Integrin Fibronectin Receptors in Matrix Metalloproteinase-1–Dependent Invasion by Breast Cancer and Mammary Epithelial Cells

Yifeng Jia, Zhao-Zhu Zeng, Sonja M. Markwart, Korrene F. Rockwood, Kathleen M. Woods Ignatowski, Stephen P. Ethier, and Donna L. Livant

Department of Radiation Oncology and Comprehensive Cancer Center, University of Michigan, Ann Arbor, Michigan

ABSTRACT

Integrins contribute to progression in many cancers, including breast cancer. For example, the interaction of αβ1 with plasma fibronectin causes the constitutive invasiveness of human prostate cancer cells. Inhibition of this process reduces tumorigenesis and prevents metastasis and recurrence. In this study, naturally serum-free basement membranes were used as invasion substrates. Immunoassays were used to compare the roles of αβ1 and αββ1 fibronectin receptors in regulating matrix metalloproteinase (MMP)-1–dependent invasion by human breast cancer and mammary epithelial cells. We found that a peptide consisting of fibronectin PHSRN sequence, Ac-PHSRN-NH2, induces αβ−β1-mediated invasion of basement membranes in vitro by human breast cancer and mammary epithelial cells. PHSRN-induced invasion requires interstitial collagenase MMP-1 activity and is suppressed by an equimolar concentration of a peptide consisting of the LDV sequence of the fibronectin connecting segment, Ac-LHGLPEILDV-PST-NH2, in mammary epithelial cells, but not in breast cancer cells. This sequence interacts with αβ1, an integrin that is often down-regulated in breast cancer cells. Immunoblotting shows that the PHSRN peptide stimulates MMP-1 production by serum-free human breast cancer and mammary epithelial cells and that the LDL peptide represses PHSRN-stimulated MMP-1 production only in mammary epithelial cells. Furthermore, PHSRN stimulates MMP-1 activity in breast cancer cells and mammary epithelial cells with a time course that closely parallels invasion induction. Thus, down-regulation of surface αβ1 during oncogenic transformation may be crucial for establishment of the αβ1–induced, MMP-1–dependent invasive phenotype of breast cancer cells.

INTRODUCTION

Integrins are a family of αβ heterodimeric receptors that mediate cell–matrix and cell–cell interactions and have important functions in cell migration, survival, and differentiation (reviewed in refs. 1 and 2). In addition to functioning in cellular adhesion to the extracellular matrix, integrins are known to mediate many diverse processes in cell migration, survival, and differentiation (reviewed in refs. 1 and 2). For example, we have shown that the interaction of plasma fibronectin receptor, αβ1, with fibronectin has been shown to play a key role in wound healing by mediating invasion of the provisional matrix by fibroblasts, endothelial cells, and keratinocytes to accomplish angiogenesis and re-epithelialization (3, 4). Also, integrins have been shown to function in the progression of a variety of cancers (reviewed in ref. 5).

One very important way that integrins can contribute to cancer progression is to mediate cancer cell invasion, thus causing metastasis. For example, we have shown that the interaction of plasma fibronectin (pFn) with αβ1 integrin induces human prostate cancer cell invasion (6–8) by using the naturally serum-free (SF), selectively permeable (9) basement membranes of sea urchin embryos [sea urchin embryo basement membranes with extracellular matrix (SU-ECM)] as in vitro invasion substrates (10). Even in the presence of serum, SU-ECM have been shown to be free of background invasion by unstimulated normal cells (3, 6, 10), which can be observed when artificial or reconstituted basement membranes are used (11). Also, these basement membranes are similar to those of mammals, both structurally and functionally (12–14). Because they are naturally SF, we used SU-ECM to identify PHSRN as the specific sequence of pFn that interacts with αβ1, to stimulate invasion by human prostate cancer cells and to show that inhibition of pFn-induced invasion reduces tumorigenesis and prevents metastasis and recurrence in both rat and human preclinical models of prostate cancer (6, 7).

Because fibronectin is found throughout the body, the proper regulation of αβ1–mediated collagenase expression, and hence invasion, is very important. This is accomplished by another integrin fibronectin receptor, αβ4. When fibronectin is intact, αβ4 integrin interacts with the LDV sequence of the fibronectin connecting segment, LHGLP-ILDVPST, to repress αβ1–mediated interstitial collagenase expression in adherent fibroblasts (15). Fragmentation of fibronectin by plasmin has been shown to de-repress αβ1–mediated invasion during wound healing (16, 17); thus, an important attribute of αβ1-induced invasion in normal cells is its regulation by αβ1 integrin.

Although still expressing abundant surface αβ1, metastatic prostate and breast cancer cells have low levels of surface αβ1, relative to prostate and mammary epithelial cells (18, 19). Loss of surface αβ1, which can result from oncogene overexpression in transformed mammary epithelial cells (19), causes constitutive invasiveness in the presence of the abundant pFn of blood, lymph, and interstitial fluid (20, 21). Because of its importance in breast cancer cell invasion and metastasis, we undertook this study to define the receptor–ligand interaction responsible for the invasive phenotype of metastatic breast cancer cells and to assess the role of interstitial collagenase matrix metalloproteinase (MMP)-1 in basement membrane invasion in vitro.

MATERIALS AND METHODS

Cell Culture. SUM-52 PE and MCF-10A cells were cultured in SF media, as described previously (22); whereas 5% serum was present in cultured SUM-149 PT and normal human mammary epithelial (HME) cells (23). When necessary, SUM-149 PT cells were serum-starved overnight before the addition of peptides.

Peptide Synthesis. NH2-terminal acetylated, COOH-terminal amidated PHSRN and LDV peptides (Ac-PHSRN-NH2 and Ac-LHGLPEILDV-PST-NH2), their randomized sequence controls (Ac-HPSPR-NH2 and Ac-PVGLSE-HPTLID-NH2), RGD peptide (Ac-GRGDS-PN-NH2), and VKNEED peptide (Ac-VKNEED-NH2) were synthesized using Fmoc/butyl-9-fluorenylmethoxycarbonyl/butyl protection strategies (24) at 25- and 100-μm scales on a Ranin Symphony peptide synthesizer. COOH-terminally amidated peptides were synthesized on Rink resin. Anhydrous trifluoroacetic acid was used to remove side chain protecting groups and to cleave peptides from the resin support. Peptides were precipitated with diethyl ether, purified by preparative

Received 1/9/04; revised 8/24/04; accepted 9/29/04.

Grant support: A research grant from Attensoon, L.L.C. (D. Livant) and National Cancer Institute R01 grant CA 70354 (S. Ethier).

The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked advertisement in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

Note: Y. Jia is currently in the Laboratory of Pathology, Center for Cancer Research, National Cancer Institute, National Institutes of Health, Bethesda, Maryland. S. Ethier is currently in the Barbara Ann Karmanos Cancer Institute, Wayne State University, Detroit, Michigan.

Requests for reprints: Donna L. Livant, Department of Radiation Oncology, Room 3007, 1331 East Ann Street Building, Ann Arbor, MI 48109-0582. Phone: 734-764-0313; Fax: 734-763-1581. E-mail: dlivant@umich.edu.

©2004 American Association for Cancer Research.
high-performance liquid chromatography, and lyophilized. Peptide purities were assessed by reverse-phase high-performance liquid chromatography and found to be 95% for Ac-PHSRN-NH2, 97% for Ac-HSPRN-NH2, 93% for Ac-LHGPEDLVPT-NH2, 92% for Ac-PGVLSEHPTLID-NH2, 95% for Ac-GRGDSP-NH2, and 91% for Ac-VKNEED-NH2 (data not shown). Peptide structures were confirmed by mass spectrometry and amino acid analysis (data not shown) using standard methods (24). Residual trifluoroacetic acid was removed by gel permeation chromatography on Sephadex G-10 in 1 N acetic acid. Peptides were lyophilized and stored in the presence of a desiccant at −20°C until solubilization in phosphate-buffered saline at 1 mg/mL at the time of use.

**Antibodies Used in Invasion Assays.** P1D6 anti-α5β1, or P1B5 anti-αβ1, integrin function-blocking monoclonal antibodies (mAbs) (25, 26), from Oncogene Research Products (Boston, MA), were prebound to suspended cells in SF or fetal calf serum (FCS)-containing media on ice for 30 minutes at 10 to 300 μg/mL before 10-fold dilution in the appropriate medium and place on SU-ECM. P1H4 and P4C2 function-blocking anti-αβ1 integrin mAbs (27, 28), from Chemicon International (Temecula, CA), were prebound to cells, and the cells were used in invasion assays as described above. Function-blocking anti–MMP-1 (COM-4A2), anti–MMP-2 (CA-4001), and anti–MMP-9 (GE-213) mAbs (29–31) from Chemicon International were prebound to cells, and the cells used in invasion assays as described above. Isotype control antibodies were as follows: IgM (BD Biosciences PharMingen, San Diego, CA); and IgG1 and IgG3 (Sigma, St. Louis, MO). All isotype control antibodies were prebound to cells, and the cells were used in invasion assays as described above.

**Invasion Assays.** Preparation of SU-ECM and in vitro SU-ECM invasion assays were performed with or without added FCS, as described previously (3, 6, 10). Plasma fibronectin-depleted (pFn-) FCS was made using gelatin affinity chromatography, as described previously (6, 19, 21). Single-cell suspensions were made with 0.25% trypsin/EDTA (Gibco Life Technologies, Inc., Grand Island, NY). Cell suspensions were rinsed by pelleting and resuspension in the appropriate medium before placement on SU-ECM invasion substrates. Invasion percentages and cellular viabilities were scored as described previously (3, 6, 10, 19). Peptides were added to the invasion assays by prebinding to suspended, rinsed cells for 5 minutes at room temperature. For assays demonstrating anti–MMP-1 inhibition of PHSRN-induced invasion, the concentration of Ac-PHSRN-NH2 in the assay medium was 1 μg/mL. Antibodies were prebound to suspended cells, as described above. Each invasion percentage is the ratio of the total number of single cells located in the interior, blastocoeic cavities of the SU-ECM substrates to the total number of single cells adhering to both their exterior and interior surfaces, and is the result of three to four independent determinations involving the scoring of the positions of all individual cells adhering to the SU-ECM invasion substrates.

**Fluorescence-Activated Cell Sorting.** Fluorescence-activated cell sorting of SUM-52 PE and MCF-10A cells was performed as described previously (19), using anti-integrin α antibody (catalog no. 12077-012; Gibco) or anti-integrin β antibody (catalog no. 23012; Oncogene Research Products) followed by fluorescein-conjugated donkey antiovine IgG (1:100; Chemicon International). Negative controls contained cells bound to fluorescein-conjugated donkey antiovine IgG without primary antibody.

**SDS-PAGE and Immunoblotting.** Each sample contained 2 × 106 adherent cells treated with the appropriate peptides for 16 hours at 37°C. The treatment groups were as follows: SF medium; SF medium + Ac-PHSRN-NH2; SF medium + Ac-LHGPEDLVPT-NH2; SF medium + Ac-HSPRN-NH2 + Ac-LHGPEDLVPT-NH2; and SF medium + Ac-PHSRN-NH2 + Ac-PGVLSEHPTLID-NH2. All cells were rinsed, and 5 mL of fresh SF medium were added to each sample before peptide addition. Samples were treated as follows: 1 μg of Ac-PHSRN-NH2 per 20,000 cells (100 μg for 2 × 105 cells in 5 mL of medium); 2.5 μg of Ac-LHGPEDLVPT-NH2 per 20,000 cells; 1 μg of Ac-PHSRN-NH2, and 2.5 μg of Ac-PGVLSEHPTLID-NH2 per 20,000 cells.

SUM-52 PE, SUM-149 PT, and MCF-10A cells were rinsed with PBS and lysed in ice-cold buffer (50 mmol/L Tris-HCl (pH 7.5), 150 mmol/L NaCl, 2 mmol/L EGTA, and 1% Triton X-100) containing protease inhibitors (1× complete protease inhibitor mixture; Roche, Indianapolis, IN). The lysate was collected and centrifuged at 12,000 × g for 10 minutes at 4°C, and the resulting pellet was resuspended in SDS sample buffer [2% SDS, 62.5 mmol/L Tris-HCl (pH 6.8), and 10% glycerol]. The amounts of protein were measured for each sample using the Bio-Rad protein assay kit (catalog no. 500-0006; Bio-Rad, Richmond, CA) with albumin standards. Before electrophoresis, samples were brought to 5% (v/v) 2-mercaptoethanol and boiled for 5 minutes.

To verify a rapid increase in latent and activated MMP-1 secreted into the SF medium after Ac-PHSRN-NH2 treatment, adherent cultures of 2 × 106 SUM-52 PE or SUM-149 PT breast cancer cells and 2 × 106 MCF-10A mammary epithelial cells in SF medium were treated with 1 μg of Ac-PHSRN-NH2 per 20,000 cells (100 μg in 5 mL of medium) for periods of time ranging from 1 hour to 6 hours. Media were collected and concentrated 50-fold using centrifugal filter devices (Amicon Ultra PL-10 device; 10,000 nominal molecular weight limit; Millipore, Bedford, MA), according to the manufacturer’s instructions. The quantitative consistency of the volume reduction was verified by using a micropipetting device for all media assayed.

For cell lysates, 30 μg of total protein per sample in SDS buffer were resolved on 10% polyacrylamide gels using a mini-PROTEAN II Electrophoresis Cell (Bio-Rad, Hercules, CA). Separated proteins were transferred onto polyvinylidene difluoride membranes (Millipore) using the submarine electrophoretic transfer unit in the same apparatus. Varying amounts (10, 5, 2.5, and 1.25 ng) of recombinant MMP-1 (catalog no. CC1031; Chemicon International) were combined with cytoskeletal actin and loaded on the same gel for generation of standard curves. Membranes were blocked for 1 hour in 0.1% (v/v) Tween 20 in PBS (PBS-T) containing 5% (w/v) nonfat dry milk and then incubated for 1 hour at room temperature with anti–MMP-1 mAb (clone COM-4A2; Chemicon International) at a 1:5,000 dilution in blocking solution.

For the analysis of secreted MMP-1 levels by immunoblotting, identical volumes of concentrated media from PHSRN-treated and untreated cells were run on 10% polyacrylamide gels, as described above. Recombinant MMP-1 was diluted 3,000-fold and run in varying amounts on each gel as positive controls. Separated proteins were transferred onto polyvinylidene difluoride membranes as described above, and membranes were incubated overnight at 4°C with rabbit anti–MMP-1 polyclonal antibody (catalog no. AB806, Chemicon International) at a dilution of 1:1,000 in Tris-buffered saline with 0.05% Tween 20.

After incubation with the primary antibody, all membranes were washed three times in PBS-T and incubated for 1 hour with goat anti–IgG antibody conjugated to horseradish peroxidase (Jackson Immunoresearch Laboratories, West Grove, PA) at a dilution of 1:5,000 in blocking solution. The membranes were then washed three times in PBS-T and processed for detection by enhanced chemiluminescence ECL reagent (Amer sham, Arlington Heights, IL). The amounts of MMP-1 and actin from the cell lysates were quantified using Quantity One software (Bio-Rad, Hercules, CA) by comparison with the recombinant MMP-1 and actin standard curves generated from the same blot. Then, the amount of MMP-1 was normalized to the amount of total cellular actin in each sample from cell lysates in immunoblots used to compare MMP-1 levels in the media of PHSRN-treated and untreated cells, equal volumes of concentrated media were loaded without normalization of protein content.

**Matrix Metalloproteinase-1 Activity Assay.** Adherent SUM-149 PT, SUM-52 PE, or MCF-10A cells were treated for various times with Ac-PHSRN-NH2 at a concentration of 1 μg per 20,000 cells in SF medium, and untreated controls were performed in parallel. Culture media were concentrated 10-fold by centrifugation through Centricon YM-10 filter units (Fisher Scientific Company, L.L.C., Pittsburgh, PA), according to the manufacturer’s instructions. The quantitative consistency of the volume reduction was verified by using a micropipetting device for all media assayed. MMP-1 activity was measured in concentrated media with the Biotrak MMP-1 activity assay system (Amersham Pharmacia Biotech Inc., Piscataway, NJ), according to the manufacturer’s instructions. The assays were read at 405 nm in a microtiter plate spectrophotometer (Dynatech Laboratories, Inc., Chantilly, VA). A standard curve was generated from a set of known aliquots of MMP-1 in the following concentration range: 0, 0.10, 0.20, 0.40, 0.80, and 1.60 ng/mL. Each time point was assayed at least three times in triplicate, and mean MMP-1 activities were determined with their first SDs. The results were analyzed using Student’s t test.
RESULTS

Invasion Induction by the PHSRN Peptide. The effects of the PHSRN peptide on invasion by HME and MCF-10A mammary epithelial cells and SUM-149 PT and SUM-52 PE breast cancer cells were evaluated on SU-ECM invasion substrates. As shown in Fig. 1A, the blocked PHSRN peptide stimulated invasion by HME with a log-linear dose-response relationship at concentrations from 10 ng/mL to 1 μg/mL (17 nmol/L to 1.7 μmol/L), whereas the scrambled, blocked PHSRN peptide, Ac-HSPRN-NH₂, was without detectable activity at 17 μmol/L. In addition to PHSRN (32), αβ₁ interacts with the RGD (8) and VKNEED sequences of the fibronectin cell binding domain (33). Thus, the acetylated, amidated derivatives of these peptides were also tested for invasion induction. As shown in Fig. 1A, neither GRGDSP nor VKNEED induced HME invasion at concentrations as high as 10 μg/mL. Very similar results were obtained for MCF-10A cells (Fig. 1B). In addition, the PHSRN peptide was equally effective at stimulating invasion by MCF-10A and HME cells, irrespective of the presence of FCS. Furthermore, no invasion occurred in the absence of the PHSRN peptide, either in SF medium or in the presence of FCS (data not shown).

Under SF conditions, the PHSRN peptide was equally effective at stimulating invasion by SUM-149 PT breast cancer cells, whereas the HSPRN, RGD, and VKNEED peptides lacked activity, as shown in Fig. 1C. Quantitatively similar results were obtained for SUM-52 PE breast cancer cells in SF medium, as shown in Fig. 1D. Thus, the PHSRN peptide was as effective at stimulating invasion by MCF-10A and HME cells, irrespective of the presence of FCS. Furthermore, no invasion occurred in the absence of the PHSRN peptide, either in SF medium or in the presence of FCS (data not shown).

Induction of Plasma Fibronectin-Dependent Invasion in Mammary Epithelial Cells by Anti-αβ₁ Antibody. Because loss of surface αβ₁ causes a pFn-dependent, constitutively invasive phenotype in SUM-149 PT cells (19), we tested whether blocking αβ₁ could render mammary epithelial cells invasive in the presence of serum. Levels of surface αβ₁ and αβ₁ are shown for mammary epithelial and breast cancer cells in Table 1. Like HME, MCF-10A cells express relatively high levels of surface αβ₁ and αβ₁. In contrast, whereas SUM-52 PE and SUM-149 PT cells express abundant surface αβ₁, fluorescence-activated cell-sorting analysis shows that they express very low levels of αβ₁.

To assess the role of αβ₁ in repressing serum-induced invasion, HME and MCF-10A cells were treated with blocking anti-αβ₁ mAb before placement on SU-ECM invasion substrates. Fig. 2A shows the relative percentages of invasive HME and MCF-10A cells when the cells were prebound with increasing amounts of P1H4 or P4C2 blocking anti-αβ₁ mAb (27, 28). Both P4C2 and P1H4 mAbs stimulated SU-ECM invasion by HME and MCF-10A cells in the presence of FCS, whereas their corresponding isotype control antibodies did not.

Because the P4C2 epitope specifically includes the region of αβ₁ that binds the LDV sequence (28, 34) to repress serum-induced invasion, we tested the extent to which blocking αβ₁ mAbs stimulated mammary epithelial cell invasion in FCS-containing medium. X axis, log antibody concentration (μg/mL); Y axis, mean percent of invasive cells relative to positive controls containing 1 μg/mL Ac-PHSRN-NH₂. Invasion by either HME or MCF-10A cells did not occur in the absence of anti-αβ₁ mAb. ○, HME cells bound purified P1H4 mAb; ●, MCF-10A cells bound to purified P1H4 mAb; ■, HME cells bound to P4C2 ascites fluid; □, HME cells bound to P1H4 isotype control; △, MCF-10A cells bound to P1H4 isotype control; □, HME cells bound to P4C2 isotype control. Symbols ■ and □ reflect the mAb concentration range of the P4C2 ascites fluid. Vertical bars, first SDs. B, dependence of P4C2 anti-αβ₁-induced invasion on pFn. X axis, mean percentages of invasive cells relative to positive controls performed in parallel: percentage of invasive cells in the presence of FCS after binding to 100 μg/mL P3C2 mAb. Y axis, media components. ■ and □ reflect the mAb concentration range of the P4C2 ascites fluid. Vertical bars, first SDs. C, dependence of P4C2 anti-αβ₁-induced invasion on FCS. X axis, log antibody concentration (μg/mL). Y axis, mean percentages of invasive cells, relative to invasion percentages observed in positive controls (invasion assays containing 1 μg/mL PHSRN and performed in parallel). ○, Ac-PHSRN-NH₂-treated cells; ●, Ac-HSPRN-NH₂-treated cells; ■, Ac-GRGDSP-NH₂-treated cells; □, Ac-VKNEED-NH₂-treated cells. Vertical bars, first SDs.

### Table 1 Cell surface integrin expression

<table>
<thead>
<tr>
<th>Cell line</th>
<th>α₅⁺</th>
<th>α₇⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>HME 1</td>
<td>54</td>
<td>62</td>
</tr>
<tr>
<td>MCF-10A</td>
<td>58</td>
<td>30</td>
</tr>
<tr>
<td>SUM-52 PE</td>
<td>5</td>
<td>28</td>
</tr>
<tr>
<td>SUM-149 PT</td>
<td>0</td>
<td>10</td>
</tr>
</tbody>
</table>

NOTE. As determined by fluorescence-activated cell-sorting analysis. * Percentage increase in the amount staining over negative control. † Data from ref. 19.
MMP-1 expression (15), the P4C2 mAb was tested for its ability to stimulate SU-ECM invasion by HME and MCF-10A cells while the presence of FCS and/or pFn was varied systematically in the media of the invasion assays. Antibody prebinding occurred in either SF medium, FCS-containing medium, or pFn-depleted FCS-containing medium, as indicated. As shown for HME and MCF-10A in Fig. 2B, the P4C2 anti-α5β1 mAb was unable to stimulate invasion in SF medium, but it stimulated invasion by HME and MCF-10A cells in FCS-containing medium. Also, P4C2-stimulated invasion was specifically dependent on pFn because no invasion occurred if P4C2-bound HME or MCF-10A cells were incubated on SU-ECM substrates in pFn− FCS. The addition of 4 μg/mL pFn, the concentration found in 10% FCS (20, 21), to medium containing pFn− FCS (pFn− FCS + pFn) restored the ability of the P4C2 mAb to stimulate HME and MCF-10A invasion. Also, the addition of 4 μg/mL pFn to SF medium (SF + pFn) was sufficient to permit P4C2-induced invasion. Thus, pFn appeared to be both necessary and sufficient for P4C2 anti-α5β1−induced invasion by both HME and MCF-10A cells.

Repression of PHSRN-Induced Invasion by the LDV Sequence. An acetylated, amidated peptide containing the LDV sequence (Ac-LHGPDEILDVPST-NH₂) was tested for its ability to repress PHSRN-induced invasion by normal and neoplastic breast epithelial cells. Because a single copy of each sequence is found in the fibronectin monomer (32, 34), the PHSRN and LDV peptides were tested at equimolar concentrations. Fig. 3A shows that whereas PHSRN stimulated HME invasion of the SU-ECM, the presence of an equimolar concentration of the LDV peptide prevented PHSRN-induced invasion. Furthermore, inhibition was sequence specific because the scrambled LDV peptide (Ac-PGVLSEHPTLID-NH₂) had no inhibitory effect on PHSRN-induced invasion. Analogous results were obtained for MCF-10A cells (Fig. 3B). Thus, the ability of the LDV peptide to inhibit PHSRN-induced HME and MCF-10A invasion is consistent with the expression of abundant surface α5β1 and α6β1 fibronectin receptors by these cells (ref. 19 and this study). In contrast, as shown in Fig. 3C and D, the LDV peptide had no inhibitory effect on PHSRN-induced invasion by either SUM-149 PT or SUM-52 PE cells. The failure of the LDV sequence to inhibit PHSRN-induced SUM-149 PT and SUM-52 PE invasion is likewise consistent with the expression of surface α6β1, but not α5β1, by these breast cancer cell lines.

Role of Matrix Metalloproteinase-1 in α5β1-Mediated Invasion. The role of MMP-1 in PHSRN-induced invasion was tested on SF SU-ECM invasion substrates in the presence of 100 ng/mL PHSRN peptide and increasing concentrations of blocking COMY-4A2 anti-MMP-1 mAb (29). As shown in Fig. 4A, the anti–MMP-1 mAb was equally effective at blocking PHSRN-induced invasion by all four cell types. Complete inhibition of invasion was achieved by a concentration of 100 to 300 μg/mL, whereas the isotype control antibody (300 μg/mL) failed to significantly affect invasion. In contrast to the role of MMP-1, results of invasion assays using inhibitory anti–MMP-2 (CA-4001) and anti–MMP-9 (GE-213) mAbs (30, 31) suggested that neither MMP-2 nor MMP-9 activities function in PHSRN-induced invasion of SU-ECM, although zymography of SF media from PHSRN-treated and untreated MCF-10A, SUM-52 PE, and SUM-149 PT cells shows that all of these cell lines secrete activated forms of both MMP-2 and MMP-9 and that the levels of both gelatinases are unchanged by exposure to the PHSRN peptide (data not shown). The anti–MMP-1 mAb was also equally effective at inhibiting serum-induced invasion by SUM-52 PE and SUM-149 PT cells (Fig. 4B). Although SUM-52 PE and SUM-149 PT cells, as well as MCF-10A cells, secrete abundant MMP-2 and MMP-9 in the presence of serum (data not shown), insignificant inhibition of serum-induced invasion was observed with high concentrations of the anti–MMP-2 and anti–MMP-9 mAbs. Thus, these data demonstrate that MMP-1 also functions specifically in serum-induced invasion by SUM-149 PT and SUM-52 PE cells. Furthermore, the specific role of MMP-1 is serum-induced invasion was also confirmed for HME and MCF-10A mammary epithelial cells because increasing concentrations of anti–MMP-1 mAb reduced anti α5β1-induced invasion, whereas anti–MMP-2 and anti–MMP-9 mAbs had no significant effect (Fig. 4C).

Stimulation of Matrix Metalloproteinase-1 Accumulation by the PHSRN Peptide. Adherent, SF breast cancer and mammary epithelial cells were treated with PHSRN and/or LDV peptides, and the cells were lysed, immunoblotted, and probed with anti–MMP-1 rabbit antiserum to compare the levels of cell-associated MMP-1. Immunoblots were run four times for each cell type, and relative band densities were determined for each treatment. Nearly all MMP-1 appeared to migrate in a single band of Mr 52,000, suggesting that cell-associated MMP-1 is predominantly the latent form. Mean MMP-1 band densities with first SDs are shown in A–C of Fig. 5, as follows: Fig. 5A, MCF-10A; Fig. 5B, SUM-52 PE; and Fig. 5C, SUM-149 PT. As shown in Fig. 5A, PHSRN stimulated MMP-1 accumulation in MCF-10A cells, whereas equimolar LDV peptide did not. However, when mixed with the PHSRN peptide, LDV blocked the accumulation of MMP-1, whereas the scrambled LDV peptide failed to reduce PHSRN-induced MMP-1 accumulation. These results are consistent with the reported role of the α5β1 receptor in fibroblasts (15). Moreover, consistent with the lack of surface α5β1 expression in SUM-52 PE and SUM-149 PT cells, the LDV peptide had no inhibitory effect on PHSRN-induced MMP-1 accumulation in these breast cancer cells (Fig. 5B and C). Fig. 5D shows an example of a typical blot used for this analysis.

Although MMP-1 has been shown to associate with surface α5β1 integrin collagen receptor to achieve activation and facilitate migration on collagen (35) and would thus be detectable in cell lysates, MMP-1 should also be secreted into the media of adherent cell
cultures treated with the PHSRN peptide. To test whether PHSRN treatment could cause a rapid increase in both the latent and activated forms of MMP-1 in the medium, adherent SF cultures of MCF-10A, SUM-52 PE, and SUM-149 PT cells were treated for 1 hour to 6 hours with the PHSRN peptide or were left untreated. All treated and untreated cultures were run in triplicate. Media from PHSRN-treated and untreated cultures were concentrated, and equal volumes of the concentrated media from each sample were analyzed by immunoblotting for the presence of MMP-1. As shown in Fig. 6A, 2 hours of PHSRN treatment induced significant increases in the mean levels of both latent and activated MMP-1 in the media from adherent MCF-10A, SUM-52 PE, and SUM-149 PT cells, relative to the amounts...
secreted by untreated cells. Fig. 6B compares examples of the M, 52,000 and M, 43,000 MMP-1 bands from PHSRN-treated and untreated MCF-10A, SUM-149 PT, and SUM-52 PE cells. The PHSRN-induced up-regulation of two anti–MMP-1 antibody-reactive proteins, a M, 52,000 latent form and a form exhibiting a M, 9,000–10,000 reduction, is illustrated. Moreover, the observed molecular weight is consistent with those reported for latent MMP-1 (M, 52,000) and activated MMP-1 (M, 43,000; ref. 36). Also, as shown in Fig. 6, both bands comigrated with the purified MMP-1 standards run on each gel. For each cell type, maximal levels of latent and activated MMP-1 were observed after 2 hours of treatment, and then they began to decline (data not shown).

Stimulation of Matrix Metalloproteinase-1 Activity by the PHSRN Peptide. MMP-1 activity levels were monitored in media of adherent cultures of MCF-10A, SUM-52 PE, and SUM-149 PT cells for up to 6 hours after the initiation of PHSRN treatment. As shown in Fig. 7A, activity reached a maximum in about 2 hours and then started to decline toward the control level after 4 hours. Differences in MMP-1 activity between PHSRN-treated cells and untreated cells incubated in parallel were highly significant (P < 0.0001). Also, the time of peak MMP-1 activity corresponded with the time of maximal MMP-1 levels in the medium, as assayed by immunoblotting (Fig. 6; data not shown). As shown in Fig. 7B, temporal changes in MMP-1 activity were also well correlated with the PHSRN-induced invasion time courses of these cell lines on SU-ECM invasion substrates. Furthermore, suspension and placement on SU-ECM for identical periods of time, without exposure to PHSRN peptide, resulted in no detectable invasion by any of the cell lines.

DISCUSSION

We report that a peptide consisting of the PHSRN sequence of fibronectin induces rapid, αβ3-mediated invasion by normal and immortalized mammary epithelial cells, as well as by metastatic breast cancer cells. Although the intact fibronectin of serum does not induce mammary epithelial cell invasion, due to αβ3 expression, invasion is induced if αβ3 is inhibited with a blocking antibody. Furthermore, αβ3 repression of αβ3-mediated invasion occurs if PHSRN-treated mammary epithelial cells are also exposed to a peptide consisting of the LDV sequence (34) of the fibronectin connecting segment. Moreover, in SF breast cancer cells lacking αβ3 on their surfaces, the LDV peptide fails to repress PHSRN-induced invasion. These results are consistent with the down-regulation of αβ3 on the surfaces of invasive breast cancer cells as well as in transformed mammary epithelial cells overexpressing the ERBB-2 oncogene (19) and in prostate cancer cells (18). They suggest that αβ3 functions to regulate αβ3-mediated invasion and that the loss of surface αβ3 may be an important event in the development of the invasive phenotype in cancer.

We also found, through the use of blocking anti–MMP-1, anti–MMP-2, and anti–MMP-9 mAbs, that PHSRN- and serum-induced invasion of basement membranes by mammary epithelial cells and breast cancer cells appears to be a MMP-1–dependent process. Consistent with the apparent dependence of invasion on MMP-1, it was also observed that PHSRN treatment induces a rapid increase in both cell-associated and secreted MMP-1 protein, as well as in MMP-1 activity in the medium. Furthermore, PHSRN-induced MMP-1 accumulation in mammary epithelial cells, but not in breast cancer cells, is prevented by the LDV peptide ligand of αβ3 integrin. Also, the rapid increase in PHSRN-induced MMP-1 activity closely parallels that of basement membrane invasion. A similarly rapid induction of MMP-1 expression has also been observed in osteoblasts treated with platelet-derived growth factor (37). Thus, MMP-1 may play an important role in αβ3-mediated invasion in breast cancer cells, and its up-regulation is likely a consequence of reduced levels of surface αβ3 fibronectin receptor relative to mammary epithelial cells. Because reduced levels of surface αβ3 integrin also give rise to αβ3-mediated, serum-dependent invasiveness in prostate cancer (6), this could be a general mechanism contributing to metastasis.

MMPs carry out most of the connective tissue destruction associ-
ated with cancer invasion and metastasis (reviewed in ref. 38). Zymography, immunoblotting, and immunohistochemistry have demonstrated increased levels of MMPs and MMP activities in human breast cancer, relative to normal breast tissues (39). However, the role of interstitial collagenases in tumor invasion and metastasis has only recently been appreciated. It has been suggested that collagenase expression is a marker for tumor progression in breast cancer, as well as in many other types of cancers (40, 41). Zymography of human breast carcinomas and normal breast tissues has also shown significant MMP-1 levels in most invasive cancers, in contrast to normal breast tissues (42). Immunohistochemistry of sectioned, invasive breast tumors and their surrounding tissues has also shown a significant correlation of MMP-1 expression with tumor stage (39). Furthermore, in most invasive breast carcinomas, in situ hybridization has demonstrated high levels of MMP-1 transcripts in both breast cancer cells and stromal cells at their invasive fronts (43). In fact, MMP-1 is also up-regulated in human breast cancer cells cocultured with fibroblasts (44), suggesting that direct interactions between these two cell types can lead to increased collagenase expression in breast cancer. In contrast, although MMP-2 transcripts have been found at high levels in invasive breast carcinomas, MMP-2 mRNA is also up-regulated in preinvasive lesions, suggesting that MMP-1 is specifically up-regulated as breast cancers become invasive (43). Interestingly, it has also been shown that interstitial collagenase also cleaves entactin, thus contributing directly to the degradation of basement membrane and hence potentially contributing to the transiting of epithelial barriers by tumor cells (45), in addition to stromal proteolysis.

In other cancers, such as colon and esophageal cancers, immunohistochemical detection of MMP-1 expression is also associated with increased invasive potential and poor prognosis (46, 47). Moreover, increased MMP-1 expression in tumor cells is significantly correlated with the depth of tumor invasion, angiogenesis, lymphangiogenesis, and the presence of local and distant metastases (48). Also, a transcription-enhancing point mutation in the MMP-1 gene promoter, which correlates with MMP-1 overexpression in tumor cells, is associated with increased malignancy in breast and lung cancer (49, 50). Thus, whereas many MMPs contribute to tumor angiogenesis and metastasis, interstitial collagenase may be critical for the development of the invasive phenotype during cancer progression.

**REFERENCES**

22. Dumin JA, Dickeson SK, Stricker TP, et al. Pro-collagenase-1 (matrix metallopro-
Integrin Fibronectin Receptors in Matrix Metalloproteinase-1–Dependent Invasion by Breast Cancer and Mammary Epithelial Cells

Yifeng Jia, Zhao-Zhu Zeng, Sonja M. Markwart, et al.

Cancer Res 2004;64:8674-8681.