TIMP-2 secreted by monocytes is a potent suppressor of invadopodia formation in pancreatic cancer cells

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Abstract

Background Monocytes are a major component of the tumor microenvironment (TME) in pancreatic ductal adenocarcinoma (PDAC). However, the complex interactions between tumor cells and monocytes and their role in tumor invasion have not been fully established. Methods In this in vitro study, to specifically test the impact of interaction on invasive potential, two PDAC cell lines PaTu8902 and CFPAC-1 were selected on their ability to form invasive adhesions, otherwise known as invadopodia. Co-culture experiments were performed using undifferentiated THP1 monocytes. Results When the PDAC cells were co-cultured with undifferentiated THP1 monocytes invadopodia formation was significantly suppressed. Moreover, conditioned media of THP1 cells (CM) was also able to suppress invadopodia formation. Further investigation revealed that both tissue inhibitor of metalloproteinase (TIMP) 1 and 2 were present in the CM. However, suppression of invadopodia formation was found that was specific to TIMP2 activity. Conclusions Our findings indicate that TIMP2 levels in the tumour microenvironment may have prognostic value in patients with PDAC. Furthermore, activation of TIMP2 expressing monocytes in the primary tumour could present a potential therapeutic opportunity to suppress cell invasion in PDAC.

Background
Pancreatic ductal adenocarcinoma (PDAC) is a devastating disease characterised by an aggressive biological tumour behaviour leading to a high mortality rate [1]. The invasive character of PDACs leads to a rapid progression of the disease [2]. In order to invade the surrounding tissue or to create distant metastases, cancer cells need to break through the basement membrane and degrade the extracellular matrix. One identified strategy is to utilise actin-rich membrane protrusions called invadopodia [3] that can degrade extracellular matrix. Initiating mechanisms to suppress such formation is a potential therapeutic target [4]. Unlike podosomes - which are present in normal myeloid cells – invadopodia are specific for invasive cancer cells [5]. The primary purpose of invadopodia is the targeted secretion of matrix metalloproteases (MMPs) to degrade the extracellular matrix (ECM) [6]. Membrane-type 1 metalloprotease (MT1-MMP, also known as MMP14), MMP9 and MMP2 have been identified as the most important MMPs in invadopodia functionality [7, 8].
MMPs can be inhibited by tissue inhibitors of metalloproteinases (TIMPs). Of the four TIMPs [9, 10], TIMP1 and 2 are the best-studied. TIMP1 has a relatively low affinity for membrane-bound MMPs [11] whereas TIMP2 is a strong inhibitor of MT1-MMP [12]. Very little is known about the effects of TIMPs on invadopodia formation, although one study in human breast cancer cells suggested TIMP2 is specifically able to decrease the formation of invadopodia [13].

In recent years, the tumour microenvironment (TME) has become of major interest to researchers and has recently been suggested to play a specific role in influencing the formation of invadopodia [14]. Around 90% of the PDAC mass are stromal cells, only 10% are carcinoma cells [15]. The main components of the TME are cancer-associated fibroblasts (CAF) and tumour-associated macrophages (TAMs). The pancreatic TME is rich in TAMs but undifferentiated monocytes [16] are also present at detectable levels. The interaction between monocytes / macrophages and the tumor cells is complex with both pro- and anti-tumorigenic effects reported [17]. This study aimed to establish whether an interaction between PDAC cells and undifferentiated monocyte-like cells presented pro- or anti-tumorigenic responses.

Results

Pancreatic cancer cells produce invadopodia

As compared to the well-described characteristics of invadopodia formation in breast cancer cells, especially in the cell line MDA-MB 231 [18, 19], less is known about invadopodia in PDAC cells [20]. We screened 10 different PDAC cell lines for invadopodia formation and compared the findings with MDA-MB 231 breast cancer cells (Table I). Our screen revealed that PaTu8902 and CFPAC-1 constitutively form a large number of invadopodia. To confirm our observations, invadopodia areas of matrix degradation were co-localised with cortactin-positive puncta as a marker of invadopodia (Figure 1A). We found that PDAC cells require longer incubation times than previously reported for breast cancer cells [18] to generate quantifiable degradation activity and thus quantification of activity was based on the total area of gelatin degradation per field of view. PaTu8902 and CFPAC-1 cells revealed high levels of degradative activity (Figure 1B). Both invadopodia-forming cell lines PaTu8902 and CFPAC-1 expressed MT1-MMP whereas PaTu8988-S did not (Figure 1C). CFPAC-1 and
PaTu8902 were further tested for MMP9 (positive, Figure 1D) and MMP2 (no expression, data not shown). Neither CFPAC-1 nor PaTu8902 were expressing TIMP2 (Figure 1E).

**Monocyte-like cells co-culture suppresses invadopodia driven matrix degradation**

We performed a number of different co-culture experiments with eGFP-tagged THP1 cells (THP1; a commonly used cell line model for undifferentiated monocyte-like cells) and PaTu8902 cells (Figure 2A&B) or CFPAC-1 cells (Figure 2C&D). Either the PDAC cells and THP1 cells were cultured together prior to the invadopodia assay, or PDAC cells and THP1 cells were cultured together during the invadopodia assay, or conditioned medium from THP1 cells was added to the PDAC cells during the invadopodia assay. In all co-culture conditions, gelatin degradation was reduced compared to control in both cell lines (Figure 2AA-D). The suppressive effect of THP-1 CM was confirmed in the spheroid invasion assay. Since CFPAC-1 is not forming adequate spheroids, PANC-1 cells were used instead (Figure 2E,F,H&I). THP-1 CM did not alter proliferation and spheroid growth (Figure 2G&J).

**Tissue inhibitor of metalloproteinases 1 and 2 are secreted by THP1 monocyte-like cells**

Given that exposure to conditioned medium was sufficient to suppress invadopodia activity we explored this phenomenon in more detail in both cell lines. The gelatin degradation assays were repeated using a control condition (control media) and three different CM conditions. To assess the nature of the inhibitory factor, CM was either boiled, used immediately after collection (fresh CM) or freeze/thawed. Incubation of cells with fresh or freeze/thawed CM significantly reduced invadopodia formation whereas incubation with boiled CM was unable to significantly reduce activity compared to control cells (Figure 3A-D). These findings suggest that the factors secreted by THP1 cells to suppress invadopodia actively are likely to be protein based.

To further elucidate the nature of THP1 CM suppression of invadopodia activity the THP1 CM was screened against a microarray of selected hormones and cytokines (Figure 2E). The only proteins detected in significant concentrations were RANTES, Interleukin 8 (IL-8) as well as tissue inhibitor of metalloproteinases 1 and 2 (TIMP1 and TIMP2). Both RANTES [21] and IL-8 are suggested to rather promote than reduce cancer cell invasion, so we focused on the further examination of TIMP1 and TIMP2. Performing a western blot with specific antibodies we were able to detect both TIMP1 and...
TIMP2 in the CM samples consistent with our array and determine that the concentration of both TIMP1 and TIMP2 in CM was between 5 and 50 ng/ml (Figure 2G-J, for full-length gels and blots see Supplementary Dataset 1).

**Inhibition of invadopodia formation is driven specifically by TIMP2**

To assess the specific effects of TIMPs on invasion, the invadopodia assays were repeated adding commercial recombinant TIMP1 (rTIMP1) or recombinant TIMP2 (rTIMP2), respectively at two different concentrations, 5 ng/ml and 50 ng/ml, to cover the range of likely concentration in conditioned medium (Figure 3E&J). Whilst the presence of rTIMP1 had no impact on degradative ability (Figure 3A&B), the presence of rTIMP2 was able to significantly reduce gelatin degradation in the treated cells even at a concentration of 5 ng/ml (Figure 4A&C). Moreover, similar results were obtained when CFPAC-1 cells were treated with rTIMP2 but not with rTIMP1 (Figure 4D-F). Our results suggest a specific inhibition of invadopodia activity in PDAC cells exposed to low concentrations of TIMP2. However, we cannot rule out a global deleterious effect of rTIMP2 exposure on cell behaviour. To address this issue we tested whether cells could recover invadopodia activity if the rTIMP2 was removed. As a control, responses were compared to incubation with a pharmacological MMP inhibitor GM6001 which is known to transiently inhibit invadopodia formation until washed out [22] (Figure 5A&B). Subsequently, we tested the recovery of cells incubated with rTIMP2 for 5 hrs prior to removal (Figure 5C&D). Similar results were obtained when CFPAC-1 cells were examined in the same protocol (Figure 5E-H). Thus, the inhibition of invadopodia activity in the presence of rTIMP2 is a specific response. Furthermore, an MTT assay confirmed that neither THP-1 CM nor rTIMP2 had an effect on viability and proliferation (Table II).

**Discussion**

There is evidence suggesting there is a strong link in some tissue types between the ability of cancer cells to form invadopodia in vitro and their invasive potential in vivo [23]. However, only one previous study screened PDAC cell lines for invadopodia formation [20]. Although in this case, the authors reported invadopodia prevalence in most cell lines, this was not comparatively quantified and degradation activity was not assessed. Indeed, we found that some of the PDAC cell lines were
extremely inconsistent in invadopodia formation and could not be relied upon for reproducible studies. Other reports focus on one PDAC cell line examining specific pathways [24, 25]. We have now identified two PDAC cell lines that reproducibly potentiate significant invadopodia activity and can be confidently used to further studies in this area. With regards to invadopodia formation, the PDAC cell lines in this study were compared to MDA-MB-231 cells – a breast cancer cell line that is well known for its ability to form invadopodia. Consistent with the findings for different PDAC lines, some breast cancer cell lines are able to form invadopodia (such as MDA-MB-231 or BT-549) [26-28] whereas others are not (e.g. MCF-7) [29].

Using these cell lines we proceeded to investigate how the TME might impact on invasive activity. The TME has a particular role in PDAC – not only because the major part of PDACs consists of fibroblast and monocytes/macrophages [16] but also because there is a complex interaction between monocytes/macrophages and PDAC cells potentially modifying the invasive potential of PDAC cells [17]. Monocytes and macrophages in the TME appear to have both pro- and antitumor effects which are suggested to be due to different cytokine profiles depending on the differentiation and polarization [30, 31]. Most studies examining mononuclear cells as part of the TME are focusing on macrophages [32, 33], however, there is evidence that monocytes are present in the TME of PDAC [16]. Nevertheless, the prognostic value of monocyte infiltration of the PDAC TME has not been extensively explored.

Taken together, the results of current studies suggest that pro-inflammatory features of the TME determined by the polarization and pro-inflammatory cytokine profile leads to a more aggressive and invasive tumour behaviour and thus decreases prognosis.

This is the first study to show that monocyte-like cells – which are an essential part of the TME [16] – have the ability to reduce the formation of invadopodia in PDAC cells and thus highly significantly decrease their invasive migratory behavior, which would inhibit their metastatic potential.

Interestingly, THP-1 cells are able to degrade the ECM and form invadopodia as well [34]. Our results indicate that the suppressive activity of undifferentiated monocyte-like cells of the PDAC cells might be mediated via secretion of TIMP2 but not TIMP1. This is consistent with a report in breast
cancer cells that suggested TIMP2 suppressed invadopodia formation. However, in this study, activity was measured by positive cortactin staining but not matrix degradation [13]. The specific inhibitory effect of TIMP2 might be due to the differential inhibition of TIMP1 and TIMP2. Indeed, MMP2, MMP9, and MT1-MMP are considered the most important MMPs needed for invadopodia activity [35]. MT1-MMP, a membrane-bound MMP, has a key role not only degrading the ECM but also activating of MMP2 [36] which in turn activates MMP9 [37]. In contrast, TIMP1 inhibits soluble MMPs and has a very low affinity to membrane-bound MMPs such as MT1-MMP. Taken together, there is evidence that the TIMP2-mediated inhibition of MT1-MMP is the key to inhibiting invadopodia formation [38]. Nonetheless, we believe that there are other mechanisms of how undifferentiated monocyte-like cells can inhibit invadopodia formation as well since we found that a monocyte-like cell/PDAC cell co-culture prior to the actual invadopodia assay was also able to reduce cancer cell invasion (Figure 1C and D). These mechanisms could include transcriptional changes in the cancer cells, e.g. upregulation of S100A8 and S100A9 which is known to increase the invasive potential of cancer cells [39]. Interestingly, in human hepatocellular carcinoma (HCC) samples, downregulation of TIMP2 expression was significantly associated with liver invasion and poorer survival outcomes [40]. Currently, PDAC tumours are considered to be immunologically “cold” [41]. We would suggest that an aspiration to increase the immune infiltrate in the PDAC setting [42] should consider specifically increasing the presence of TIMP2 secreting monocyte-like cells.

We found that THP1 monocyte-like cells produce TIMP2 in relevant concentrations without being co-cultured with PDAC cells. THP1 CM is able to suppress invadopodia formation, a direct interaction between THP1 cells and PDAC cells is not necessary for this effect. However, there is evidence that a direct co-culture of mononuclear cells and cancer cells leads to changes both in MMP and TIMP expression levels [43]. Besides the examination of these interactions, further experiments could also use human monocyte derived macrophages in the stages of M0, M1, and M2 to see the differential effects on invasion depending on the polarisation of the macrophages. To translate these findings into an in vivo model, monocyte-like cells or differentiated macrophages could be injected and isolated in nude/SCID mice.
In conclusion, our findings are indicative that TIMP2 could both be a potential prognostic marker and a therapeutic target in PDAC. A high ratio of TIMP2-secreting monocyte-like cells could be associated with reduced metastatic potential and better prognosis. Furthermore, the application of synthetic TIMP2 agonists could possibly lead to a reduction of cancer cell invasion in vivo.

**Methods**

**Cell culture**

Patu8902 and Capan2 were obtained from “Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH” [DSMZ], Germany), Capan1 and MiaPaCa2 were kindly provided by Prof. H. Kocher at Barts Cancer Institute, UK. The human breast cancer cell line MDA-MB-231 was purchased from ATCC. All these cells were maintained in Dulbecco’s modified Eagle’s media (DMEM, Sigma Aldrich UK) supplemented with 10% v/v fetal bovine serum (FBS) and 1 mM penicillin/streptomycin. CFPAC-1 cells (kindly provided by Dr. A. Pessina at Universita degli studi di Milano, Italy) were cultured in Iscove’s modified Eagle’s media (IMEM, Sigma Aldrich UK) supplemented with 10% v/v fetal bovine serum (FBS) and 1 mM penicillin/streptomycin. Colo-357 (kindly provided by Prof. Michalski University Hospital Heidelberg, Germany) and AsPC1 (kindly provided by Dr Stéphanie Kermorgant, Barts Cancer Institute, UK) were cultured in Roswell Park Memorial Institute-1640 (RPMI-1640, Sigma Aldrich UK) media supplemented with 10% v/v FBS with 1 mM penicillin/streptomycin. THP-1 cells (obtained by the European cell collection bank) were tagged with enhanced green fluorescent protein (eGFP) using lentiviral vector technique as described before [44]. Cells were maintained in RPMI-1640 media supplemented with 10% v/v FBS with 1 mM penicillin/streptomycin. For inhibitor treatment cells were incubated with 10 mM GM6001 as previously described [45]. All cell lines were regularly screened for mycoplasma contamination using DAPI staining.

**THP-1-conditioned media (CM)**

THP-1 cells were seeded in serum-free, antibiotic-free RPMI media at a density of 200,0000 cells/ml and incubated at 37°C and a 5% CO₂ atmosphere. After 24 hours, the cell suspension was centrifuged at 1,500 rpm at room temperature for 10 minutes. CM was either directly used for experiments (fresh
Antibodies

Anti-TIMP1 antibody (D10E6) produced in rabbit was purchased from Cell Signalling Technology, U.S.A., anti-TIMP2 antibody produced in rabbit antibody (SAB4502972) was purchased from Sigma. Anti-MT1-MMP, Anti-MMP2 and Anti-MMP9 antibody were produced in rabbit and purchased from Cell Signalling Technology, U.S.A.

Invadopodia Assay

For the invadopodia assays, the QCM™ Gelatin Invadopodia Assay (Red) (Chemicon® / Millipore) was used. Briefly, coverslips were inverted onto poly-L-lysine in deionized water for 20 minutes at room temperature (RT). The slides were then washed with PBS three times before incubation with glutaraldehyde: PBS for 15 minutes at RT. After washing three times with PBS, each coverslip was placed on gelatin in PBS in a 1:5 ratio of fluorescently-labelled - unlabelled gelatin and incubated for 10 minutes at RT and subsequently washed in PBS three times. The. Patu8902 and CFPAC-1 cells were detached using non-enzymatic Cell Dissociation Solution (Sigma Aldrich UK), resuspended in DMEM F-12 growth media (10% FBS, 1 mM penicillin/streptomycin) and seeded onto the prepared coverslips. For co-culture experiments cells were seeded in the presence of control media (serum-free RPMI media and DMEM-F12 media in a 1:1 ratio), in the presence of THP1-CM (frozen CM mixed with DMEM-F12 media with 10% FBS and 1% P/S in a 1:1 ratio) or in the presence of 50,000 THP1 cells. Alternatively, PDAC cells were co-cultured with THP1 cells for 24 hours prior to dissociation and seeding on prepared coverslips.

Spheroid assay

Spheroids were formed in black walled 96-wells clear black round bottom ultra-low attachment spheroid microplates (Corning). First, 1000 cancer cells were seeded in 200µl DMEM F-12 growth media (10% FBS, 1 mM penicillin/streptomycin). Then, the was centrifuged at 200g for 8min at room temperature. Cells were then cultured at 37°C with 5% CO2 for 72 hours. After spheroid assembly was achieved, 170ul medium was removed by multichannel pipetting. A collagen mixture was
prepared on ice with a final concentration of 1.3 mg/ml rat tail collagen I (Corning). A collagen matrix of 100µl was then added to each well with a multichannel pipette. The spheroid plate was then incubated at 37°C with 5% CO2 for 2 hrs. A 1:1 mixture of serum-free RPMI and DMEM-F12 with 10% FBS and 1 mM penicillin/streptomycin was then added on top of the collagen matrix to initiate the assay. Brightfield images were obtained with a 4x objective at 0hrs and 48hrs, respectively.

**MTT viability assay**

For the MTT assay, Patu8902 cells in 100µl full growth media (DMEM supplemented with 10% FBS and 1 mM penicillin/streptomycin) containing 4000 cells were seeded in triplicates in three 96 well plates. Plate 1 was used as a treatment-naïve control. Growth medium in plate 2 and 3 were removed from the wells and replaced with a) normal growth medium (DMEM supplemented with 10% FBS and 1 mM penicillin/streptomycin), b) DMEM supplemented with 10% FBS and 1 mM penicillin/streptomycin mixed with serum-free DMEM (1:1), c) (DMEM supplemented with 10% FBS and 1 mM penicillin/streptomycin mixed with frozen and rewarmed THP1-CM (1:1), d) DMEM supplemented with 10% FBS and 1 mM penicillin/streptomycin containing rTIMP2 (50ng/ml). The MTT assay is performed after 24 hrs (plate 1, control), 72 hrs (plate 2) and 120 hrs (plate 3).

The MTT powder was made up to a working concentration of 500μg/ml in DMEM media prior to each experiment. First, media was removed from each well. Cells were then washed with sterile PBS twice and then 1ml MTT:DMEM solution was added to the wells. Cells were incubated for four hours. The MTT:DMEM was removed and 500uL of DMSO was added, the plates were then incubated at 37°C for 10 minutes. Absorbance was measured at a wavelength of 562nm using a Nanodrop.

**Western Blot**

Serum-free THP-1 conditioned media (CM), serum-free RPMI containing recombinant TIMP1 and TIMP2 (expressed in CHO cells Sigma Aldrich UK) in various concentrations ranging from 5 to 6000 ng/ml, respectively, and serum-free RPMI control media, respectively, were filled in a Spin-X® UF concentrators (Spin-X UF 6 10K MWCO, Corning) and centrifuged at 4,000 rpm for 18 minutes at RT. Subsequently, gel sample buffer was added to the concentrated CM and the sample was boiled at 90°C for 3 min. Equal amounts of protein were electrophoresed on 10% SDS-polyacrylamide gels then
transferred to nitrocellulose membranes as described elsewhere [46]. Nitrocellulose were incubated with primary antibodies using the recommended concentrations and HRP-conjugated secondary antibodies (Dako Ltd). Cell lysates were seeded in 6-well plates and cultured for 24 hours. Lysates were generated when cells were 70-80% confluent. First, cells were washed with PBS and then lysates were generated with 100µl NP40 based lysis buffer per well. Lysates were scrapped and centrifuged at 13 000 x g for 15min at 4°C. The Supernatant was then transferred to an Eppendorf tube and boiled for 3 min at 95°C in 6x laemmli buffer. Samples were stored at -20°C. Equal amounts of protein were then electrophoresed on 7.5% SDS-polyacrylamide gels and immunoblotting was performed as described above.

**Determination of rTIMP1 and rTIMP2 concentrations**

Since there are no comparable studies using rTIMP1 and 2, respectively, in an invadopodia model, the approximate concentration had to be estimated. Physiologic TIMP1 and 2 levels, respectively, range between 5 and 1000 ng/ml [47, 48]. In a second step, CM and RPMI control media containing either TIMP1 or TIMP2 in various concentrations ranging from 5 to 6000 ng/ml was concentrated as described above and immunoblotted for TIMP1 and TIMP2, respectively. The blots of the CM and the rTIMP1 and rTIMP2 containing RPMI control media were compared by eye. TIMP1 and TIMP2 levels in the CM varied between the experiments and were ranging between 5 and 50 ng/ml. Therefore, these concentrations were used for the later rTIMP1 and rTIMP2 experiments.

**Micro Array**

THP1 CM was screened for proteins using the RayBio® C-Series Human Cytokine Antibody Array C5 (RayBiotech Norcross, USA) according to the manufacturer’s protocol.

**Immunofluorescence**

Following 24h-incubation, cells were fixed and stained as previously described. Cells were stained for F-actin (Alexa fluor 488-conjugated phalloidin,Invitrogen), DAPI (Sigma Aldrich UK) and cortactin (Anti-Cortactin (p80/85) Antibody, clone 4F11; Millipore).

**Image analysis**

Images were analysed using ImageJ 1.51h (National Institutes of Health, USA). For gelatine
degradation analysis, the total amount of degradation per image was measured in a total of 10 images per tested condition. The amount of degradation was computed automatically using ImageJ.

For the spheroid assay, the invasive front was determined as the area of all cells invading from the spheroid. The invasive front was calculated as the area of the invading cells relative to the size of the spheroid after 48 hrs. Relative spheroid growth was determined by measuring the total area of the spheroids at 0 hrs and 48 hrs, respectively.

Statistical analyses

For data collection and statistical analysis, Microsoft Excel (Microsoft, Redmond, USA) and Prism 5.0 (GraphPad Software, La Jolla, CA, USA) were used. To test for significant differences, the two-tailed Student’s t-tests was used. Data are presented as mean ± standard error of the mean (SEM). A difference was considered significant at p < 0.05.

Declarations

Ethics Approval – not relevant

Consent for publication – not relevant

Acknowledgement – not relevant

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Author Contributions

C.B. and C.M.W. planned the experiments and wrote the manuscript, C.B. and C.M.W. prepared the figures and performed the statistics, C.B., H.L., A.R., C.M.T, Y.A.B, and Y.C.P. performed the experiments, C.M.W. supervised the project. All authors interpreted the data and reviewed the manuscript

Competing interests

The authors declare no competing interests.

Data availability

The datasets generated during and/or analysed during the current study are available from the
corresponding author on reasonable request.

References

1. Oettle H, Neuhaus P, Hochhaus A, Hartmann JT, Gellert K, Ridwelski K, Niedergethmann M, Zülke C, Fahlke J, Arning MB et al: Adjuvant chemotherapy with gemcitabine and long-term outcomes among patients with resected pancreatic cancer: the CONKO-001 randomized trial. *JAMA* 2013, 310(14):1473-1481.

2. Le Large TYS, Bijlsma MF, Kazemier G, van Laarhoven HWM, Giovannetti E, Jimenez CR: Key biological processes driving metastatic spread of pancreatic cancer as identified by multi-omics studies. *Seminars in cancer biology* 2017, 44:153-169.

3. Hoshino D, Branch KM, Weaver AM: Signaling inputs to invadopodia and podosomes. *J Cell Sci* 2013, 126(Pt 14):2979-2989.

4. Stoletov K, Lewis JD: Invadopodia: a new therapeutic target to block cancer metastasis. *Expert review of anticancer therapy* 2015, 15(7):733-735.

5. Gimona M, Buccione R, Courtneidge SA, Linder S: Assembly and biological role of podosomes and invadopodia. *Current opinion in cell biology* 2008, 20(2):235-241.

6. Egeblad M, Werb Z: New functions for the matrix metalloproteinases in cancer progression. *Nature reviews Cancer* 2002, 2(3):161-174.

7. Jacob A, Prekeris R: The regulation of MMP targeting to invadopodia during cancer metastasis. *Frontiers in cell and developmental biology* 2015, 3:4.

8. Clark ES, Whigham AS, Yarbrough WG, Weaver AM: Cortactin is an essential regulator of matrix metalloproteinase secretion and extracellular matrix degradation in invadopodia. *Cancer research* 2007, 67(9):4227-4235.

9. Fortunato SJ, Menon R, Lombardi SJ: Presence of four tissue inhibitors of matrix
metalloproteinases (TIMP-1, -2, -3 and -4) in human fetal membranes. American journal of reproductive immunology 1998, 40(6):395-400.

10. Jackson HW, Defamie V, Waterhouse P, Khokha R: TIMPs: versatile extracellular regulators in cancer. Nature reviews Cancer 2017, 17(1):38-53.

11. Lang R, Braun M, Sounni NE, Noel A, Frankenne F, Foidart JM, Bode W, Maskos K: Crystal structure of the catalytic domain of MMP-16/MT3-MMP: characterization of MT-MMP specific features. Journal of molecular biology 2004, 336(1):213-225.

12. Zucker S, Drews M, Conner C, Foda HD, DeClerck YA, Langley KE, Bahou WF, Docherty AJ, Cao J: Tissue inhibitor of metalloproteinase-2 (TIMP-2) binds to the catalytic domain of the cell surface receptor, membrane type 1-matrix metalloproteinase 1 (MT1-MMP). The Journal of biological chemistry 1998, 273(2):1216-1222.

13. Artym VV, Zhang Y, Seillier-Moiseiwitsch F, Yamada KM, Mueller SC: Dynamic interactions of cortactin and membrane type 1 matrix metalloproteinase at invadopodia: defining the stages of invadopodia formation and function. Cancer research 2006, 66(6):3034-3043.

14. Gould CM, Courtneidge SA: Regulation of invadopodia by the tumor microenvironment. Cell adhesion & migration 2014, 8(3):226-235.

15. Xie D, Xie K: Pancreatic cancer stromal biology and therapy. Genes & diseases 2015, 2(2):133-143.

16. Komura T, Sakai Y, Harada K, Kawaguchi K, Takabatake H, Kitagawa H, Wada T, Honda M, Ohta T, Nakanuma Y et al: Inflammatory features of pancreatic cancer highlighted by monocytes/macrophages and CD4+ T cells with clinical impact. Cancer Science 2015, 106(6):672-686.
17. Cui R, Yue W, Lattime EC, Stein MN, Xu Q, Tan XL: Targeting tumor-associated macrophages to combat pancreatic cancer. Oncotarget 2016, 7(31):50735-50754.

18. Wang Z, Liang X, Cai M, Du G: Analysis of invadopodia formation in breast cancer cells. Methods in molecular biology (Clifton, NJ) 2016, 1406:203-210.

19. Paz H, Pathak N, Yang J: Invading one step at a time: the role of invadopodia in tumor metastasis. Oncogene 2014, 33(33):4193-4202.

20. Neel NF, Rossman KL, Martin TD, Hayes TK, Yeh JJ, Der CJ: The RalB small GTPase mediates formation of invadopodia through a GTPase-activating protein-independent function of the RalBP1/RLIP76 effector. Molecular and cellular biology 2012, 32(8):1374-1386.

21. Singh SK, Mishra MK, Eltoum IA, Bae S, Lillard JW, Jr., Singh R: CCR5/CCL5 axis interaction promotes migratory and invasiveness of pancreatic cancer cells. Sci Rep 2018, 8(1):1323.

22. Revach O-Y, Weiner A, Rechav K, Sabanay I, Livne A, Geiger B: Mechanical interplay between invadopodia and the nucleus in cultured cancer cells. Scientific Reports 2015, 5:9466.

23. Gil-Henn H, Patsialou A, Wang Y, Warren MS, Condeelis JS, Koleske AJ: Arg/Abl2 promotes invasion and attenuates proliferation of breast cancer in vivo. Oncogene 2013, 32(21):2622-2630.

24. Quattrochi B, Gulvady A, Driscoll DR, Sano M, Klimstra DS, Turner CE, Lewis BC: MicroRNAs of the mir-17~92 cluster regulate multiple aspects of pancreatic tumor development and progression. Oncotarget 2017, 8(22):35902-35918.

25. Siret C, Dobric A, Martirosyan A, Terciolo C, Germain S, Bonier R, Dirami T, Dusetti N, Tomasini R, Rubis M et al: Cadherin-1 and cadherin-3 cooperation determines
the aggressiveness of pancreatic ductal adenocarcinoma. *Br J Cancer* 2018, 118(4):546-557.

26. Wang Z, Liang X, Cai M, Du G: **Analysis of Invadopodia Formation in Breast Cancer Cells.** *Methods in molecular biology (Clifton, NJ)* 2016, 1406:203-210.

27. Meirson T, Genna A, Lukic N, Makhnii T, Alter J, Sharma VP, Wang Y, Samson AO, Condeelis JS, Gil-Henn H: **Targeting invadopodia-mediated breast cancer metastasis by using ABL kinase inhibitors.** *Oncotarget* 2018, 9(31):22158-22183.

28. Chevalier C, Cannet A, Descamps S, Sirvent A, Simon V, Roche S, Benistant C: **ABL tyrosine kinase inhibition variable effects on the invasive properties of different triple negative breast cancer cell lines.** *PloS one* 2015, 10(3):e0118854.

29. Beghein E, Devriese D, Van Hoey E, Gettemans J: **Cortactin and fascin-1 regulate extracellular vesicle release by controlling endosomal trafficking or invadopodia formation and function.** *Scientific reports* 2018, 8(1):15606.

30. Mytar B, Baj-Krzyworzeka M, Majka M, Stankiewicz D, Zembala M: **Human monocytes both enhance and inhibit the growth of human pancreatic cancer in SCID mice.** *Anticancer Res* 2008, 28(1a):187-192.

31. Helm O, Held-Feindt J, Grage-Griegenow E, Reiling N, Ungefroren H, Vogel I, Kruger U, Becker T, Ebsen M, Rocken C et al: **Tumor-associated macrophages exhibit pro-and anti-inflammatory properties by which they impact on pancreatic tumorigenesis.** *International journal of cancer* 2014, 135(4):843-861.

32. Wang X, Luo G, Zhang K, Cao J, Huang C, Jiang T, Liu B, Su L, Qiu Z: **Hypoxic tumor-derived exosomal miR-301a mediates M2 macrophage polarization via PTEN/PI3Kgamma to promote pancreatic cancer metastasis.** *Cancer Res* 2018.

33. Farajzadeh Valilou S, Keshavarz-Fathi M, Silvestris N, Argentiero A, Rezaei N: **The
role of inflammatory cytokines and tumor associated macrophages (TAMs) in microenvironment of pancreatic cancer. Cytokine Growth Factor Rev 2018, 39:46-61.

34. Starnes TW, Cortesio CL, Huttenlocher A: Imaging podosome dynamics and matrix degradation. Methods in molecular biology (Clifton, NJ) 2011, 769:111-136.

35. Clark ES, Weaver AM: A new role for cortactin in invadopodia: regulation of protease secretion. European journal of cell biology 2008, 87(8-9):581-590.

36. Lebeau A, Nerlich AG, Sauer U, Lichtinghagen R, Lohrs U: Tissue distribution of major matrix metalloproteinases and their transcripts in human breast carcinomas. Anticancer Res 1999, 19(5b):4257-4264.

37. Toth M, Chvyrkova I, Bernardo MM, Hernandez-Barrantes S, Fridman R: Pro-MMP-9 activation by the MT1-MMP/MMP-2 axis and MMP-3: role of TIMP-2 and plasma membranes. Biochem Biophys Res Commun 2003, 308(2):386-395.

38. Itoh Y: Membrane-type matrix metalloproteinases: Their functions and regulations. Matrix Biology 2015, 44-46:207-223.

39. Lim SY, Yuzhalin AE, Gordon-Weeks AN, Muschel RJ: Tumor-infiltrating monocytes/macrophages promote tumor invasion and migration by upregulating S100A8 and S100A9 expression in cancer cells. Oncogene 2016, 35:5735.

40. Kai AK, Chan LK, Lo RC, Lee JM, Wong CC, Wong JC, Ng IO: Down-regulation of TIMP2 by HIF-1alpha/miR-210/HIF-3alpha regulatory feedback circuit enhances cancer metastasis in hepatocellular carcinoma. Hepatology 2016, 64(2):473-487.

41. Winograd R, Byrne KT, Evans RA, Odorizzi PM, Meyer AR, Bajor DL, Clendenin C, Stanger BZ, Furth EE, Wherry EJ et al: Induction of T-cell Immunity Overcomes
Complete Resistance to PD-1 and CTLA-4 Blockade and Improves Survival in Pancreatic Carcinoma. *Cancer Immunol Res* 2015, **3**(4):399-411.

42. Guo S, Contratto M, Miller G, Leichman L, Wu J: *Immunotherapy in pancreatic cancer: Unleash its potential through novel combinations*. *World Journal of Clinical Oncology* 2017, **8**(3):230-240.

43. Morishita A, Gerber A, Gow CH, Zelonina T, Chada K, D'Armiento J: *Cell Specific Matrix Metalloproteinase-1 Regulates Lung Metastasis Synergistically with Smoke Exposure*. *J Cancer Res Forecast* 2018, **1**(2):1014.

44. Ramasamy K, Khatun H, Macpherson L, Caley MP, Sturje J, Mufti GJ, Schey SA, Calle Y: *Fluorescence-based experimental model to evaluate the concomitant effect of drugs on the tumour microenvironment and cancer cells*. *Br J Haematol* 2012, **157**(5):564-579.

45. Kikuchi K, Takahashi K: *WAVE2- and microtubule-dependent formation of long protrusions and invasion of cancer cells cultured on three-dimensional extracellular matrices*. *Cancer Sci* 2008, **99**(11):2252-2259.

46. Wells CM, Abo A, Ridley AJ: *PAK4 is activated via PI3K in HGF-stimulated epithelial cells*. *J Cell Sci* 2002, **115**(Pt 20):3947-3956.

47. Shankavaram UT, Lai WC, Netzel-Arnett S, Mangan PR, Ardans JA, Caterina N, Stetler-Stevenson WG, Birkedal-Hansen H, Wahl LM: *Monocyte membrane type 1-metalloproteinase. Prostaglandin-dependent regulation and role in metalloproteinase-2 activation*. *The Journal of biological chemistry* 2001, **276**(22):19027-19032.

48. Tandara AA, Mustoe TA: *MMP- and TIMP-secretion by human cutaneous keratinocytes and fibroblasts--impact of coculture and hydration*. *J Plast Reconstr Aesthet Surg* 2011, **64**(1):108-116.
Table I: Tumor characteristics and invadopodia formation\(^1\) of tested cell lines

| Cell line    | Tumor entity | Source of tumour cells | Invadopodia formation |
|--------------|--------------|-------------------------|-----------------------|
| PaTu8902     | PDAC         | Primary tumor           | yes                   |
| CFPAC-1      | PDAC         | Liver metastasis        | yes                   |
| MiaPaCa-2    | PDAC         | Primary tumor           | inconsistent          |
| CAPAN-2      | PDAC         | Primary tumor           | inconsistent          |
| Panc-1       | PDAC         | Primary tumor           | inconsistent          |
| CAPAN-1      | PDAC         | Liver metastasis        | no                    |
| PaTu8988-T   | PDAC         | Liver metastasis        | no                    |
| PaTu8988-S   | PDAC         | Liver metastasis        | no                    |
| Colo357      | PDAC         | Lymph node metastasis   | no                    |
| AsPC1        | PDAC         | Ascites                 | no                    |
| MDA-MB-231\(^2\) | Breast cancer | Pleural effusion     | yes                   |

PDAC = Pancreatic Ductal Adenocarcinoma

\(^1\) Active invadopodia formation was defined as cortactin puncta corresponding with black dots on the gelatin.

\(^2\) MDA-MB-231 cells are well known for producing robust invadopodia formation and were used as a positive control.

Table II: MTT assay results of PaTu8902 cells with different treatment conditions

| Treatment                  | Mean % gain in growth | Standard deviation |
|----------------------------|-----------------------|--------------------|
| G                          | 52                    | 12                 |
| G+SF                       | 52                    | 14                 |
| G+CM                       | 52                    | 12                 |
| G+rTIMP2 (50ng/ml)         | 56                    | 16                 |

G = DMEM supplemented with 10% v/v fetal bovine serum (FBS) and 1 mM penicillin/streptomycin. SF=serum-free DMEM, CM = THP1 conditioned medium, rTIMP2 = recombinant tissue inhibitor of metalloproteinases. ANOVA of raw data reported no significant difference between any of the groups.
A

Invadopodia – PaTu8902 after 24 hrs
Cortactin Gelatin
Invadopodia – CFPAC-1 after 24 hrs
Cortactin Gelatin

B

Degradation in \( \mu \text{m}^2 \) field

C

CFPAC-1 PaTu8902 Patu8988-S
GAPDH
MTMMP1

D

CFPAC-1 PaTu8902
GAPDH MMP9

E

CFPAC-1 PaTu8902
GAPDH TIMP2
PaTu8902 and CFPAC-1 cells exhibit robust invadopodia activity that by expressing MT1-MMP and MMP9.

A) Pancreatic ductal adenocarcinoma (PDAC) cell lines were analysed for seeded on fluorescent gelatin for 24 hours fixed and stained for cortactin and DAPI.

Cortactin puncta corresponding with black dots on the gelatin were considered active invadopodia.

B) Quantification of degradation per field of view for (A), MDA-MB-231 cells were also seeded on fluorescent gelatin for 24 hours and the degradation per field of view calculated as a positive control.

C) PaTu8902, CFPAC-1 and PaTu8988-S cell lysates were blotted for MT1-MMP. PaTu8902, CFPAC-1 show high levels of MT1-MMP expression whereas PaTu8988-S shows no MT1-MMP expression.

D) Western blot of PaTu8902 and CFPAC-1 cell lysates showing high levels of MMP9 expression and no expression of MMP2 (data not shown).

E) Western blot of PaTu8902 and CFPAC-1: no expression of TIMP2.
Invadopodia formation can be suppressed by co-culturing PaTu8902 and CFPAC-1 cells with monocyte-like cells. A) Representative images from a PaTu8902 invadopodia assay where the cells were either incubated with control medium (control), incubated with THP1 conditioned medium (THP1-CM), or were cultured with THP1 cells during the invadopodia assay (THP1-DC), or cultured with THP1 cells prior to the invadopodia assay (THP1-PC). For the THP1-PC condition, growth medium (GM) containing the THP1 cells was evacuated and washed with PBS. B) Quantification of degradation per field of view for experimental conditions described above (A). C) Representative images from a CFPAC-1 invadopodia assay where the cells were either incubated with control medium (control), incubated with THP1 conditioned medium (THP1-CM), or were cultured with THP1 cells during the invadopodia assay (THP1-DC), or cultured with THP1 cells prior to the invadopodia assay (THP1-PC). D) Quantification of degradation per field of view for experimental conditions described above (C). E) Representative images from a PaTu8902 spheroid assay where the cells were either incubated with control medium (control) or incubated with THP1 conditioned medium (THP1-CM) at 0hrs and 48 hrs. F) Quantification of spheroid invasion for experimental conditions described above (E). G) Quantification of relative spheroid growth after 48 hrs for experimental conditions described above (E). H) Representative images from a Panc1 spheroid assay where the cells were either incubated with control medium (control) or incubated with THP1 conditioned medium (THP1-CM) at 0hrs and 48 hrs. I) Quantification of spheroid invasion for experimental conditions described above (H). J) Quantification of relative spheroid growth after 48 hrs for experimental conditions described above (I). In all cases **** = p < 0.0001, *** = p < 0.001, ** = p < 0.01, * = p < 0.05.

Experiments were repeated three times.
THP1 cells secrete TIMP1 and TIMP2  A) Representative images from a PaTu8902 invadopodia assay where the cells were either incubated with control medium, conditioned medium that had been applied fresh, boiled or freeze/thawed as indicated. B) Quantification of degradation per field of view for experimental conditions described above (A). C) Representative images from a CFPAC-1 invadopodia assay where the cells were either incubated with control medium, conditioned medium that had been applied fresh, boiled or freeze/thawed as indicated. D) Quantification of degradation per field of view for experimental conditions described above (C). E) THP1 conditioned media protein micro array (see methods for details). F) concentrated samples of THP1 CM and RPMI 1640 control were immunoblotted for TIMP1 using specific antibodies. G) control media containing recombinant TIMP1 was immunoblotted in comparison to aliquots of concentrated samples of THP1 CM for TIMP1 using specific antibodies. H) concentrated samples of THP1 CM and RPMI 1640 control were immunoblotted for TIMP2 using specific antibodies. I) control media containing recombinant TIMP2 was immunoblotted in comparison to a concentrated sample of THP1 CM for TIMP2 using specific antibodies. Experiments were repeated three times.
Figure 4

TIMP2 inhibits invadopodia formation in PDAC cells. A) Representative images from a PaTu8902 invadopodia assay where the cells were either incubated with THP1 conditioned medium (THP1-CM), or were cultured with recombinant TIMP1 (rTIMP1) or recombinant TIMP2 (rTIMP2) at indicated concentrations B and C) Quantification of degradation per field of view for experimental conditions described above (A). D) Representative images from a CFPAC-1 invadopodia assay where the cells were either incubated with THP1 conditioned medium (THP1-CM), or were cultured with recombinant TIMP1 (rTIMP1) or recombinant TIMP2 (rTIMP2) at indicated concentrations E and F) Quantification of degradation per field of view for experimental conditions described above (D). In all cases **** = p < 0.0001, ** = p < 0.01, *** = p < 0.001. Experiments were repeated three times.
Figure 5

TIMP2-mediated inhibition of invadopodia is reversible after washout in pancreatic cancer cells A) Representative images from a PaTu8902 invadopodia assay where the cells were either incubated for 48 hrs with control medium, control medium with GM6001 (10µM) washed out after 5 hrs and replaced with fresh control media or GM6001 (10µM) left for 48 hrs. B) Quantification of degradation per field of view for experimental conditions described above (A). C) Representative images from a PaTu8902 invadopodia assay where the cells were either incubated for 48 hrs with control medium, control medium with recombinant TIMP2 (rTIMP2, 50 ng/ml) washed out after 5 hrs and replaced with fresh control medium or control medium with rTIMP2 (50 ng/ml) left for 48 hrs. D) Quantification of degradation per field of view for experimental conditions described above (C). There was no significant difference in the amount of gelatin degradation between the control cells and the cells where rTIMP2 had been removed post 5 hours incubation indicating that PaTu8902 cells were able to recover invadopodia activity once rTIMP2 had been removed. In contrast, again, those cells continuously exposed to rTIMP2 revealed significantly reduced invadopodia formation. E) Representative images from a CFPAC-1 invadopodia assay where the cells were either incubated for 48 hrs with control medium, control medium with GM6001 (10µM) washed out after 5 hrs and replaced with fresh control media or GM6001 (10µM) left for 48 hrs. F) Quantification of degradation per field of view for experimental conditions described above (E). G) Representative images from a CFPAC-1 invadopodia assay where the cells were either incubated for 48 hrs with control medium, control medium with recombinant TIMP2 (rTIMP2, 50 ng/ml) washed out after 5 hrs and replaced with fresh control media or rTIMP2 (50 ng/ml) left for 48 hrs. H) Quantification of degradation per field of view for experimental conditions described above In all cases **** = p < 0.0001, *** = p < 0.001, ** = p < 0.01. Experiments were repeated three times.