Mapping Proteolytic Processing in the Secretome of Gastric Cancer-Associated Myofibroblasts Reveals Activation of MMP-1, MMP-2, and MMP-3

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ABSTRACT: Cancer progression involves changes in extracellular proteolysis, but the contribution of stromal cell secretomes to the cancer degradome remains uncertain. We have now defined the secretome of a specific stromal cell type, the myofibroblast, in gastric cancer and its modification by proteolysis. SILAC labeling and COFRADIC isolation of methionine containing peptides allowed us to quantify differences in gastric cancer-derived myofibroblasts compared with myofibroblasts from adjacent tissue, revealing increased abundance of several proteases in cancer myofibroblasts including matrix metalloproteinases (MMP)-1 and -3. Moreover, N-terminal COFRADIC analysis identified cancer-restricted proteolytic cleavages, including liberation of the active forms of MMP-1, -2, and -3 from their inactive precursors. In vivo imaging confirmed increased MMP activity when gastric cancer cells were xenografted in mice together with gastric cancer myofibroblasts. Western blot and enzyme activity assays confirmed increased MMP-1, -2, and -3 activity in cancer myofibroblasts, and cancer cell migration assays indicated stimulation by MMP-1, -2, and -3 in cancer-associated myofibroblast media. Thus, cancer-derived myofibroblasts differ from their normal counterparts by increased production and activation of MMP-1, -2, and -3, and this may contribute to the remodelling of the cancer cell microenvironment.

KEYWORDS: myofibroblast, secretome, COFRADIC, degradome, neo-N-termini

INTRODUCTION

Elucidation of the dynamic changes in secretomes, i.e., the secreted subset of the proteome, underlies a systems approach to understanding the mechanisms controlling cell–cell and cell–matrix interactions in health and disease. In cancer it is now clear that changes in the cellular microenvironment determine disease progression, and that these include two-way interactions between cancer cells and surrounding stromal cells.1−3

Several recent studies have employed proteomic techniques to define the secretomes of cancer cells.4−10 Recent studies have also defined the secretome in breast and colon cancer fibroblasts,11,12 but in general the interrogation of stromal cell secretomes by proteomic methods has been limited. Changes in secretomes may reflect alterations in secretory protein abundance due to variations in gene expression, rates of exocytosis or presecretory post-translational processing. But in addition, there may also be extensive postsecretory proteolysis that defines the extracellular tumor degradome. In cancer, alterations in extracellular proteolysis through differential secretion of proteases or their inhibitors is important, not least because it underlies disease progression and sensitivity to protease targeted therapies.13−15

Cancer-associated fibroblasts, of which myofibroblasts are a subset, are important stromal cells that exhibit an altered phenotype in many cancers.16−19 Myofibroblasts play important roles in defining the tissue microenvironment through secretion of extracellular matrix components, growth factors, proteases and their inhibitors.20,21 Differences between normal and cancer-associated myofibroblasts (CAMs) have been linked to tumor progression by mechanisms including recruitment from circulating mesenchymal stromal cells, global DNA hypomethylation and changes in gene expression profiles.22−24
Since myofibroblasts stimulate cancer cell invasion, in the present study we sought to define the differences between gastric CAMs compared with adjacent tissue-derived myofibroblasts (ATMs) with respect to proteolytic processing of their secretomes. The data indicate both upregulation and activation of matrix metalloproteinases (MMPs)-1, -2, and -3 are characteristic of the CAM secretome.

MATERIALS AND METHODS

Human Primary Myofibroblasts

Myofibroblasts from gastric cancers (CAMs) and adjacent tissue (ATMs) from two patients have been described previously (Supporting Information). The work was approved by the Ethics Committee of the University of Szeged, Hungary, and subjects provided informed consent.

SILAC Labeling

Myofibroblasts were cultured in DMEM SILAC media (Pierce, Thermo Scientific, Rockford, IL, USA) for 6 population doublings in the presence of either natural (light) or heavy $^{13}$C$_6$-labeled l-arginine (0.94 mM) and $^{13}$C$_6$ l-lysine (0.46 mM) (Invitrogen, Paisley, Renfrew, U.K.). Media were further supplemented with 10% dialyzed fetal bovine serum (Pierce), 2% antibiotic/antimycotic (Sigma-Aldrich, Poole, U.K.) and 1% penicillin/streptomycin (Sigma).

Sample Preparation

Media (10 mL, serum-free) obtained from 1 × 10$^6$ myofibroblasts plated in 10 cm diameter dishes (80–90% confluency) were collected after 24 h. Samples were concentrated to approximately 0.5 mL using Amicon Ultra-15 3 kDa centrifugal filter devices (Millipore, Watford, U.K.), precipitated with 20% TCA and resuspended in 50 mM HEPES, pH 7.4, 100 mM NaCl, 0.8% w/v CHAPS, 1% v/v Protease Inhibitor Cocktail Set III, EDTA-Free (Calbiochem, Merck Biosciences, Beeston, U.K.). Equal amounts (35 μg each) of light and heavy SILAC-labeled secretome samples from CAMs and ATMs were mixed following determination of protein concentration by the Bradford assay (Bio-Rad Lab, Inc., Hemel Hempstead, U.K.).

COFRADIC Isolation of Methionine Containing Peptides

Methionyl-COFRADIC was performed as described previously (see Supporting Information Methods). Samples were reduced and S-alkylated, and following trypsinization (trypsinːprotein 1ː100), peptides were fractionated by reversed-phase HPLC (RP-HPLC) using an Agilent 1100 HPLC system with a Zorbax 300SB-C$_{18}$ column (2.1 mm (internal diameter) × 150 mm, Agilent Technologies, Wokingham, U.K.). The resulting HPLC fractions were further processed by incubating for 30 min with 0.1% w/v hydrogen peroxide at 30 °C. Following oxidation of methionines, reaction mixtures were immediately reinjected onto the RP-HPLC column for secondary RP-HPLC separations under identical conditions. Fractions with methionine containing peptides displayed a hydrophilic shift and were collected ($n = 90$) and analyzed by LC−MS/MS.

COFRADIC Isolation of N-Terminal Peptides

N-terminal COFRADIC was performed as described previously (see Supporting Information Methods). Proteins were reduced and alkylated, and primary α- and ε-amines were blocked by tri.deutero-acetylation. Samples were then trypsinized, and N-terminal peptides were pre-enriched by strong cation exchange chromatography at low pH. Following a pyroglutamate removal step, peptides were separated by RP-HPLC as described above. Primary fractions were incubated with 2,4,6-trinitrobenzenesulphonic acid (TNBS) to modify internal tryptic peptides with free α-N-termini. A series of secondary RP-HPLC runs was then performed on each individual primary fraction, and N-terminal peptides (which did not display a hydrophobic shift) were collected ($n = 36$) for LC−MS/MS analysis.

Non-COFRADIC Experiments

Samples prepared for “shotgun” analysis of the secretomes were processed following the same method as for Met-COFRADIC up to the stage immediately before the first RP-HPLC run. At this point the sample was processed for LC−MS/MS analysis (see Supporting Information Methods).

For neo-N-terminal enrichment, the non-COFRADIC method employed a SCX-only enrichment of N-terminal peptides. Samples were prepared as for N-terminal COFRADIC up to the first RP-HPLC run. At this stage 60 fractions of 1 min interval were collected, pooled to give a total of 20 fractions, dried and prepared for LC−MS/MS analysis.

LC−MS/MS Analysis and Peptide Identification by Mascot

Peptides were analyzed using a LTQ Orbitrap XL mass spectrometer (Thermo Electron, Bremen, Germany) as described previously. Mascot generic files (mgf) were created using the Mascot Distiller software (version 2.2.1.0, Matrix Science, Ltd., London, U.K.). When generating peak lists, grouping of spectra was performed with a maximum intermediate retention time of 30 s and maximum intermediate scan count of 5. Grouping was further done with 0.1 Da precursor ion tolerance. A peak list was only generated when the spectrum contained more than 10 peaks. There was no desalting, and the relative signal-to-noise limit for both precursor and fragment ions was set to 2. The peak lists were then searched with Mascot using the Mascot Daemon interface (version 2.2.0, Matrix Science, Ltd.) against human proteins in the Swiss-Prot database (Uniprot release 15.0, containing 20 333 human protein sequences). Spectra were searched with semiArgC/P enzyme settings, allowing no missed cleavages for the N-terminal peptide experiments, and with trypsin/P settings allowing no missed cleavages for the Met-COFRADIC/shotgun experiments. Mass tolerance of the precursor ions was set to 10 ppm (with Mascot’s C13 option set to 1) and of fragment ions to 0.5 Da. The instrument setting was ESI-TRAP. Variable modifications were acetylation of alpha-N-termini and pyroglutamate formation of N-terminal glutamine residues; fixed modification was oxidation of methionine (sulfoxide). Additionally, for N-terminal peptide experiments, tri.deutero-acetylation of the N-terminus was set as variable peptide modification, and tri.deutero-acetylation of lysine side chains was included as fixed modification. Only peptides that were ranked one and scored above the identity threshold score set at 99% confidence were withheld. The FDR was calculated for every search as described previously (see Supporting Information Methods, Table SM3). Identified peptides were quantified using the Mascot Distiller Quantitation Toolbox (www.matrixscience.com) in the "precursor" mode as described previously. Ratios for all peptides of interest were validated by manual inspection of spectra. For processing of all MS data, the ms_lims software platform was used. Protein ratios were inferred using the mean of the peptide group ratios for each protein. A peptide group represents all quantifications of a single peptide sequence in an experiment. The distribution of
protein ratios as determined by the Met-COFRADIC and shotgun experiments was plotted using Rover.31 This was used to define thresholds to give the 5% of proteins with the largest fold changes in CAMs relative to ATMs. All spectra have been stored in the PRIDE database (http://www.ebi.ac.uk/pride/, accession numbers 27157−27161) using PRIDE converter.32

Protein subcellular localizations and functional classifications were manually curated, using the UniProt and HPRD online databases.

In Vivo Imaging

Immunocompromised mice (6 weeks old, BALB/c nu/nu, Charles River, Wilmington, MA) with xenografts of MKN45 cells with or without CAMs (Supporting Information Methods) were used for imaging MMP-activity using MMPSense 750 FAST. These experiments were approved by the University of Liverpool Animal Welfare Committee and were conducted in compliance with the U.K. Animals (Scientific Procedures) Act 1986.

Western Blot

Myofibroblast cell extracts were prepared in RIPA buffer containing protease and phosphatase inhibitors. Cell extracts or media were resolved by SDS-PAGE and processed for Western blotting as previously described.33 Blots were probed with antibodies against MMP-1 (BAF901, R&D Systems, Minneapolis, MN, USA), MMP-2 (BAF902, R&D Systems) and MMP-3 (BAF513, R&D Systems). Membranes were reprobed with anti-GAPDH antibody (Biodesign, Saco, Maine, USA) for assessing equal loading where appropriate.

Enzyme Assays

Fluorogenic assays for MMP enzyme activity were performed using selective substrates: DNP-Pro-Leu-Ala-Leu-Trp-Ala-Arg-OH (MMP-1; Calbiochem, Bedfont Cross, U.K.), DNP-Pro-Leu-Gly-Met-Trp-Ser-Arg-OH (MMP-2/9; Calbiochem), MCA-Pro-Leu-Nva-Dpa-Ala-Arg-NH₂ (MMP-2; Calbiochem), DNP-Pro-Tyr-Ala-Tyr-Trp-Met-Arg-OH (MMP-3; AnaSpec, Fremont, CA, USA) and 5-FAM-Arg-Pro-Lys-Pro-Val-Glu-Nva-Trp-Arg-Lys(TQ2W)-NH₂ (MMP-3; Enzo Life Sciences, Exeter, U.K.) as previously described.33,34 Briefly, equal volumes of assay buffer and media from 10⁵ myofibroblasts were incubated with 12 μM substrate as appropriate.
Table 1. Neo-N-terminal Peptides of Extracellular Proteins Unique to CAMs (i.e., Singletons), Excluding Those Representing Removal of a Signal Sequence

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<th>peptide</th>
<th>patient 1</th>
<th>patient 2</th>
<th>patient 1 (N-terminal COFRADIC)</th>
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<td>Extracellular domain</td>
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<td>Nonhelical region</td>
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<td>Alpha-1 region</td>
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<td>SLKKEECPAGSHR</td>
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"Functional classification and cleavage sites were manually annotated using Uniprot database. Start and end position number of the N-terminally labeled peptide are relative to unprocessed forms of the protein."
RESULTS

Different Secretomes in Cancer-Derived and Adjacent Tissue Myofibroblasts

When myofibroblasts derived from two gastric cancers and corresponding ATMs were SILAC labeled and the secretomes analyzed by LC−MS/MS, approximately 350 unique proteins (310 and 392) were identified in each pair on the basis of one or more peptide unique assignments with validated quantification (Supporting Information Tables S1, S2). Of these, 42 and 48% were characterized as extracellular proteins on the basis of annotations in the UniProt and HPRD databases. One of the paired samples was further analyzed using the COFRADIC technology that enriches for methionine-containing peptides in an attempt to increase overall proteome coverage by reducing sample complexity. This approach more than doubled the number of unique peptides and proteins identified in the secretome, although a comparable proportion (31%) of the identifications were attributable to extracellular proteins (Figure 1A; Supporting Information Table S3). The Met-COFRADIC analysis identified the majority (72%) of proteins identified in the initial experiments (Figure 1A). In total across the three experiments, 1460 unique proteins were identified, of which 364 were annotated to be extracellular proteins.

Extracellular proteins making up the secretome were grouped into 11 functional classifications (Figure 1B). There was a broadly similar distribution across these groups in both pairs of CAMs and ATMs, with binding proteins and extracellular matrix proteins the two largest groups (Figure 1B). When 95% confidence limits were determined and applied to individual experiments, 9−47 proteins were identified that were above or below these limits (Supporting Information Figure S1; Supporting Information Table S4). Interestingly, MMP-1, MMP-3, MMP-10 and uPA were identified as upregulated in CAMs in both patients (Supporting Information Table S4). The largest groups showing differential abundance were “binding” proteins, e.g., the insulin-like growth factor binding proteins (IGFBPs), “receptor” proteins, e.g., epidermal growth factor receptor, and proteases, e.g., MMP-1 (Supporting Information Table S4).

Proteolytic Processing of the Secretome

We then extended the analysis to the identification of neo-N-termi generated as a consequence of proteolytic cleavage. Thus, SILAC-labeled media samples were enriched for N-terminal peptides using a strong cation exchange (SCX) step to remove nonterminal peptides. As part of this procedure, neo-N-terminal peptides had been trideutero-acetylated prior to mass spectrometry, and the results were filtered so that only peptides that were trideutero-acetylated and had a valid Mascot identification were used in subsequent analyses. Of unique quantified peptides, 20−42% were trideutero-acetylated (Supporting Information Table S5). This process identified peptides starting at residues 1 or 2 of the protein or immediately after removal of the signal sequence, which were excluded from further analysis as they were considered uninformative for present purposes. For each of the remaining peptides, the ratio of relative abundance between the CAM and ATM samples was manually validated by inspecting the spectra and calculating the area under the peaks of the heavy and light isotopes.

One of the paired samples was further analyzed using N-terminal COFRADIC in addition to the SCX enrichment step in an attempt to increase coverage of N-terminal peptides by reducing sample complexity. Similar numbers of peptides were identified by N-terminal COFRADIC and SCX-only enrichment, but over 2-fold more of these identifications

Figure 2. Increased MMP activity in vivo in xenografts containing CAMs. (A) In BALB/c nu/nu mice, MMP activity revealed by FMT imaging of MMPSense 750 FAST is increased in xenografts of gastric cancer MKN45 cells containing two different CAMs compared with MKN45 cells alone. (B) Representative FMT images of mice with xenografts of MKN45 cells alone (top row), MKN45 cells + CAM2 (middle row), and MKN45 cells + CAM1 (bottom row).
corresponded to neo-N-termini in the COFRADIC data set (Supporting Information Table S5). Approximately 50% of all unique proteins identified corresponded to putative secreted proteins (Supporting Information Table S6). In the functional classification, “binding proteins” and “ECM” proteins again predominated (Figure 1B). Furthermore, 41 proteins were identified that were not seen in the first set of SILAC experiments, of which 24 were known extracellular proteins.

CAM-Restricted Proteolytic Events

In the data set as a whole, neo-N-termini corresponding to putative proteolytic cleavage sites were identified in a total of 94 unique proteins, of which collagens alpha1(I) and alpha-2(I) and IGFBP-5 had the most cleavage sites (Supporting Information Table S7). In order to refine cancer-related changes, we then sought those proteins for which unique neo-N-termini were identified in CAM secretomes relative to their ATM counterparts. Applying this criterion, we identified 13 proteins that exhibited CAM-restricted proteolytic cleavage (Table 1): strikingly, these included cleavages in the propeptide domains of MMP-1 (interstitial collagenase) and MMP-3 (stromelysin-1) (Table 1 and Supporting Information Figure S2); moreover, N-terminal COFRADIC also identified increased prodomain cleavage in MMP-2 (72 kDa type IV collagenase) (Supporting Information Table S7).

In Vivo MMP Activity

To determine the in vivo relevance of myofibroblasts for MMP activity in tumors, we made use of FMT imaging of a MMP fluorescent substrate in a xenograft model consisting of MKN45 gastric cancer cells alone or with human CAMs. In matched tumors of similar size generated with and without coadministration of CAMs, the MMPSense substrate revealed significantly increased activity when human CAMs were present (Figure 2).

Activation of MMP-1, -2, and -3 in CAM Media

The substrate used for in vivo imaging does not distinguish individual MMPs, and subsequent studies therefore made use of in vitro techniques to assess the relative contribution of
MMPs1−3 in CAM media. Western blots of MMP-1 revealed increased abundance of a 42 kDa form corresponding to the active enzyme in CAM compared with ATM media; in contrast, in cell extracts of the two cell types there was similar abundance of the precursor proteins of 57 and 52 kDa (Figure 3A). For MMP-2, we identified a precursor form of 72 kD in cells and media of both CAMs and ATMs, again in similar abundance in the two cell types. When blots of media were overexposed it was possible to identify a minor band of 63 kDa corresponding to the active enzyme in CAM but not ATM media (Figure 3B).

Finally, for MMP-3 we identified precursor proteins of 59 and 54 kDa in cells and media of both CAMs and ATMs; a band of 45 kDa corresponding to the active enzyme was found in media, and the abundance was greater in CAMs compared with ATMs (Figure 3C).

To establish the functional significance of these results, we then studied MMP enzyme activity in CAM and ATM media. Using a fluorogenic substrate for MMP-1, we found significantly greater activity in CAM compared with corresponding ATM media (Figure 4A). Similarly, substrates selective for MMP-2, or for MMP-2/MMP-9 (Figure 4A), revealed greater activity in CAM than ATM media, and the same pattern was observed with two different MMP-3 substrates (Figure 4A).

**Figure 4.** Increased MMP-1, MMP-2, and MMP-3 activity in CAM media. (A) MMP-1, MMP-2, and MMP-3 enzyme activities (using the selective substrates indicated in brackets) are increased in CAM compared with ATM media; left, assays based on Trp fluorescence and right on FRET. (B) CAM-CM media stimulates AGS cell migration and is inhibited by neutralizing antibody to MMP-1 (2.5 μg mL−1) or inhibitors (see text) of MMP-2 (6 μM) and MMP-3 (3 μM). Horizontal bars, p < 0.05.

### DISCUSSION

The tumor microenvironment reflects the secretomes of both cancer and stromal cells including myofibroblasts, fibroblasts, pericytes, endothelial cells, inflammatory and immune cells. Crucially, interactions between different secretomes influence cancer cell migration, invasion and metastasis by multiple mechanisms including the activation or inhibition of proteases.
with consequences for the proteolytic cleavage of ECM proteins, growth factors, cytokines and chemokines. The secretomes of cancer cells have attracted increasing attention in recent years, but little is yet known of stromal cell secretomes. Differences in the secretomes of myofibroblasts recovered from gastric cancers and those recovered from adjacent tissue have been reported recently using iTRAQ. We have now used SILAC and COFRADIC to determine the extent to which proteolysis influenced myofibroblast secretomes in gastric cancer. Our study identified neo-N-termini derived from 94 proteins in CAM secretomes including evidence of cleavage of the prodomains of MMP-1, -2, and -3 leading to increased extracellular proteolytic activity. The data indicate that a distinguishing feature of cancer myofibroblasts is increased expression and increased activation of these MMPs in an autonomous manner in gastric cancer with the potential for promoting cancer progression.

The identification of secreted biomarkers by cancer cells has been a focus of interest for several generations and has been stimulated more recently by the development and refinement of proteomic methods and the prospect of rigorously defining the cancer secretome. Extracellular proteolysis presents additional challenges in defining the relevant secretomes; it is important not least because it underlies multiple mechanisms implicated in cancer progression including angiogenesis, tumor cell migration and invasion. A number of proteomic methods have recently been used to identify neo-N-termini generated in complex samples including COFRADIC and terminal amine isotopic labeling of substrates (TAILS), which has been applied to the identification of substrates of MMP-2 and MMP-9 in fibroblast secretomes. Since SILAC has previously been used successfully for secreteme studies in a range of cell types including stromal cells from other tissues, and COFRADIC coupled with SILAC labeling is considered to offer a rigorous approach to N-terminomics these methods were selected for the present studies.

The present study identified proteins in media released by the endoplasmic reticulum–Golgi secretory pathway; however, as commonly found in secretome studies, there were also proteins likely to be released by other mechanisms including cytoplasmic proteins liberated through cell death, shedding of membrane proteins and release of exosomes. Previous studies have identified similar rates of apoptosis in CAMs and ATMs suggesting that differential cell death is unlikely to account for differences in the CAM and ATM secretomes. For proteins released through classical secretory mechanisms, we were able to identify many previously reported in the secretomes of fibroblasts and of the stem cells that may give rise to them, including extracellular matrix proteins, IGFBPs, MMPs and TIMPs. The differences between CAM and ATM secretomes may reflect alterations in gene expression, post-translational processing, relative rates of secretion and in proteolysis following release. We have now defined the latter though identification of neo-N-termini in CAMs compared with their corresponding ATM. For example, we found multiple neo-N-termini in collagens alpha-2 (I) and alpha-1(I), and IGFBP5, as well as at limited sites in 91 other proteins. In a small subset of proteins we identified neo-N-termini that were unique to CAMs, and these included six neo-N-termini in the prodomains of MMP-1 and MMP-3, suggesting that activation of these MMPs might be functionally important in CAMs.

It is well established that MMP activity is increased in tumors and promotes cancer cell migration and invasion; the present in vivo imaging data indicate that CAMs contribute to this increased activity in an animal model. It is likely that MMPs have multiple roles in different tumor functions accounting for the fact that MMP inhibitors have not yet led to successful anticancer therapies. The present demonstration of increased expression and activation of MMP-1, MMP-2 and MMP-3 in CAM secretomes nevertheless suggests a novel dimension to the role of these enzymes. In vivo there may be activation of myofibroblast MMPs by epithelial or cancer-derived proteases, e.g., MMP-7. Importantly, however, the present data indicate that increased MMP activity in CAM media occurs independently of a cancer or epithelial cell stimulus. This is nevertheless relevant to cancer cell function not least because MMP-1, MMP-2, and MMP-3 stimulate cancer cell migration and make a substantial contribution to the chemotactic properties of CAM conditioned media. The prodomain cleavages of MMP-1, -2, and -3 identified here are all on the N-terminal side of the conserved cysteine switch sequence, and we think it is possible that these facilitate activation by exposing the site for autolysis in the same way that trypsin activates MMPs. However, the precise protease(s) responsible for CAM-autonomous prodomain cleavages is presently unclear and should now be investigated. In this context it is worth noting that there was decreased abundance of protease inhibitors, including TIMPs-1, -2, and -3 in one patient (Supporting Information Table S4), which may contribute to increased protease activity.

The present study using SILAC–COFRADIC has provided the most detailed analysis of gastric cancer myofibroblast secretomes to date. It extends previous studies, but in addition, it defines differences in the extracellular degradomes of CAMs and ATMs. At least some of the differences between CAFs or CAMs and their normal tissue counterparts are thought to reflect interactions that occur in the presence of cancer cells. The present data indicate that selective MMP-activation occurs in the CAM secretome even when these cells are cultured in the absence of cancer cells. This property reflects a cell-autonomous mechanism by which CAMs might contribute to a cancer-promoting microenvironment. The possibility of targeting anticancer therapies to stromal cells has emerged recently, and the present data indicate how these could be refined to include the stromal degradome.

**ASSOCIATED CONTENT**

**Supporting Information**

Supporting methods, tables, and figures. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**

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