422). CAF-free collagen was used as control. After polymeric inserts were put on plates (for CAF-S1), at least four cultures were performed in duplicates. After 24 h, inserts were transferred into new 24-well plates containing 0.5% FBS and further incubated for 6 days. All conditions were done in duplicates. After 7 days of invasion, inserts were gently washed in PBS, fixed in 4% PFA for 15 min, washed again and mounted with Mounting medium (Aqua-Poly/Mount #18606) on glass-bottom dishes (FisherScientific #15199112). Images were then acquired on an inverted Laser Scanning Confocal Microscope with Spectral Detection and Multi-photon Laser (LSM880NLO/MaiTro/Coherent Inc.) using a 25×/1.0 NA water immersion objective. The microscope was set up so that membrane pores were in focus and highest Z level where no MDA-MB-231-GFP cells could be detected anymore). Fluorescence channels were recorded using the excitation wavelengths 405 nm (for CAF-S4), 488 nm (for BC cells) and 590 nm (for CAF-S1 and CAF-S4) in a custom-made Inverted Transwell assay. 

For DAPT (Sigma, #D5942) treatment, when CAF-S4 were used, they were labeled and included in collagen as described above. MDA-MB-231-GFP cells were seeded as described above (below CAF-S4-free collagen for control experiments or CAF-S4-embedded collagen). 10 µm DAPT or DMSO was added on both sides of the inserts and media were changed at day 3. All conditions were done in duplicates. Inserts were fixed, acquisition and analysis were performed as described above with the following Imaris parameters: surface detail 2 µm; threshold absolute intensity 2.00; option “enable splitting of touching objects” was selected; seed points diameter 15 µm; filtered on voxels above 350. 5 independent control experiments were performed and 6 independent experiments including 5 different CAF-S4 cell lines were performed to assess DAPT effect on CAF-S4-induced invasion of BC cells.

Tumor cell density, E-cadherin in BC cells, F-actin and Vinculin in CAF. For tumor cell density and assessment of staining in CAF subsets, quantifications were performed in two steps on Fiji software. (1) Masks were manually drawn on each merged image to either keep tumor cell zones of CAF zones according to the analyses. Masks were then applied to the corresponding single channel (Dapi, E-cadherin, F-actin or Vinculin depending on the analyses) images. (2) For BC cell density, evaluation of cell number was performed by setting a threshold filter (0, 20) on Dapi channel (raw images opened and auto-scaled with Bio-formats plug-in, saved in jpg format) followed by a mask conversion, segmentation and then particles were analyzed (size 2000-25,000; circularity > 0.2). Cell number was then divided by the tumor zone area. For F-actin or Vinculin signal extraction, as CAFs were negative for E-cadherin, step (1) was performed and step (2) was automatically implemented (as shown in Figs. 5, 7 was defined on ImageJ), and next divided by tumor cell surface. Images were thresholded (single threshold per experiment ranging from 50 to 225 applied on raw images opened and auto-scaled with Bio-formats plug-in, saved in jpg format), converted to masks, integrated density was retrieved and then divided by tumor zone area.

Inverted Transwell assay. For inverted Transwell assays, CAF were first labeled with fluorescent dyes (Thermo Fisher Scientific, for CAF-S1: CellTrace Yellow #C34573, for CAF-S4: CellTrace Violet, #C34571) according to manufacturer’s protocol; 2 × 10⁵ cells were then embedded in 100 µl of a 2.5-mg/ml collagen solution (Corning, #354249) and seeded inside 24-well cell culture inserts (Corning, 9 µm pore size, #3422). CAF-free collagen was used as control. After polymeric inserts were put on plates (for CAF-S1), at least four cultures were performed in duplicates. After 24 h, inserts were gently transferred into new 24-well plates containing 0.5% FBS and further incubated for 6 days. All conditions were done in duplicates. After 7 days of invasion, inserts were gently washed in PBS, fixed in 4% PFA for 15 min, washed again and mounted with Mounting medium (Aqua-Poly/Mount #18606) on glass-bottom dishes (FisherScientific #15199112). Images were then acquired on an inverted Laser Scanning Confocal Microscope with Spectral Detection and Multi-photon Laser (LSM880NLO/MaiTro/Coherent Inc.) using a 25×/1.0 NA water immersion objective. The microscope was set up so that membrane pores were in focus and highest Z level where no MDA-MB-231-GFP cells could be detected anymore). Fluorescence channels were recorded using the excitation wavelengths 405 nm (for CAF-S4), 488 nm (for BC cells) and 590 nm (for CAF-S1 and CAF-S4) in a custom-made Inverted Transwell assay. 

For DAPT (Sigma, #D5942) treatment, when CAF-S4 were used, they were labeled and included in collagen as described above. MDA-MB-231-GFP cells were seeded as described above (below CAF-S4-free collagen for control experiments or CAF-S4-embedded collagen). 10 µm DAPT or DMSO was added on both sides of the inserts and media were changed at day 3. All conditions were done in duplicates. Inserts were fixed, acquisition and analysis were performed as described above with the following Imaris parameters: surface detail 2 µm; threshold absolute intensity 2.00; option “enable splitting of touching objects” was selected; seed points diameter 15 µm; filtered on voxels above 350. 5 independent control experiments were performed and 6 independent experiments including 5 different CAF-S4 cell lines were performed to assess DAPT effect on CAF-S4-induced invasion of BC cells.

Tumor-on-chip assay. The devices were made of PDMS (polydimethylsiloxane), a biocompatible, gas-permeable and transparent silicone rubber. The devices were

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micro-fabricated at the Institut Pierre-Gilles de Gennes pour la Microfluidique (IPGG, Paris) using standard soft-lithography or advanced micro-milling for the mold fabrication and soft-lithography methods to prepare the chips. The micro-fluidic device were sterilized in UV-oven (365 nm wavelength) for 30 min, coated with 10 µg/ml human fibronectin (Sigma, #F0895) overnight at 37 °C, 5% CO2, rinsed with PBS and then dried for at least 5 h in a sterile room. Cells were embedded in 2.3 mg/ml collagen hydrogels (PureCol, Advanced Matrix, #5005 diluted in MEM medium (Sigma Aldrich, #M0275) containing 0.28% NaHCO3 (pH = 8) to adjust the final pH to 7). MCF7-GFP or T47D-GFP targeted density was 1.2 × 106 cells/ml, CAF density was 3 × 105 cells/ml. When DAPT (Sigma, #D5942) was used, CAF were pre-treated overnight before inclusion in collagen and a final concentration of 10 μM was added to the medium. We verified that DMSO did not interfere with BC cell velocity in the system. Time-lapse videos were then acquired with an inverted motorized Leica video-microscopes, equipped with motorized stage, 37 °C incubator and CO2 controller (Leica DMi8 with a Retiga R6 camera and illumination by Lumenor SOLA SE 365) in a 20% O2 environment. The automated imaging system was controlled by the software Metamorph (Universal Imaging). Transmission and fluorescent (GFP) images were acquired every 2 h over 30 h and at least four positions per condition were recorded with a 10x objective. For data analyses, Fiji software was used. In each position, cells were manually tracked over all frames (Manual tracking’ plugin, at least 65 cells per condition). Cell velocity (distance of migration divided by the interval time) was calculated for each successive pair of time points. Means per cell over the global tracking are shown. For CAF-S1 versus CAF-S4 experiments, for each BC cell line: two independent experiments were performed, with three different CAF-S1 and CAF-S4 pairs. For DAPT experiments, for each BC cell lines: three independent experiments were performed, including two different CAF-S4 cell lines.

Cytokine antibody-pair-based assay. Cytokine antibody-pair-based assays were performed using human cytokine array kit (R&D Systems, #ARY008) according to the manufacturer’s protocol. In brief, 80,000 CAF-S1 or CAF-S4 fibroblasts were cultured alone or in presence of 80,000 BC cells (MCF7 or T47D) in 1 ml of DMEM medium in 12-well plates at 1.5% O2 during 48 h. Culture supernatants were next collected (CM) and centrifuged at 13,000 rpm during 10 min to eliminate debris. Array membranes, previously spotted with capture antibodies by the manufacturer, were incubated with 0.5 ml of CM overnight at 4 °C. Membranes were then washed three times with 50 ml of washing buffer at RT, and incubated with a streptavidin-horseradish peroxidase-coupled antibody (1:2000) for 30 min at RT and revealed using Chemi-Reagent Mix. The Immunoblot images were captured and visualized using the ChemiDoc Imaging System (Bio-Rad) and intensity of each spot in the captured images was measured using ImageJ software. Two spots per membrane were used to measure the levels of each cytokine. Intensity mean of the two spots—revealing each cytokine level per experiment—was reported to the intensity mean of three internal controls spotted on the membrane and assessed by the following ratio: specific intensity mean / control intensity mean, to take into consideration variation in hybridization efficiency in the different experiments.

Software. Software used for the above analyses are: R (https://cran.r-project.org, R versions 3.3.1, 3.4.0 and 3.5.0), Fiji (ImageJ v2.0.0–rc-19/1.49m, v2.0.0–rc-65/1.52b), IMARIS (v8.4.1), Matlab.

Statistical analyses. Data graphical representation and statistical analyses were done using R or GraphPad Prism environment. Barplots represent means ± standard errors of the mean (SEM). Boxplots: box limits indicate the inter-quartile range (IQR) (25th to 75th percentiles), with a center line indicating the median. Whiskers show value ranges up to 1.5 × IQR above the 75th or below the 25th percentiles, while outliers beyond those ranges shown as individual points. The number of independent experiments is specified in each figure legend, with at least three independent experiments, unless otherwise specified. The statistical tests used are in agreement with the data distribution: normality was first checked using the Shapiro–Wilk test and parametric or non-parametric two-tailed tests were applied according to normality test results (indicated in figure legend). Survival analyses were carried out with survival R package by Kaplan–Meier analyses were carried out with survival R package by Kaplan–Meier curves (p-values from Log-rank test) and by univariate and multivariate Cox regressions using additive hazards regression models. Stratification of patients for Kaplan–Meier analyses were performed using median value of the stromal content distributions of CAF-S1 or CAF-S4 enrichments (defined by our algorithm, see ‘IH’ analyses). Differences were considered statistically significant when p ≤ 0.05.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
RNAseq data from EP-CAM+, CAF-S1 and CAF-S4 sorted from BC samples, without or with culture, generated in this study are available on European Genome-Phenome Archive platform (https://ega-archive.org) under accession number: EGA89000/103238.

A source data file corresponding to all panels of the figures and supplementary figures that support the findings of this study is provided with the paper.

Code availability
R scripts used to generate panels of the figures and supplementary figures are provided within the data source file of the paper, available at https://doi.org/10.6084/m9. figshare.9924275.

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**Additional information**

*Supplementary information* is available for this paper at https://doi.org/10.1038/s41467-019-14134-w.

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