Cancer-associated fibroblasts lead tumor invasion through integrin-β3–dependent fibronectin assembly

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Cancer-associated fibroblasts (CAFs) are the most abundant cells of the tumor stroma. Their capacity to contract the matrix and induce invasion of cancer cells has been well documented. However, it is not clear whether CAFs remodel the matrix by other means, such as degradation, matrix deposition, or stiffening. We now show that CAFs assemble fibronectin (FN) and trigger invasion mainly via integrin-αvβ3. In the absence of FN, contractility of the matrix by CAFs is preserved, but their ability to induce invasion is abrogated. When degradation is impaired, CAFs retain the capacity to induce invasion in an FN-dependent manner. The level of expression of integrins α and β3 and the amount of assembled FN are directly proportional to the invasion induced by fibroblast populations. Our results highlight FN assembly and integrin-αvβ3 expression as new hallmarks of CAFs that promote tumor invasion.

Introduction

Metastasis formation is a complex multistep process that requires cancer cells to move from their tissue of origin to neighboring and distant organs (Joyce and Pollard, 2009; Wirtz et al., 2011). Through their journey, cancer cells cooperate with the tumor microenvironment, which promotes their invasion (Mueller and Fusenig, 2004; Joyce and Pollard, 2009). Cancer-associated fibroblasts (CAFs) are the most abundant cells in the tumor microenvironment (Kalluri and Zeisberg, 2006). They are a unique cell population, as they can modulate cancer cell invasion directly by secreting proinvasive stimuli and indirectly by remodeling the matrix (Attieh and Vignjevic, 2016).

During the progression of carcinoma, after breach of the basement membrane, cancer cells reach the tumor stroma, encountering CAFs and the ECM. At this stage, the influence of CAFs on tumor invasion is still debated (Kalluri, 2016): it has been shown that depleting CAFs from the stroma induces tumor invasion (Özdemir et al., 2014; Rhim et al., 2014), but most studies agree that an enrichment in CAFs stimulates cancer cell invasion (De Wever et al., 2004; Orimo et al., 2005, Gaggioli et al., 2007; Goetz et al., 2011). There is also disagreement concerning the mechanism by which CAFs act: Do they enhance the invasive capacity of cancer cells through diffusible molecules (De Wever et al., 2004; Orimo et al., 2005)? Or is their physical presence required to contract and align the matrix (Gaggioli et al., 2007; Goetz et al., 2011; Sanz-Moreno et al., 2011; Calvo et al., 2013), facilitating cancer cell invasion (Riching et al., 2014)?

Most new studies highlight the importance of contractility in CAFs in stimulating invasion. However, the ability of CAFs to remodel the matrix by other mechanisms (degradation, stiffening, or deposition of new ECM) and the interdependence between those mechanisms have been poorly studied. For example, highly contractile cells are characterized by stable and long-lived fibrillar adhesions that deposit and assemble new ECMs (Zaidel-Bar et al., 2007). Therefore, ECM deposition by CAFs is a direct consequence of their contractility. The tumor stroma is known to be enriched in matrix proteins like fibronectin (FN) and tenascin C that favor tumor progression (De Wever et al., 2004; Oudin et al., 2016), but it is still not known which of the two functions, contractility or matrix deposition, is responsible for cancer cell invasion.

Here, we investigate how CAFs induce invasion of cancer cells through the ECM. Using a combination of pharmacological and genetic perturbations, we modulated the ability of CAFs to contract, deposit, and degrade the matrix. We found that FN assembly by CAFs via integrin-αvβ3 is critical to stimulate cancer cell invasion.

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Results and discussion

The physical presence of CAFs in the matrix is required to induce cancer cell invasion

To investigate the role of CAFs in cancer invasion, we isolated CAFs and noncancer-associated fibroblasts (NAFs) from the tumor and the neighboring healthy tissue of the colon of patients, respectively, ending with a couple of NAFs and CAFs per patient. We characterized all cell populations using markers of activated fibroblasts (see Materials and methods; Fig. S1 A). In all patients, CAFs and NAFs expressed α smooth muscle actin (α-SMA), fibroblast activation protein (FAP), and PDGF receptor β (PDGFR-β; Fig. S1 A), indicating that even though they were isolated from seemingly healthy tissue, NAFs exhibit features of activated fibroblasts.

To assess the role of CAFs and NAFs in cancer cell invasion of the ECM, we embedded spheroids of CT26 cancer cells in a collagen I matrix either alone or together with CAFs or NAFs (Fig. 1 A). This 3D model recapitulates the scenario of a tumor mass invading the stroma. The invasion capacity of cancer cells was quantified using a custom analysis software 3 d after embedding (see Materials and methods; Fig. 1 B). CT26 is an invasive cancer cell line (Geraldo et al., 2013), and in this assay, cells invaded the collagen matrix even when cultured alone (Fig. 1, A and C). However, in the presence of fibroblasts, the invasion of cancer cells was further enhanced, as previously shown for other noninvasive cancer cell lines (Fig. 1, A and C; Gaggioli et al., 2007; Goetz et al., 2011; Labernadie et al., 2017). CAFs were also more potent in increasing invasion compared with their paired NAFs (Fig. 1, A and C). To validate that this phenotype was not caused by an increased attraction of CAFs compared with NAFs by the tumor, we quantified the number of fibroblasts and their distance from the spheroid. Both CAFs and NAFs accumulated at 50–120 µm from the spheroid edge, and they were found in similar numbers (Fig. 1 D). This suggests that the increased invasion index in the presence of CAFs is most likely caused by an increased potential to secrete proinvasive diffusible molecules or to remodel the matrix.

We next investigated whether CAFs have to be present in collagen gels to stimulate invasion of cancer cells or whether their diffusible molecules are sufficient. In one of the conditions, we added CAF-conditioned media (CAFcm) to spheroids embedded in collagen. Alternatively, we cultured CAFs in the distant presence of cancer cell spheroids (CAF’s diffusible molecules [CAFdm]); in the latter condition, CAFs were not present in the matrix to remodel it, but the secretome cross talk of both cell types was maintained (Fig. 1 E, schemes). In both cases, cancer cells invaded collagen gels to a similar extent as in control conditions (Fig. 1 E), indicating that the physical presence of CAFs in the matrix is necessary to increase cancer cell invasion.

These data show that CAFs induce more cancer cell invasion compared with their paired NAFs and that diffusible molecules of CAFs are not sufficient. Interestingly, the overall ability of fibroblasts to induce cancer cell invasion did not correlate with the expression levels of commonly used CAF markers (Fig. S1 B).

FN deposition by CAFs induces cancer cell invasion

The necessity of CAFs to be physically present in the matrix to induce invasion points toward their role in matrix remodeling. Although NAFs were embedded into the ECM, they did not induce cancer cell invasion. These findings indicate that CAFs, and not NAFs, can remodel the matrix to induce invasion. Proteomic data analysis of two fibroblast couples from colon cancer patients show enrichment in FN in the secretome and proteome of CAFs compared with their paired NAFs (ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD003670). In addition, FN is known to be enriched in the tumor microenvironment and is a proinvasive ECM protein (Wolanska and Morgan, 2015; Oudin et al., 2016). FN could thus be deposited by CAFs to promote cancer cell invasion. To test this hypothesis, we inhibited FN expression in CAFs using small interfering RNA (Fig. S1 C). Depletion of FN in CAFs from all patients abrogated their ability to stimulate the invasion of cancer cells, indicating that FN is necessary for CAFs to induce cancer cell invasion (Fig. 2 A and Fig. S1 D).

This result was surprising, as it has been shown that CAFs mainly stimulate cancer cell invasion by contracting and aligning the matrix (Gaggioli et al., 2007; Goetz et al., 2011; Calvo et al., 2013). Indeed, live imaging of cancer cell spheroids and CAFs in collagen revealed that CAFs were active in remodeling the matrix (Video 1). CAFs aligned and pulled collagen fibers perpendicularly to the edge of the spheroids, facilitating the migration of cancer cells (Video 2 and Fig. 2, B and C), whereas in the absence of CAFs, collagen fibers were oriented parallel to the spheroid edge (Fig. 2 C), which is not favorable for cancer cell invasion (Provenzano et al., 2006; Kopanska et al., 2016). However, FN-depleted CAFs (CAFsiFN) retained the ability to align collagen fibers in the same fashion (Fig. 2 C). Analysis of the width and length of collagen fibers remodeled by CAFsiCtrl and CAFsiFN also showed no difference between the two cell populations, demonstrating that FN depletion had no effect on the overall topography of the collagen matrix (Fig. 2 D). CAFsiFN also contracted and applied mechanical forces on the matrix similarly to control CAFs, indicating that depletion of FN in CAFs has no consequence on their ability to apply mechanical forces on the matrix (Fig. 2 E and F).

When inhibiting the contractility of CAFs using myosin II inhibitor blebbistatin (Fig. S1 E), collagen alignment and contraction were abrogated, as well as downstream FN assembly, as previously shown (Fig. 2 G; Zaidel-Bar et al., 2007). In this condition, cancer cells did not invade the matrix, either alone or in the presence of CAFs (Fig. S1 F). Together, these results demonstrate that both contractility and FN are important for CAF-mediated cancer cell invasion. However, the overall ability of fibroblasts to induce cancer cell invasion did not significantly correlate with their capacity to contract collagen, especially in the case of couple 3, where NAFs and CAFs displayed similar collagen contraction (Fig. S1 G). This suggests that mechanical forces are important for CAF-mediated cancer cell invasion, as they induce FN assembly. However, if not followed by FN deposition, mechanical forces by CAFs are not sufficient to promote invasion.

Finally, it has been suggested that invadopodia, the actin-rich structures responsible for matrix degradation, could also exert mechanical forces on the matrix and switch on a contractile phenotype (Aung et al., 2014). As contraction and degradation of the matrix could be interdependent, we also checked for the role of proteolysis in CAF-mediated cancer cell invasion. Inhibition of matrix proteolysis using broad spectrum matrix metalloproteinase (MMP) inhibitors GM6001 and
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Figure 1. CAFs induce invasion of cancer cells through matrix remodeling. (A) Maximum intensity projections of cancer cell spheroids in collagen I, with or without fibroblasts, at day 3. CT26 cancer cells express LifeAct-GFP (green); F-actin (red) and DNA (cyan) were stained with phalloidin-rhodamine and DAPI, respectively. Bar, 100 µm. (B) 3D rendering of spheroids at day 3. Invasion is quantified using the invasion counter software. Red dots represent nuclei of invading GFP cancer cells (migrated out of the spheroid). (C) Quantification of cancer cell invasion alone (blue box) and in the presence of NAFs (green boxes) or CAFs (red boxes) for three different patients. P-values are compared with cancer cells alone (in gray) and to cancer cells with CAFs (in black) using Newman-Keuls multiple comparisons test (*, P < 0.05; **, P < 0.01; ***, P < 0.001). (D, left) Quantification of the density of NAFs and CAFs from all three patients around the spheroid. Fibroblast density is defined as the number of nuclei of non-GFP cells normalized to the surface area of the spheroid contour in 3D. P-value is calculated using Mann–Whitney test. (Right) Quantification of the mean distance of NAFs and CAFs from the spheroid. The mean distance from the spheroid was defined as the distance from the nuclei of non-GFP cells to the closest point along the cancer cell spheroid contour. P-value is calculated using Mann–Whitney test. (E, left) Schematic representation of the experiment. Cancer cells were embedded in collagen gels (a), CAFs were either mixed with cancer cells in the collagen droplet (b), their conditioned media was added to cancer cells (c), or they were plated around the collagen droplet (d). (Right) Quantification of cancer cell invasion alone, in the presence of CAFcm or CAFdm, or in the presence of CAFs for three different patients. Invasion index is defined as the ratio between the number of invading nuclei of GFP cancer cells and the area of the spheroid contour. Quantification results are expressed as box and whiskers (minimum to maximum) of at least n = 3 separate experiments. P-values are compared with cancer cells alone (in gray) and to cancer cells with CAFs (in black) using Newman-Keuls multiple comparisons test (*, P < 0.05; **, P < 0.01; ***, P < 0.001). CC, cancer cells.
Figure 2. CAFs deposit FN to induce cancer cell invasion. (A) Quantification of cancer cell invasion alone or in the presence of CAFs from patient 2, depleted or not for FN. P-values are compared with cancer cells alone (in gray) and to cancer cells with CAFs (in black) using Newman-Keuls multiple comparisons test (***, P < 0.001). (B) Time-lapse sequence of CT26 cancer cells and CAFs from patient 1 in collagen. CT26 cancer cells express LifeAct-GFP (green), CAFs are stained with a lyophilic carbocyanine dye (red), and collagen is acquired by reflection (blue). Time is in hours and minutes (HH:mm). Bar, 200 µm. The magnified region is represented by the white square. Bar, 100 µm. [C, top] Overlaid images of collagen I matrices containing cancer cells. [C, bottom] Gel contraction (%) of CAFs from patient 1. [D] Overlaid images of Collagen I matrices containing cancer cells with F-actin, FN, and CAFs. [E] Overlaid images of collagen I matrices containing cancer cells with Strain Energy and CAFs from patient 1. [F] Overlaid images of collagen I matrices containing cancer cells with FN, F-actin, and CAFs from patient 1. [G] Overlaid images of collagen I matrices containing cancer cells with FN, F-actin, and CAFs from patient 1.
BB94 abrogated the spontaneous invasion of cancer cells, as previously shown (Poincloux et al., 2009; Wolf et al., 2013). Interestingly, CAFs rescued cancer cell invasion in the presence of MMP inhibitors (Fig. S1 H), indicating that CAFs can stimulate an invasion of cancer cells that is independent of MMPs. When treated with BB94, CAFs still contracted collagen plugs; thus, CAFs’ ability to contract the matrix did not depend on their ability to degrade it (Fig. S1 I). Interestingly, FN-depleted CAFs failed to rescue cancer cell invasion in the presence of MMP inhibitors (Fig. 2 H and Fig. S1 J).

Together, these data show that in the presence of CAFs that are able to deposit FN, cancer cells can invade the matrix in an MMP-independent manner. These findings could provide an explanation of the failure of MMP inhibitors in clinics. As a major constituent of the tumor microenvironment, CAFs can provide an alternative escape mechanism for cancer cells by aligning collagen fibers and assembling the FN that enables cancer cell invasion.

CAFs secrete and assemble more FN than NAFs

FN fibrillogenesis is a complex multistep process. Cells secrete FN as soluble dimers, which then bind to integrin receptors, unfold, and associate with each other to form a fibrillar matrix (Wolanska and Morgan, 2015). It is possible that CAFs are more efficient in assembling FN than NAFs and consequently induce more invasion of cancer cells. To address this, we compared the capacity of CAFs and NAFs to secrete and assemble FN (Fig. 3, A and B).

The analysis of the conditioned media showed that CAFs secrete more FN compared with NAFs (Fig. 3 A). Similarly, CAFs assembled more FN fibrils with their paired NAFs (Fig. 3 B). Moreover, the amounts of secreted and assembled FN by fibroblasts significantly correlated with the invasion index (Fig. 3 C). This result indicates that the ability of fibroblasts to induce cancer cell invasion directly correlates with the amount of FN they produce.

To further discriminate between the roles of secreted and assembled FN in cancer invasion, based on the estimation of the amount of FN secreted by CAFs (Fig. 3 A), we added 250 ng/ml of soluble FN to cancer cell spheroids. We observed that supplementing collagen with soluble FN did not induce invasion (Fig. 3 D). This was not surprising, as CAFs’ secreted molecules did not promote cancer cell invasion (Fig. 1 C).

Together, these results show that CAFs secrete and assemble FN more efficiently than NAFs and point toward the importance of FN assembly in CAF-mediated cancer cell invasion. As the invasion induced by all fibroblast populations most significantly correlated with the amount of assembled FN, we uncover a signature of CAFs and a link between ECM remodeling by CAFs and cancer cell invasion. These results are in agreement with a new study in which the analysis of FN expression in tumors from 435 head and neck cancer patients revealed an inverse correlation between high levels of FN and patient prognosis (Gopal et al., 2017). Besides serving as a cue in the ECM that cancer cells haptotact toward (Oudin et al., 2016), FN also establishes chemotactic gradients by modulating the bioavailability of growth factors such as hepatocyte growth factor (Rahman et al., 2005; Hynes, 2009). Therefore, by being in close proximity to each other, CAFs could stimulate the invasion of cancer cells toward FN and the growth factors FN matrices retain.

Integrin-αvβ3 is necessary for FN assembly

As soluble FN did not stimulate cancer cell invasion, we addressed the role of assembled FN. FN is assembled via transmembrane proteins, integrins; more specifically, mostly via integrin-α5β1 and -αvβ3 (Wolanska and Morgan, 2015). We correlated the amounts of integrin isoforms α5, αv, β1, and β3 in CAFs to their ability to induce invasion. Integrin-αv and -β3 expression showed the highest correlation, hinting at the importance of integrin-αvβ3 in CAF-mediated cancer cell invasion (Fig. S2, A and B).

We next tested whether β3- or α5-depleted CAFs (Fig. S2, C and D) could induce invasion. In this condition, the amount of secreted FN by CAFs was unchanged (Fig. S2 E). In the presence of both CAFsi-α5 and CAFsi-β3, invasion was significantly reduced (Fig. 4 A). This effect was confirmed using cilengitide, an inhibitor of integrin-β3 (Fig. 4 B). FN assembly by CAFsi-β3 was markedly reduced compared with control CAFs and α5-depleted CAFs (Fig. 4 C and Fig. S2 F) and was not caused by an impaired migration of CAFs toward cancer cells (Fig. S2 G). These results indicate that integrin-β3 and -α5 are necessary for CAF-mediated cancer cell invasion. They also validate the requirement of FN assembly for cancer cell invasion, as silencing of integrins or blocking of integrin-β3 or -α5 by contractility in CAFs does not affect FN secretion but abrogates cancer cell invasion (Fig. S2 E).

Because the fluorescence signal in 3D assays is nontrivial to quantify, we assessed the amount of assembled FN on 2D cell spheroids alone or together with control or FN-depleted CAFs from patient 1 generated using CurveAlign. The yellow line indicates the edge of the spheroid, and the green lines indicate fiber orientation with respect to the closest point on the spheroid edge. Bar, 100 μm. (Bottom) Rose plots representing the frequency of distribution of the absolute angles of collagen fibers within the range of 0–90° with respect to the closest point on the spheroid edge. (D, left) Maximum intensity projections of CAFs from patient 1 treated with siRNA scrambled control (CAFsCtrl) or with siRNA targeting FN (CAFsFN). F-actin is stained with phalloidin-rhodamine (green), FN is immunostained (magenta), and collagen is acquired using second harmonic generation (cyan). Bar, 20 μm. (Right) Quantification of collagen fiber width and length using CFIRE. P-value is calculated using Mann–Whitney test for at least n = 3 separate experiments. E, left) Control and FN-depleted CAFs from patient 1 cultured in collagen I gels 1 d after embedding. (Right) Percentage of gel contraction of control and FN-depleted CAFs from patient 1 calculated using the formula 100 x [gel area (T0) – gel area(T)] / gel area (T0). P-value is calculated using Mann–Whitney test for n = 3 over n = 6 separate experiments. F, left) Traction force map of control and FN-depleted CAFs from patient 1 on collagen-coated polycarbonate gels with Young’s modulus of 5 kPa. Color code gives the magnitude of traction stress in Pa, which corresponds to forces of piconewton/squared micrometers. (Right) Corresponding mean force (strain energy) exerted by CAFs over a 30-min time lapse. P-value is calculated using Mann–Whitney test for n = 10 cells over n = 2 separate experiments. (G) Maximum intensity projections of cancer cell spheroids in collagen I gels with CAFs from patient 1 treated with siRNA scrambled control, siRNA against FN, or blebbistatin at day 3. Bar, 100 μm. Magnified regions are represented by the white squares. CT26 cancer cells express LifeAct-GFP (green), F-actin is stained with phalloidin-rhodamine (red), FN is immunostained (cyan), and collagen is acquired using reflection [white]. Bar, 50 μm. (H) Quantification of cancer cell invasion in either the presence of control or FN-depleted CAFs treated with BB94 from patient 3. Invasion index is defined as the ratio between the number of invading nuclei of GFP cancer cells and the area of the spheroid contour. All quantification results are expressed as box and whiskers [minimun to maximum] of at least n = 3 separate experiments. P-values are compared with cancer cells alone (in gray) and to cancer cells with CAFs (in black) using Newman-Keuls multiple comparisons test (*, P < 0.05; **, P < 0.01; ***, P < 0.001).
substrates. Surprisingly, depletion of integrin-α5 did not alter FN assembly by CAF 1 d after plating, though the ability to assemble FN was reduced upon depletion of integrin-β3 (Fig. S3 A). In contrast, when fibroblasts were cultured for 3 d and allowed to reach confluency, depletion of integrin-α5 or -β3 reduced FN fibrillogenesis (Fig. 4 D). These results suggest that integrin-β3 mediates FN assembly at an early stage, whereas α5 could be required later on for sustained assembly of FN fibers.

Integrin-α5 and -β3 are required at different stages of FN fibrillogenesis

As our results indicate that both integrin-α5β1 and -αvβ3 are required for FN fibrillogenesis, we next wondered about their localization in CAFs and with respect to FN fibers. In 2D, integrin-α5 was found in the center of the cells in mature fibrillar adhesions colocalizing with FN fibers. αvβ3 was present at the cell periphery, at focal adhesions partially colocalizing with FN puncta (Fig. 5 A). In 3D collagen matrices, integrin-αvβ3 accumulated at the cell poles along with FN deposits (Fig. 5 B), whereas integrin-α5 was localized all along the cell periphery (Fig. S3 B).

The localization of integrin-αvβ3 at the cell periphery suggests its requirement during initial cell–matrix interactions. Indeed, after letting the cells adhere for 2 h, integrin-αvβ3 already clustered, whereas integrin-α5 was not yet recruited (Fig. 5 C). However, as blocking of integrin-β3 did not alter the CAFs’ contractility (Fig. S3 C), these results indicate that integrin-αvβ3 is required at the initial steps of FN fibrillogenesis, downstream of contractility.
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Integrin-β3 specifically accumulates at areas of high-traction force and is stationary within focal adhesions, whereas integrin-β1 is more mobile and thus prone to relocate to fibrillar adhesions (Rossier et al., 2012; Schiller et al., 2013). Inhibition of myosin disassembles β3 clusters without affecting levels and localization of integrin-β1 (Schiller et al., 2013). This could explain why treatment of CAFs with blebbistatin abrogates FN assembly. However, depletion of αvβ3 does not alter CAFs’ ability to contract collagen, indicating that αvβ3 is downstream of contractility.

Integrin-αvβ3 is necessary for FN assembly. (A) Quantification of cancer cell invasion alone or in the presence of control CAFs, α5-depleted CAFs, and β3-depleted CAFs from patient 3. (B) Quantification of cancer cell invasion in the presence of CAFs from patient 2, with or without cilengitide treatment. (A and B) Invasion index is defined as the ratio between the number of invading nuclei of GFP cancer cells and the area of the spheroid contour. All quantification results are expressed as box and whiskers [minimum to maximum] of at least n = 3 separate experiments. P-values are compared with cancer cells alone (in gray) and to cancer cells with CAFs (in black) using Newman-Keuls multiple comparisons test (*, P < 0.05; **, P < 0.01; ***, P < 0.001). (C) Maximum intensity projections of CAFs from patient 2 treated with siRNA scrambled control, siRNA against integrin-α5 or integrin-β3, or cilengitide. Bar, 100 µm. Magnified regions are represented by the white squares. F-actin is stained with phalloidin-rhodamine (green), FN is immunostained (magenta), and collagen is acquired using second harmonic generation (cyan). Bar, 20 µm. (D, left) Immunostaining of FN (green) in control CAFs, FN-depleted CAFs, α5-depleted CAFs, and β3-depleted CAFs from patient 2, 3 d after plating. F-actin was stained with phalloidin-rhodamine (red), and DNA was stained with DAPI (blue). Bar, 40 µm. (Right) Graph represents the percentage of assembled FN relative to control conditions defined as the amount of fluorescence in a monolayer (integrated density) normalized to the number of nuclei. Quantification results are expressed as column bars with mean ± SEM. Depleted CAFs were compared with control CAFs for n = 10 frames over n = 4 separate experiments. P-value is calculated using Newman-Keuls multiple comparisons test (*, P < 0.05; **, P < 0.01; ***, P < 0.001).
In cells plated on RGD-rich substrates, αvβ3 localizes at focal adhesions and, unlike β1, is responsible for force-dependent focal adhesion maturation (von Wichert et al., 2003; Roca-Cusachs et al., 2013; Chagede et al., 2015). Thus, αvβ3 could be involved in mediating the signal, from contractility to formation of fibrillar adhesions. This hypothesis is supported by a study showing that pulling on vascular smooth muscle cells using anti-β3 antibody–coated beads, and not α2 or β1, resulted in the activation of downstream signaling (Goldschmidt et al., 2001). Similarly, a blockade of integrin-β3, but not -β1, abrogates DNA synthesis induced by mechanical strain in vascular smooth muscle cells (Wilson et al., 1995).

Because in our study we used collagen, which is not an RGD substrate, αvβ3 localization at focal adhesions is
somewhat surprising. The possible explanation is that because of
the presence of serum and cellular FN secreted by the cells
themselves, cells are constantly exposed to RGD motifs. It has
also been shown that αβ3 localizes to the cell’s edge inde-
pendently of RGD upon stimulation by growth factors such as
FGF (Kiosses et al., 2001). In the context of a tumor where
CAFs, blood vessels, or cancer cells constantly secrete FN and
growth factors, the nature of the matrix would not really matter,
as it is constantly enriched in RGD peptides.

In conclusion, our study shows that diffusible molecules
secreted by CAFs are not sufficient to induce cancer cell inva-
sion. Instead, CAFs’ remodeling of the matrix is the key player.
We propose a model in which contractility of CAFs is neces-
sary for downstream activation of the integrin-αβ3 and assem-
by of FN puncta. αβ1 becomes critical only at later stages
of FN assembly at fibrillar adhesions (Fig. 5 C). Because the
ability of all fibroblast populations to assemble FN directly
 correlates with their ability to induce cancer cell invasion, we
demonstrate that ECM deposition (more specifically, FN fibril-
logenesis) is the key component for CAF-mediated cancer cell
invasion. Finally, we show that integrin-β3 is the major driver
for FN assembly in CAFs, as its inhibition abrogates CAF-
mediated cancer cell invasion.

**Materials and methods**

**Cell lines**

CT26 mouse intestinal cancer cells were obtained from American Type
Collection. Cells were cultured in DMEM (Life Technologies)
supplemented with 10% FBS (Invitrogen) and 5% CO2. Cells were in-
fected with a lentiviral GFP plasmid according to standard procedures.

**Isolation and culture of primary fibroblasts**

Human primary fibroblasts were isolated from fresh colon tumors
(CAFs) and adjacent noncarcinoma tissue (NAFs) samples from pa-
tients treated at Lariboisière Hospital, Paris, with the written consent
of the patients and approval of the local ethics committee. Samples were
treated as previously described (Amatangelo et al., 2005). In brief, tis-
nues were collected after surgical resection in RPMI buffer and washed
in PBS supplemented with 10% antibiotic–antimycotic (AA; Gibco).
100-mm2 tissue culture plates were scratched using a scalpel, and tis-
ue pieces of ~1–2 mm2 were cut and placed on the junctions. After
isolation, the tumor pieces were kept in 10 ml DMEM supplemented
with 10% FBS (Life Technologies) and 10% AA. Cells were in-
fected with a lentiviral GFP plasmid according to standard procedures.

**Western blotting**

Protein lysates were obtained from fibroblasts seeded on soft plates
at passage 3. Protein lysates were processed according to standard
procedures. In brief, cells were washed with PBS, lysed in radio-
immunoprecipitation assay buffer (1% Triton X-100, 50 mM Tris,
ph 7.5, 1 mM EDTA, and 150 mM NaCl) supplemented with prote-
ase and phosphatase inhibitor cocktails (Sigma-Aldrich), and boiled in
Laemmli buffer for 5 min. The samples were separated in SDS-PAGE
gradient gels (4–15%), transferred to a nitrocellulose membrane using
the BioRad system, and blocked in 5% nonfat dried milk dissolved in
PBS supplemented with 0.1% Tween for 30 min at RT. The membranes
were incubated with primary antibodies overnight at 4°C followed by
incubation with peroxidase-conjugated secondary antibodies for 1 h
at RT. Immunoreactive bands were detected using an ECL-plus kit
(Roche). Quantifications were done using ImageJ (National Institutes
of Health) by normalizing the protein amount to α-tubulin or GAPDH
amounts (loading controls). Antibody description and working dilu-
tions can be found in Table S1.

**Invasion assay**

Agarose (Invitrogen) was dissolved in water to a concentration of
0.01 g/ml and boiled. 150 µl of the solution was added to a 48-well
plate, and agarose was left to polymerize for at least 10 min at RT. A
solution of 104 cells/ml of CT26 cancer cells was made, and 75–100 µl
of the solution was added to the wells. The wells were subsequently
filled with DMEM supplemented with 10% FBS, and spheroids were
left to form for 3–4 d.

30-mm2 tissue culture plates were specifically fashioned for
the invasion assay: three holes ~3–4 mm in 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 square
coverslips were glued to the dish overnight at RT. A d before the experi-
ment, the dishes were silanized with 3-aminopropyl-trimethoxysilane
(Sigma-Aldrich). The dishes were washed extensively with water and
treated for 30 min with 0.5% glutaraldehyde followed by a final wash.
This treatment was made to avoid collagen detachment from the plastic
holes because of the high contractility exerted by CAFs.

2 mg/ml of rat tail collagen I (Corning) was prepared in DMEM,
10x PBS, and 1 M NaOH, to a pH of 7. The solution was kept on ice
to avoid collagen polymerization. Spheroids were embedded in 15 µl
collagen drops containing 5 × 103 fibroblasts, positioned in the hole
of the culture plate. After filling all three holes, the plate was flipped
every 2–3 wk. When having reached confluency, fibroblasts were trypsinized and plated on 30 KPa
30-mm2 soft plates (Excellness) that were previously coated with
5 µg/ml of rat tail collagen I (Corning) in DMEM polymerized at 37°C
for at least 24 h. Soft plates were used to avoid activation of fibro-
blasts by matrix rigidity. Unless stated otherwise, all fibroblasts were
cultured on soft plates and kept in their primary nontransformed state
until passage 10. After isolation, all cell populations were character-
ized for the presence of markers of activated fibroblasts: α-SMA, FAP,
and PDGFR-β (Fig. S1 A).

**Collagen contraction assay**

1.5 × 106 fibroblasts were suspended in 1.5 ml of 2-mg/ml rat tail col-
lagen I (Corning) and added to a 24-well plate in triplicates (500 µl
well). After 30 min of incubation at RT, collagen plugs were detached
from the walls of the well with a scalpel, and DMEM supplemented
with 10% FBS was added. Images of the collagen plugs were acquired
at time 0 (T0) and after 24 h (T1) using a microscope (M165FC; Leica). To obtain the gel contraction value, the relative area of the gel was measured using ImageJ software at T0 and T1, and the percentage of contraction was calculated using the formula 100 × [gel area (T0) – gel area (T1)]/gel area (T0).

Collagen topography measurements
Fibers’ alignment and their angles with respect to the spheroid edge were measured on images acquired using reflection microscopy with the available software, CurveAlign in MatLab. The angles of collagen fibers compared with the spheroid edge were determined for 10 slices per condition. Fiber width and length were measured on images acquired using second harmonic generation with the available software, CT-FIRE in MatLab.

Inhibitors and siRNA
Blebbistatin (Sigma-Aldrich), BB94 (AbCam), and cilengitide (Selleckchem) were used at 15, 5, and 1 µM, respectively. They were mixed with the media and added to the invasion assay after collagen polymerization. GM6001 (Millipore) was used at 20 µM and was mixed with both the collagen before its polymerization and added to the media as previously described (Wolf et al., 2013).

For protein knockdowns using siRNA, CAFs were cultured in standard conditions and transfected using HiPerFect (301704; Qiagen). 6 × 10⁵ CAFs were plated in a well of a 6-well plate and subjected to transfection using 100 nM siRNA. siRNA was purchased from Qiagen, and sequences are listed in Table S2.

3D immunofluorescence
Spheroids embedded in collagen were fixed using 4% PFA in PBS for 30 min at RT and washed with PBS. Anti-FN antibody was used in 500 µl PBS, and dishes were left under agitation at RT for 2 d. Spheroids were then washed with PBS and permeabilized with 0.1% Triton X-100 in PBS for 30 min at RT. DNA and F-actin were stained using DAPI and phalloidin, respectively (Life Technologies). Collagen was imaged using confocal reflection microscopy.

For integrin stainings, CAFs were embedded in collagen drops for 3 d in identical culture conditions and then fixed and stained similarly to spheroids. Permeabilization was performed for 15 min to avoid damaging the cell membrane. Antiintegrin antibodies were added in 500 µl PBS, and the dishes were left under agitation at RT for 2 d. Appropriate secondary antibodies were added in 500 µl PBS along with phalloidin-rhodamine. Collagen was imaged using second harmonic generation. In all conditions, cells were imaged at a similar distance from the coverslip using the same settings.

Quantification of FN secretion and expression
CT26 cancer cells were incubated in 1 ml of serum-deprived DMEM at a density of 10⁶ cells for 24 h. The media were collected, passed through 0.2-µm filters to eliminate cell debris, and added on soft plates and sequences are listed in Table S2.

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CT26 cancer cells were incubated in 1 ml of serum-deprived DMEM at a density of 10⁶ cells for 24 h. The media were collected, passed through 0.2-µm filters to eliminate cell debris, and added on soft plates containing 1.5 × 10⁵ fibroblasts for 3 d. The media were collected, through 0.2-µm filters to eliminate cell debris, and added on soft plates and sequences are listed in Table S2.

Transfection using 100 nM siRNA. siRNA was purchased from Qiagen, and sequences are listed in Table S2.

For quantification of FN secretion and expression in CAFs and NAFs, fibroblasts were plated on glass coverslips in CT26-conditioned media. For the remaining 2D stainings, fibroblasts were plated on glass coverslips in DMEM supplemented with 10% FBS. For staining of early FN assembly and integrin localization, cells were fixed 24 h after plating. For staining of mature FN fibers on confluent monolayers, cells were fixed 3 d after plating. Cells were fixed using 4% PFA in PBS for 20 min at RT and washed with PBS. Anti-FN antibody was added for 1 h at RT. Cells were then washed and permeabilized with 0.1% Triton X-100 in PBS for 5 min at RT. Antiintegrin antibodies were added for 1 h at RT.

For quantification of FN assembly in CAFs and NAFs, fibroblasts were plated on glass coverslips in DMEM supplemented with 10% FBS. For staining of early FN assembly and integrin localization, cells were fixed 24 h after plating. For staining of mature FN fibers on confluent monolayers, cells were fixed 3 d after plating. Cells were fixed using 4% PFA in PBS for 20 min at RT and washed with PBS. Anti-FN antibody was added for 1 h at RT. Cells were then washed and permeabilized with 0.1% Triton X-100 in PBS for 5 min at RT. Antiintegrin antibodies were added for 1 h at RT. After an additional round of PBS washing, secondary antibodies along with DAPI and phalloidin were added for 1 h at RT to stain DNA and F-actin, respectively. Coverslips were mounted on slides in AquaPolymount (Polysciences) and imaged using an upright wide-field microscope (DM6000; Leica) with a 63x/1.32 NA oil immersion objective. The images were processed and quantified with ImageJ. The amount of assembled FN per cell is calculated by normalizing the amount of fluorescence in a cell (integrated density) to the area of the cell and the background fluorescence.

Traction force microscopy
Traction force microscopy experiments were performed as previously described (Elkhatib et al., 2014). For traction force measurements, Fluosphere bead solution (0.2 µm, 505–515 nm; Invitrogen) was added at 2.5% volume. For time-lapse imaging, we used an inverted wide-field confocal spinning-disk microscope (40x oil immersion objective, NA 1.3; Roper/Nikon). A fluorescent image of beads and a phase contrast image of the cells were recorded every 3 min during 30 min. At the end of the measurement, cells were detached by adding 10% Triton X-100 (EuroMedex), and a reference image without cells was recorded. To ensure good quality imaging of fluorescent beads, we performed Z stacks of 30 images with a distance of 1 µm and automatically chose the optimal focus (MetaMorph software). We used a previously described correlation algorithm to extract the bead displacement fields (Elkhatib et al., 2014).

Imaging 3D samples
For time-lapse experiments, CAFs were stained with a lyophilic carbocyanine dye (Vybrant Dil-Cell Labeling Solutions; ThermoFisher) according to the manufacturer’s recommendation. Cells were embedded in collagen as described in the Invasion assay section. The dish was incubated at 5% CO₂ and 37°C in the on-stage incubator (Okolab). For fixed and live 3D samples, images were acquired with an inverted AOB5 two-photon laser-scanning confocal microscope (SP8; Leica) coupled with a femtosecond laser (Chameleon Vision II; Coherent Inc.)
using a 25×1.0 NA water immersion objective. The microscope was equipped with three nondescanned HyD detectors: NDD1 (500–550 nm), NDD2 (≥590 nm), and NDD3 (405 nm). Fluorescence channels were recorded simultaneously using the excitation wavelength 980 nm. Collagen was visualized by either second harmonic generation using the excitation wavelength 910 nm or by confocal reflectance microscopy that does not interfere with DAPI staining, using light at a wavelength of 488 nm and a standard photomultiplier tube detector at a low gain (500 V). Images were recorded every 10 min up to 72 h. 3D stacks were obtained at a step size of 2-µm intervals. The images were processed with the Leica Application Suite, ImageJ, and Imaaris (Bitplane).

Invasion counter software
Quantification of cell invasion from spheroids was performed using a custom semiautomated image analysis program written in Python using the following packages: numpy, sciPy, matplotlib, scikit-image, and PyQ4. Image stacks of nuclei were first loaded into a custom GUI, and the spheroid contour was determined using adjustable Gaussian filtering, thresholding, and 3D morphological operations. The nuclei of invading cells were then automatically detected using adjustable Gaussian filtering, thresholding, and size exclusion. Centroid positions were determined by taking a weighted mean of the intensity for each nucleus. The positions of invading cancer cell nuclei were then manually verified and modified as necessary. Based on the LifeAct-GFP signal (expressed in cancer cells only), the nuclei of cancer cells were discriminated from the nuclei of fibroblasts. The invasion index, defined as the number of invading cancer cells normalized to the surface area of the spheroid contour in 3D, was then determined. This normalization is necessary to control for the slight variability in spheroid size. Because of the high optical density of the spheroids, only the bottom half of the spheroid was visible. To quantify the distance of fibroblasts from the spheroid, the distance from the nuclei of non–LifeAct-GFP cells to the closest point along the cancer cell spheroid contour was determined. Fibroblast density was defined as the number of nuclei of non-GFP cells normalized to the surface area of the spheroid contour in 3D.

Statistical analyses
All experiments were performed in triplicates in two to six independent experiments. All statistical analysis and graphic representations were performed using Prism software (GraphPad). Data are represented as box and whiskers (minimum to maximum). To show protein amounts and percentages, data are represented as column bars (mean ± SEM). Statistical significance was determined with one-way ANOVA. The Newman-Keuls test was applied for multiple comparisons of different conditions. The Mann-Whitney t test was applied for paired comparisons.

Online supplemental material
Fig. S1 shows (a) the levels of CAF markers in all fibroblast populations, (b) the effect of FN depletion on cancer cell invasion in CAFs from all couples, (c) that blocking of contractility inhibits cancer cell invasion but that contractility of fibroblasts does not correlate with invasion, and (d) the effect of MMP inhibition on cancer cell invasion. Fig. S2 shows the levels of integrins in all fibroblast populations and the effects of integrin-α5 and -β3 depletion on fibroblast viability, migration, FN secretion, and cancer cell invasion. Fig. S3 shows that depletion of integrin-β3 inhibits FN assembly in CAFs 24 h after plating, but not contractility and integrin-α5 localization in CAFs embedded in collagen. Videos 1 and 2 show live imaging of cancer cell spheroids and CAFs embedded in collagen. Tables S1 and S2 list antibodies and siRNA sequences, respectively, used in this study.

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Author contributions: Y. Attieh and D.M. Vignjevic conceived of the study. Y. Attieh performed the majority of the experiments. C. Grass helped with experiments. N. Elkhatib and T. Betz assisted with traction force microscopy. A.G. Clark wrote the program to quantify cell invasion. P. Mariani, M. Pocard, and S. Richon collected clinical samples for the study. B. Gurchenkov assisted with live experiments. Y. Attieh and D.M. Vignjevic wrote the manuscript with input from A.G. Clark.

D.M. Vignjevic supervised the project.

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