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TITLE: Proteolytic Processing of Laminin-332 by Hepsin and Matriptase and Its Role in Prostate Cancer Progression

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Laminin-332 is lost in prostate cancer progression. Laminin-332 is known to be cleaved by various cell surface proteases, including MMPs, and this cleavage facilitates migration of cancer cells. We have found that Laminin-332 is individually cleaved by two serine proteases, hepsin and matriptase, and that this cleavage enhances migration of human prostate cancer cells in vitro. Hepsin is over-expressed in more than 90% prostate cancer cases. Similarly, matriptase is over-expressed in human prostate cancer cases and expression of both proteases correlates with tumor progression. However, the mechanism(s) by which these two serine proteases play a role in prostate cancer is unknown. We found that Hepsin/matriptase overexpressing prostate cancer cells, LNCaP show significantly increased migration on Ln-332. This project aims to define the role played by the cleavage of Laminin-332 by hepsin and matriptase in prostate cancer.

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**Introduction:**

The American Cancer Society estimated that there will be approximately 217,7300 new prostate cancer cases in the USA in 2010 (Jemal, Siegel et al. 2010). Extracellular matrix molecules such as laminin-332 (Ln-332) play a very important role in cancer progression. Ln-332 expression is decreased during prostate cancer progression. Conversely, type II transmembrane serine protease hepsin is over-expressed in more than 90% prostate cancer cases. Similarly, another type II transmembrane serine protease matriptase is over-expressed in human prostate cancer and expression of both proteases correlates with tumor progression. However, the mechanism(s) by which these two serine proteases play a role in prostate cancer is unknown. This project aims to define the role played by the cleavage of Ln-332 by hepsin and matriptase in prostate cancer. We have found (in the preliminary data submitted in the original DOD grant application) that Ln-332 is individually cleaved by two serine proteases, hepsin and matriptase, and that this cleavage enhances migration of human prostate cancer cells *in vitro*. Based on this preliminary data our central hypothesis of the proposal is that type II transmembrane serine proteases, Hepsin and Matriptase cleave Laminin-332 and that the proteolytic cleavage of laminin-332 plays a role in the progression of prostate cancer. We will explore this with implementation of two specific aims:

Specific Aim 1: Determine the role of cleavage of Laminin-332 by Hepsin and Matriptase in invadopodia formation and characterize these properties in prostate cancer cell migration and invasion.

Specific Aim 2a: Determine the effect of cleavage of Ln-332 by Hepsin and Matriptase on the formation of prostate tumors *in vivo*.

Specific Aim 2b: Determine the role of Hepsin in the formation of prostate cancer bone lesions and elucidate the role of cleavage of Ln-332 by Hepsin on the formation of bone lesions in vivo.
Task 1. Determine the role of cleavage of Laminin-332 by Hepsin and Matriptase in invadopodia formation and characterize these properties in prostate cancer cell migration and invasion. (Months 1-12)

We proposed that we will study the role of cleavage of Ln-332 by hepsin and matriptase in invadopodia formation; however our multiple attempts to see invadopodia formation in PC3 cells failed as discussed in detail in the previous annual report. Due to failure of this experiment, we designed a different strategy as detailed below to determine the role of cleavage of Ln-332 by hepsin and matriptase and to characterize these properties in prostate cancer cell migration.

As discussed in the previous annual report, we published a paper in the Journal of Biological Chemistry, Tripathi et al. JBC 2008 Nov 7;283 (45):30576-84 (Tripathi, Nandana et al. 2008). The published article was attached in the previous annual report.

There are some cell adhesion data that was not published in that paper, we would like to discuss them here,

Cell adhesion of DU145 cells exhibit no significant difference on hepsin-cleaved Ln-332

To determine if cellular adhesion of DU145 cells was different on hepsin cleaved Ln-332 versus untreated Ln-332 substrate, we performed cell adhesion assay as described in the Materials and Methods Chapter. We coated Ln-332 alone (L), Ln-332 + hepsin (LH), Ln-332 + hepsin + KD1 and PBS onto 96-well plates. DU145 cells were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained (crystal violet), and solubilized; and absorbance was read at 560 nm. We found that DU145 cells had no significant difference in cell adhesion on hepsin cleaved Ln-332 as compared to untreated Ln-332 or on mixture
with Ln-332, hepsin and KD1 (Figure 1). Results represent mean ± S.D. (N= 2, in duplicates; P < 0.05).

**Cell adhesion of hepsin-overexpressing prostate cancer cells exhibit no significant difference on Ln-332**

To determine if cellular adhesion of hepsin-overexpressing LNCaP-34 prostate cancer cells on Ln-332 was different as compared to low hepsin-expressing LNCaP-17 cells on Ln-332, we performed cell adhesion assay as described in Materials and Methods Chapter. We coated Ln-332 or PBS onto 96-well plates. Both Cell types were allowed to adhere for 1 h at 37ºC in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained (crystal violet), and solubilized; and absorbance was read at 560 nm. We found that hepsin-overexpressing LNCaP-34 cells had no significant difference in cell adhesion on Ln-332 as compared to low hepsin-expressing LNCaP-17 cells (Figure 2). Results represent mean ± S.D. (N= 3, in triplicates; P < 0.05).

As part of the proposed aims we also studied Ln-322 processing by another TTSP, matriptase. We have published these findings in The Prostate Journal (Manisha Tripathi, Alka Potdar, Hironobu Yamashita, Brandy Weidow Peter T. Cummings, Daniel Kirchhofer, and Vito Quaranta. Laminin-332 cleavage by Matriptase alters motility parameters of prostate cancer cells. Prostate. 2011 Feb 1;71(2):184-96 and the published paper is attached at the end of this report. In this study, we provide the first direct evidence that matriptase proteolytically processes Ln-332, a key extracellular matrix macromolecule found in the basement membrane of many epithelia, including prostate. Western blot and mass spectrometry indicated that proteolysis occurs in the β3 chain of Ln-332, near a domain involved with binding to collagen. Substrate specificity was confirmed by blocking cleavage with a matriptase inhibitor, Kunitz domain-1 (KD1). Transwell migration assays showed that DU145 prostate cancer cell motility was significantly enhanced when plated on matriptase-cleaved Ln-332.
Similarly, Transwell migration of matriptase-overexpressing LNCaP prostate cancer cells was increased on Ln-332 and, as determined by live single-cell microscopy, two of their motility parameters, speed and directional persistence, were also higher. Ln-332 has a demonstrated role in maintaining epithelial integrity. Its deregulated expression has been reported in several cancers and, especially relevant to this study, it is extinguished in advanced prostate cancer. Based on these findings, we proposed in the paper that proteolytic processing of Ln-332 could be a possible mechanistic role for matriptase in prostate cancer progression via altered migration parameters and subsequent basement membrane transgression (published article attached).

Again the cell adhesion data that was not published in this manuscript is discussed here:

Cell adhesion of DU145 cells exhibit no significant difference on matriptase-cleaved Ln-332

To determine if cellular adhesion of DU145 cells was different on matriptase cleaved Ln-332 versus untreated Ln-332 substrate we performed cell adhesion assay as described in Materials and Methods. We coated Ln-332 alone (L), Ln-332 + matriptase (LM), Ln-332 + matriptase + KD1 and PBS onto 96-well tissue culture treated plates. DU145 cells were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were removed by washing and adherent cells were fixed, stained with crystal violet, solubilized; and absorbance was read at 560 nm. We found that DU145 cells had no significant difference in cell adhesion on matriptase cleaved Ln-332 as compared to untreated Ln-332 or on mixture with Ln-332, matriptase and KD1 inhibitor. Results represent mean ± S.D. (N= 2, in duplicates; P < 0.05) (Figure 3).

Task 2A. Determine the effect of cleavage of Ln-332 by Hepsin and/or Matriptase on the formation of prostate tumors in vivo. (Months 12-24)
We proposed that we will perform orthotopic grafts into the prostates of nude mice to study the progression of tumors formed by genetically modified PC3 cells but we got a better approach to study this aim. To determine the effect of cleavage of Ln-332 by hepsin on prostate cancer in vivo we were fortunate to be able to study it in a hepsin overexpressing mouse model.

To this end we studied degradation of Ln-332 by hepsin in vivo by using a double transgenic mouse model of prostate cancer in collaboration with Dr. Robert Matusik at Vanderbilt University. Dr. Matusik’s group crossed c-myc overexpressing mouse model of prostate cancer with the hepsin transgenic mice. Interestingly, the hepsin/myc transgenic mice showed progression of tumorigenesis at the primary site including development of higher grade tumors as compared to the myc alone mice (Nandana, Ellwood-Yen et al. 2010). In collaboration with Dr. Matusik’s group, we studied whether the increase in tumorigenesis in the hepsin/myc transgenic mice is correlated with Ln-332 degradation. We found that Hepsin/myc tumors mice display increased degradation of Laminin-332 as compared with myc alone tumors.

In an effort to correlate increased hepsin expression with increased hepsin activity in terms of Ln-332 cleaved in vivo, we performed Western blot analysis with the tumor tissues derived from wild type, myc and hepsin/myc mice. The Western blot results showed intact Ln-332 (145 kDa), in the tissue lysate of wild type mice. The myc tumor displayed a 100 kDa band. In the hepsin/myc tumor this 100 kDa band was further diminished (Figure 3). These data correlate with the fact that hepsin cleaves laminin-332 in vitro. The tissue lysate from wild type mice showed intact Ln-332 beta3 chain whereas our data indicates that increased hepsin levels in hepsin/myc tumors are correlated with degradation of Ln-332, a component of the basement membrane that is lost during human prostate cancer.

We also performed immunohistochemical analysis for Ln-322 on tumors derived from myc mice, that develops adenocarcinoma by 6 month of age (Ellwood-Yen, Graeber et al. 2003) and
hepsin/myc mice, that develops invasive adenocarcinoma at 4.5 months and develops a higher grade adenocarcinoma compared with age-matched myc mice tumors (Nandana, Ellwood-Yen et al. 2010). We found that 12 month old hepsin/myc tumor displayed increased degradation of Ln-332 as compared with age-matched myc tumor. Although, the immunohistochemistry data (Figure 4) are not very clear as we see more staining of Ln-332 on hepsin/myc mice as compared to myc mice but it is possible that the staining appears to be more due to diffused Ln-332.

**Task 2B. Determine the role of Hepsin in the formation of prostate cancer bone lesions and elucidate the role of cleavage of Ln-332 by Hepsin on the formation of bone lesions *in vivo* (Months 24-36).**

We have not addressed task 2B partly or fully as yet.
Figure 1: DU145 prostate cancer cells exhibit no change in cell adhesion on hepsin-cleaved Ln-332. Ln-332 alone (L), Ln-332 + hepsin (LH), Ln-332 + hepsin + KD1 and PBS were coated onto 96-well plates. DU145 cells were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained, dried and solubilized; and absorbance was read at 560 nm. Results represent mean ± S.D. (N= 2, in duplicates; P < 0.05).

Figure 2: Hepsin overexpressing LNCaP cells exhibit no change in cell adhesion on Ln-332. Ln-332 or PBS was coated onto 96-well plates. LNCaP17 and LNCaP cells were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained (crystal violet), and solubilized; and absorbance was read at 560 nm. Results represent mean ± S.D. (N= 3, in triplicates; P < 0.05).

Both Cell types were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained (crystal violet), and solubilized; and absorbance was read at 560 nm. We found that hepsin-overexpressing LNCaP-34 cells had no significant difference in cell adhesion on Ln-332 as compared to low hepsin-expressing LNCaP-17 cells. Results represent mean ± S.D. (N= 3, in triplicates; P < 0.05).
Figure 3: DU145 cells exhibit no change in cell adhesion on matriptase-cleaved Ln-332. Ln-332 alone (L), Ln-332 + matriptase (LM), Ln-332 + matriptase + KD1 (LMI) and PBS were coated onto 96-well plates. DU145 cells were allowed to adhere for 1 h at 37°C in serum-free medium. Unbound cells were washed off; adherent cells were fixed, stained (crystal violet), and solubilized; and absorbance was read at 560 nm. Results represent mean ± S.D. (N= 2, in duplicates; P < 0.05)

Figure 4: Hepsin/myc tumors mice display increased degradation of Laminin-332 as compared with myc alone tumors. Western blot analysis showing Laminin-332 β3 chain expression in 12 month wildtype, myc and hepsin/myc mice. Western blot is representative of the experiment performed three times, two mice for each group has been used. Lane WT shows intact Laminin-332 in 12 month wildtype mice (145 kDa). Lanes Myc and Myc/Hep show cleaved Laminin-332 (100kDa) in 12 month myc and hepsin/myc mice respectively.
Figure 5: Immunohistochemical staining of tumor sections of lateral prostate of hepsin/myc and Myc overexpressing Mouse models of Prostate Cancer. Panel A and B showing hepsin staining in hepsin/myc (A) and myc (B) mouse and panel C and D showing Ln-332 staining in hepsin/myc (C) and myc (D) mouse.

**Key Research Accomplishments:**
1) We found that matriptase proteolytically processes Ln-332. Western blot and mass spectrometry indicated that proteolysis occurs in the β3 chain of Ln-332. Substrate specificity was confirmed by blocking cleavage with a matriptase inhibitor, Kunitz domain-1 (KD1). Transwell migration assays showed that DU145 prostate cancer cell motility was significantly enhanced when plated on matriptase-cleaved Ln-332. Similarly, Transwell migration of matriptase-overexpressing LNCaP prostate cancer cells was increased on Ln-332 and, as determined by live single-cell microscopy, two of their motility parameters, speed and directional persistence, were also higher. These results were published in The Prostate Journal (Manisha Tripathi, Alka Potdar, Hironobu Yamashita, Brandy Weidow Peter T. Cummings, Daniel Kirchhofer, and Vito Quaranta. Laminin-332 cleavage by Matriptase alters motility parameters of prostate cancer cells. Prostate. 2011 Feb 1;71(2):184-96 Cell adhesion of DU145 cells exhibit no significant difference on hepsin-cleaved Ln-332.

2) Cell adhesion of hepsin-overexpressing prostate cancer cells exhibit no significant difference on Ln-332.

3) Cell adhesion of DU145 cells exhibit no significant difference on matriptase-cleaved Ln-332.

4) Hepsin/myc tumors mice display increased degradation of Laminin-332 as compared with myc alone tumors.

**Reportable outcomes:**


2) **Poster Presentation:** Manisha Tripathi, Alka Potdar, Hironobu Yamashita, Brandy Weidow Peter T. Cummings, Daniel Kirchhofer, and Vito Quaranta. Laminin-332 cleavage by
matriptase alters motility parameters of prostate cancer cells. Symposium on Basement Membranes in Tissue Development and Regeneration, July 7-9 2010, Center for matrix biology, Vanderbilt University, Nashville TN.


4) **Oral Presentation:** May 2010: Role of Proteolytic Processing of Laminin-332 by Type II Transmembrane Serine Proteases, Hepsin and Matriptase in Prostate Cancer Progression. Presented at UCLA, Los Angeles, California.

**Conclusion:**

1) Hepsin and matriptase cleave Ln-332 in vitro;

2) The cleavage site of Ln-332 is located on the N-terminus of the domain VI of beta 3 chain;

3) This processing of Ln-332 by hepsin and matriptase is inhibited by an inhibitor of these proteinases;

4) Migration of DU145 and LNCap prostate cancer cells is increased on hepsin- and matriptase-cleaved Ln-332 in vitro;

5) Matriptase overexpression causes increased persistence of LNCap cell migration on Ln-332;

6) Cell adhesion of prostate cancer cells is not affected by hepsin or matriptase cleavage of Ln-332;

7) Hepsin overexpression induces Ln-332 degradation in vivo.

**So what section:**
In summary, we have identified a new substrate for matriptase and hepsin, the processing of which has been directly linked to increased cell migration \textit{in vitro} and correlated with increased tumorigenesis \textit{in vivo}. Our findings contribute to unraveling the roles of matriptase and hepsin in prostate cancer. By understanding the mechanisms of action of these two proteinases, we can better target their inhibition, towards developing novel therapeutic strategies in the cure of human prostate cancer. In a broader prospective, our study should promote additional studies aimed at molecular mechanisms of interaction between epithelial cells and their immediate microenvironment, the basement membrane.

\textbf{References:


\textbf{Appendices

Ln-332 Laminin-332 (Laminin-5)

TTSP Type II transmembrane serine proteases

BM Basement membrane

ECM Extracellular matrix

mRNA Messenger ribonucleic acid

PBS Phosphate Buffered Saline

siRNA Small interfering RNA

SDS Sodium dodecyl sulphate
FITC fluorescein isothio- Cyanate
Laminin-332 Cleavage by Matriptase Alters Motility Parameters of Prostate Cancer Cells

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BACKGROUND. Matriptase, a type II transmembrane serine protease, has been linked to initiation and promotion of epidermal carcinogenesis in a murine model, suggesting that deregulation of its role in epithelia contributes to transformation. In human prostate cancer, matriptase expression correlates with progression. It is therefore of interest to determine how matriptase may contribute to epithelial neoplastic progression. One approach for studying this is to identify potential matriptase substrates involved in epithelial integrity and/or transformation like the extracellular matrix macromolecule, laminin-332 (Ln-332), which is found in the basement membrane of many epithelia, including prostate. Proteolytic processing of Ln-332 regulates cell motility of both normal and transformed cells, which has implications in cancer progression.

METHODS. In vitro cleavage experiments were performed with purified Ln-332 protein and matriptase. Western blotting, enzyme inhibition assays, and mass spectrometry were used to confirm cleavage events. Matriptase overexpressing LNCaP prostate cancer cells were generated and included in Transwell migration assays and single cell motility assays, along with other prostate cells.

RESULTS. We report that matriptase proteolytically cleaves Ln-332 in the β3 chain. Substrate specificity was confirmed by blocking cleavage with the matriptase inhibitor, Kunitz domain-1. Transwell migration assays showed that DU145 cell motility was significantly enhanced when plated on matriptase-cleaved Ln-332. Similarly, Transwell migration of matriptase-overexpressing LNCaP cells was significantly increased on Ln-332 and, as determined by live single-cell microscopy, two motility parameters of this cell line, speed and directional persistence, were also higher.


KEY WORDS: laminin-332; matriptase; type II transmembrane serine protease; proteolysis; prostate cancer; cell migration

INTRODUCTION

The American Cancer Society estimated that there would be ~192,280 new prostate cancer cases in the USA in 2009 alone [1]. In general, prostate cancer morbidity and mortality is due to metastasis, or the spread of cancer cells from their primary site to other tissues or organs in the body [2–5]. In order for cells to metastasize, they must first degrade nearby tissue barrier, the basal lamina (BL), and then migrate to secondary sites via the bloodstream or lymphatic...
Degradation of BL takes place by cleavage of its extracellular matrix (ECM) components, like collagens and laminins, by enzymes like matrix metalloproteinases (MMPs) or type II transmembrane serine proteases (TTSPs) [5,6]. These protease–ECM interactions have also been shown to contribute to cell migration, a critical component of invasion and metastasis [7–9].

One essential component of BL is laminin-332 (Ln-332; previously known as Ln-5 [10]), which is known to play an important role in development, wound healing, and tumorigenesis [11]. Ln-332 is reportedly overexpressed in many tumor types, including: esophageal, cutaneous, oral, laryngeal, colon, tracheal, and cervical cancers [12,13]. Interestingly, Ln-332 expression is instead decreased or lost in prostate cancer [14–17]. As depicted in Figure 1A, Ln-332 is a large multi-domain glycoprotein consisting of α3, β3, and γ2 subunits. It has a cross-shaped structure, with its long arm consisting of domains I and II, which holds the intact molecule together [13]. At its C-terminus, the α3 chain has five large globular (LG) domains that interact with various cell surface receptors, including integrins α3β1, α6β4, and heparin proteoglycan syndecans, which leads to establishment of cell adhesion and migration phenotype [9,13,18,19].

Through its N-terminus, the β3 chain of Ln-332 interacts with other ECM molecules like Ln-6, Ln-7, and collagen VII, affecting BL assembly and cell-survival signaling [9,13]. The γ2 chain contains epidermal growth factor (EGF)-like domains III-V [13] and interacts with epidermal growth factor receptor (EGFR), which also stimulates cell migration [9,20,21]. Of note, several studies have shown that all three Ln-332 chains can be processed by different protease systems [9]. In addition, some reports have suggested that these proteolytic events regulate cell motility of both normal and transformed cells [13], which may have implications in cancer progression.

Previously, our group has shown that Ln-332 γ2 chain is cleaved by MMP2 and MT1-MMP [22,23]; others have shown that this chain is also cleaved by MMPs 3, 8, 12, 13, 14, and 20 [24], cathepsin S [25], mTLD [26], BMP-1 [27], and neutrophil elastase [28]. Furthermore, most of these studies have shown that processing of Ln-332 γ2 chain by these proteases regulates cell migration [22,24,28]. Studies have also shown that Ln-332 β3 chain is processed by MT1-MMP, which enhances prostate cancer cell migration compared to activity on uncleaved Ln-332 [29]. Another study reported that Ln-332 β3 chain is a ligand for MMP7, which enhances cell motility of a colon carcinoma cell line [30]. It has also been reported that Ln-332 β3 chain is cleaved at its N-terminus by endogenous proteinase(s) in human keratinocytes and other cell lines [31]; however, the specific protease involved in this cleavage has not been pinpointed.

Directly relevant to this study, we previously demonstrated that Ln-332 β3 chain is also cleaved by hepsin, one of 20 known TTSPs, which results in increased migration of prostate cancer cells in vitro [32]. Taken together, these studies establish that proteolytic processing of Ln-332 occurs physiologically and can alter cell adhesion and migration phenotype [9,13,18,19].

![Fig. 1](image-url)

**Fig. 1.** Ln-332 is cleaved by matriptase. A: Schematic of Ln-332 structure, which is composed of α3, β3, and γ2 chains. B: Purified Ln-332 from 804G rat bladder cells (0.2 μM) was incubated with the recombinant protease domain of matriptase (0.6, 2, and 6 μM) for 2 hr. After incubation, the mixtures and Ln-332 alone were electrophoresed on SDS–PAGE and then stained with Coomassie blue. Ln-332 alone revealed four primary bands representing the α3 (190 kDa), β3 (145 kDa), and γ2 chains (155 and 80 kDa; lane 1). Of note, a unique band was resolved at ~100 kDa in the lanes containing Ln-332 and matriptase, particularly at higher concentrations of the latter (lanes 3 and 4, arrows). C: Ln-332 (0.8 μM) and matriptase (24 μM) were coincubated for 0, 3, 6, and 12 hr. No cleavage product of Ln-332 was observed from the 0 hr mixture, however, the cleaved ~100 kDa band was present in lanes containing the mixtures from 3, 6, and 12 hr (arrow).
The Prostate

In this report, we demonstrate for the first time that matriptase is an important step in targeting this protease further identification of critical substrates for matriptase-activated receptor-2 [46,50–55]. However, (transmembrane and associated with src kinases), and plasminogen activator, prostasin zymogen, Trask phage-stimulating protein 1 (MSP-1), urokinase-type plasminogen activator (uPA) and matriptase can cleave the pro form of HGF, macrophage-derived chemokine, and correlates with disease progression [36–38]. In addition, Forbs et al. [39] reported that inhibition of matriptase in prostate cancer cells by siRNA reduces their invasive growth potential in vitro; this group also reported a similar effect using a synthetic matriptase inhibitor. Another recent study found that a small molecule inhibitor against matriptase reduced growth of tumors in prostate cancer xenograft models [40]; these authors of this report also showed that tumor growth inhibition was through the attenuation of cancer cell invasion, rather than cell proliferation. A similar study showed reduction of cell migration and invasion using both in vitro and xenograft models by inhibiting matriptase by siRNA [41]. Matriptase is inhibited by the Kunitz domain (KD) of hepatocyte growth factor (HGF) activator inhibitor-1 (HAI-1), which is a Kunitz-type transmembrane serine protease inhibitor [42]. Using this inhibitor, it has been shown that inhibition of HAI-1 expression in prostate cancer cells results in increased cell invasion and migration in vitro [43]. Most recently, the strong oncogenic potential of matriptase has been firmly established by a report that showed transgenic mice with matriptase (MT-SP1, TADG-15, epithin, and ST-14), would also cleave Ln-332 and in turn might have similar biological significance. The importance of matriptase in physiology is underscored by the fact that matriptase-deficient mice die shortly after birth due to a severely impaired water barrier function in the epidermis of skin and oral epithelium [35]. Furthermore, matriptase has recently been recognized as a potential marker for prostate cancer progression [36], as many studies have shown that matriptase expression is significantly increased in prostate tumor samples compared to normal tissue, and correlates with disease progression [36–38]. In addition, Forbs et al. [39] reported that inhibition of matriptase in prostate cancer cells by siRNA reduces their invasive growth potential in vitro; this group also reported a similar effect using a synthetic matriptase inhibitor. Another recent study found that a small molecule inhibitor against matriptase reduced growth of tumors in prostate cancer xenograft models [40]; these authors of this report also showed that tumor growth inhibition was through the attenuation of cancer cell invasion, rather than cell proliferation. A similar study showed reduction of cell migration and invasion using both in vitro and xenograft models by inhibiting matriptase by siRNA [41]. Matriptase is inhibited by the Kunitz domain (KD) of hepatocyte growth factor (HGF) activator inhibitor-1 (HAI-1), which is a Kunitz-type transmembrane serine protease inhibitor [42]. Using this inhibitor, it has been shown that inhibition of HAI-1 expression in prostate cancer cells results in increased cell invasion and migration in vitro [43]. Most recently, the strong oncogenic potential of matriptase has been firmly established by a report that showed transgenic mice with matriptase overexpression in the skin exhibited malignant transformation and had potentiated chemical carcinogenesis [44].

One of the key regulatory mechanisms of physiological and pathological functions by matriptase is by proteolytically processing or cleaving its substrates [45–49]. A few studies have demonstrated that matriptase can cleave the pro form of HGF, macrophage-stimulating protein 1 (MSP-1), urokinase-type plasminogen activator, prostasin zymogen, Trask (transmembrane and associated with src kinases), and protease-activated receptor-2 [46,50–55]. However, further identification of critical substrates for matriptase is an important step in targeting this protease therapeutically for the treatment of prostate cancer. In this report, we demonstrate for the first time that matriptase cleaves Ln-332, which affects prostate cancer cell migration parameters. This study provides new insight into possible mechanisms for matriptase and Ln-332 in cancer progression.

MATERIALS AND METHODS

Cell Culture

The human prostate cancer cell line DU145 (American Type Culture Collection, Manassas, VA) and the rat bladder squamous carcinoma cell line 804G [43] were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Life Technologies, Inc., Rockville, MD) supplemented with 10% fetal bovine serum (FBS; Gemini, Irvine, CA) and 1% glutamine/penicillin/streptomycin (g/p/s) antibiotics (Life Technologies, Inc.) in a humidified atmosphere containing 5% CO2 at 37°C.

LNCaP prostate cancer cells expressing low (LNCaP-wt) and high levels (LNCaP-mt) of matriptase were created by Dr. Daniel Kirchhofer as detailed below (Genentech, San Francisco, CA), and were cultured in RPMI 1640 medium supplemented with 10% FBS, 500 μg/ml Geneticin (Invitrogen, Carlsbad, CA), 0.5 μg/ml puromycin (Sigma, St. Louis, MO), and 1% g/p/s antibiotics and incubated with 5% CO2 at 37°C.

Purification of Rat Ln-332

Ln-332 was purified from spent medium of 804G rat bladder squamous carcinoma cells as previously described [32].

Cleavage of Ln-332 by Matriptase

To study the cleavage of Ln-332 by matriptase, purified rat Ln-332 (0.2 μM) was incubated with the human recombinant protease domain of matriptase (0.6, 2, and 6 μM). The reaction was performed in a buffer containing 250 mM NaCl and 50 mM Tris (pH 7.5) for 2–3 hr (as indicated) at 37°C. For the time course experiment, Ln-332 (0.8 μM) was incubated with matriptase (24 μM) and reaction buffer containing 250 mM NaCl and 50 mM Tris, pH 7.5, for 0, 3, 6, and 12 hr at 37°C. After incubation, sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) was performed under reducing conditions on the matriptase and Ln-332 mixture (4–12% precasted gradient gels were used). Protein bands were visualized using SimplyBlue™ Safe Coomassie Blue stain (Invitrogen). Precision Plus™ protein dual color standard (BioRad, Hercules, CA) was used as a protein marker for comparison.
Western blot analysis was performed after transferring the reaction mixtures of Ln-332 with and without matriptase onto a PVDF membrane (Perkin Elmer, Waltham, MA) from the reducing gel. Polyclonal antibody (pAb) against the C-terminus of Ln-332 β3 chain (1:500; sc-20775; H-300; Santa Cruz Biotechnology, Inc., Santa Cruz, CA), and secondary anti-rabbit IgG HRP monoclonal antibody (1:5,000; GE Healthcare, UK) were used for Western blots. Protein bands were visualized with an ECL Plus system (Perkin Elmer). ImageJ was used for quantification of bands in the scanned Western blot film. Briefly, in the “Analyze” function of ImageJ, we “Set Measurements” for area, mean gray value, and integrated density. The bands of interest were selected and the parameters were measured. The raw intensity measurement for each band was normalized to its corresponding β-actin control.

**Mass Spectrometry**

The cleaved product of Ln-332 by matriptase was identified using mass spectrometry analysis performed by the Mass Spectrometry Research Center at Vanderbilt University (Nashville, TN). Briefly, after digestion, the proteins in the reaction mixture were separated by SDS–PAGE and visualized using Coomassie Blue stain. The protein bands of interest were excised from the SDS–PAGE gel. After further processing, trypsin digestion was performed and peptides were extracted and concentrated. For the preparation of sample for matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS), 0.4 μl of the sample was applied to a target plate and overlaid with 0.4 μl alpha-cyano-4-hydroxycinnamic acid matrix (10 mg/ml in 60% acetonitrile, 0.1% trifluoroacetic acid). MALDI-TOF MS and tandem TOF/TOFMS/MS were performed using a Voyager 4700 mass spectrometer (Applied Biosystems, Foster City, CA). TOF/TOF fragmentation spectra were acquired in a data-dependent fashion based on the MALDI-TOF peptide mass map for the protein. Both types of mass spectral data were collectively used to examine the protein databases to generate statistically significant candidate identification using GPS Explorer software (Applied Biosystems) running the MASCOT database search algorithm (Matrix Science, Inc., Boston, MA). Searches were performed against the SWISS PROT and the NCBI databases.

**Matriptase Inhibition Assay**

HAI-1-derived KD1 was used to inhibit enzymatic matriptase activity. KD1 is the N-terminal Kunitz domain of HAI-1 and was produced in *Escherichia coli* as described in Shia et al. [42]. As described in the

**Generation of Matriptase Overexpressing LNCaP Cells**

The cDNA of full-length matriptase was inserted into a mammalian expression vector containing the puromycin resistance gene for antibiotic selection (Genentech). The LNCaP-luc clone [56] was transfected with the construct encoding full-length matriptase with a C-terminal FLAG tag, and the cells were selected with 0.5 μg/ml puromycin (Sigma–Aldrich). Clones were analyzed by FACS for matriptase surface expression using an anti-FLAG monoclonal antibody (Sigma–Aldrich). Two clones, one for the high matriptase expressor LNCaP-mt cells and one for the low matriptase expressor LNCaP-wt cells, were selected for further experiments.

**LNCaP Cell Protein Isolation and Western Blot**

LNCaP-wt and LNCaP-mt cells were lysed with RIPA buffer (25 mM Tris, pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 1% sodium deoxycholate, and 0.1% SDS) containing protease inhibitor cocktail (Roche, Indianapolis, IN). Total protein levels of samples were measured using a bicinchoninic acid assay (Pierce, Thermo Fisher Scientific, Rockford, IL) and normalized. Denatured proteins were separated by SDS–PAGE and transferred for Western blotting. Non-specific binding to membranes was blocked for 1 hr with blocking buffer (5% milk in PBS). Blots were incubated overnight in primary antibody (1:1,000 pAb matriptase; Bethyl Laboratories, Inc., Montgomery, TX) or 1:1,000 monoclonal anti β-actin antibody (AC-74, Sigma) in blocking solution at 4°C, and subsequently in HRP conjugated anti-rabbit IgG and anti-mouse IgG secondary antibody (NA493v; GE Healthcare) for 1 hr at room temperature. Protein bands were visualized with an ECL Plus system (Amersham Pharmacia Biotech, Piscataway, NJ).

**Transwell Cell Migration Assay**

Modified Boyden chamber cell migration assays were performed using 8.0 μm pore size Transwell™ permeable supports (Corning Costar, Lowell, MA). The lower side of the filters were coated with either untreated or matriptase-treated rat Ln-332 with and without KD1, phosphate-buffered saline solution (PBS), or Ln-332 coincubated with matriptase and
KD1 overnight at 4°C. Transwells were then blocked with 5% milk in PBS for 1 hr. Cells (DU145 or LNCaP) were trypsinized, resuspended in serum-free medium, and washed twice with serum-free medium. DU145 or LNCaP cells (20,000 or 50,000, respectively) were seeded in the upper chamber of inserts. After 5 hr (DU145) or 24 hr (LNCaP) incubation in cell culture conditions, cells remaining on the upper filter were scraped off using a cotton swab and the inserts were washed with PBS. Cells that migrated to the lower chamber were fixed with 400 μl of fixation solution (Hema-3® stain kit, cat. # 122-911, Fisher Scientific Company LLC, Kalamazoo, MI) for 10 min, and stained with 400 μl of staining solution for 20 min. Cells were manually counted using an inverted microscope equipped with a 10× objective (Zeiss, Germany). Results are presented as the mean number of cells counted per field ± standard deviation. Student’s t-tests were performed on final data to test significance of effects, with P < 0.05 accepted as significant.

**Single Cell Motility Assay**

LNCaP-wt or LNCaP-mt cells were plated (20,000 cells) overnight on 60-mm dishes coated with Ln-332 (10 μg/ml). Cell density was kept low to avoid interacting cell populations. Cells were monitored using phase-contrast optics in a Zeiss Axiovert 200 M inverted microscope with a monochrome, cooled CCD camera (CoolSNAP HQ, Roper Scientific, Trenton, NJ) equipped with a temperature-controlled, humidified chamber. All cells were equilibrated in the humidified, temperature-controlled (37°C) microscope chamber for 30 min and media was replaced with fresh growth media prior to data collection. Images were both acquired and cells tracked using Metamorph (Molecular Devices Corporation, Sunnyvale, CA). Time-lapse images were collected at a magnification of 10× (1 pixel = 0.98 μm) using a sampling time interval of 1 min, for at least 12 hr in all experiments.

**Cell Tracking**

Each cell was tracked by following the cell nucleus using the “track objects” function in Metamorph. Only single cells were considered for the analysis; cells that remained stationary, moved outside the viewing area, underwent cell division, did not migrate over a distance of at least two cell bodies (<20 μm), or that adhered to other cells during the course of the experiment were excluded from the tracking procedure. Applying this criterion, ~60% cells were retained. Results are presented in combined box-and-whisker and scatter plots, which show the mean speed per population (dark horizontal line), 25th and 75th quartiles (box), 95% confidence intervals (whiskers), and raw data points (scatter). Mann–Whitney U-tests were performed on final data to test significance of effects, with P < 0.05 accepted as significant.

**RESULTS**

**Ln-332 Is Cleaved by Matriptase**

To determine whether matriptase cleaves Ln-332, purified Ln-332 from 804G rat bladder cells was incubated with three concentrations of the recombinant protease domain of matriptase for 2 hr. After incubation, the mixtures and Ln-332 alone were electrophoresed on SDS–PAGE and then stained with Coomassie Blue. Ln-332 alone revealed four primary bands representing the α3 (190 kDa), β3 (145 kDa), and γ2 chains (155 and 80 kDa; Fig. 1B, lane 1) of the structure and matriptase alone revealed one strong band at ~27.5 kDa (lane 5). Of note, the lanes containing Ln-332 and matriptase mixes revealed a unique band at ~100 kDa, particularly at higher concentrations of the latter (lanes 3 and 4, arrows). These lanes also resolved bands at ~27.5, ~18, and ~10 kDa, which were determined to be matriptase by mass spectrometry (data not shown). An additional SDS–PAGE gel that was instead run for 3 hr is shown in Supplemental Figure 1, which confirms the presence of a unique band at ~100 kDa after incubating Ln-332 with matriptase. These results suggest that the recombinant protease domain of matriptase cleaved Ln-332.

To determine if this cleavage was time-dependent, we also performed a time-course experiment whereby Ln-332 and matriptase were coincubated for 0, 3, 6, and 12 hr and the mixtures were again resolved using SDS–PAGE. As expected, no cleavage product of Ln-332 was observed in the 0 hr mixture, however, the cleaved ~100 kDa band was present in lanes containing the mixtures from 3, 6, and 12 hr (Fig. 1C, arrow).

**Matriptase Cleaves the Ln-332 β3 Chain**

To determine that the cleavage event of Ln-332 was due to addition of matriptase, and not another contamination protease, we also added a known inhibitor of matriptase, KD1, to the mixture. As in earlier experiments, purified rat Ln-332 alone revealed four bands, representing its respective chains (Fig. 2A, lane 1) and Ln-332 treated with matriptase again revealed the ~100 kDa cleavage product (lane 2). Of interest, the mixtures containing KD1 lacked the ~100 kDa cleavage product (lanes 3, 4; arrow), suggesting that matriptase was responsible for cleavage of
Matriptase was again resolved as an ~27.5 kDa band, which matches up with the previous SDS–PAGE results, and KD1 was resolved as an ~11 kDa band, in line with our previous report [32].

To further determine the identity of the unique band that appeared after matriptase treatment of Ln-332, we used both Western blotting with an antibody to Ln-332 and a proteomics approach. As shown in Figure 2B, a pAb against the C-terminal sequence of Ln-332 β3 chain reacted with both the full-length β3 chain in Ln-332 alone and the ~100 kDa matriptase-cleaved fragment. This result suggests that matriptase cleaves the β3 chain of Ln-332, possibly removing an N-terminal sequence. To explore this, we then performed mass spectrometric analysis. The protein bands of Ln-332 and the ~100 kDa product were excised from a gel from SDS–PAGE. After trypsin digestion, MALDI-TOF MS and tandem TOF/TOF MS were performed, and data from both methods were collectively used to examine the protein databases. Statistically significant candidates were identified using GPS Explorer software running the MASCOT database search algorithm. Searches were performed against the SWISS PROT and NCBI databases. The Ln-332 protein band contained its three chains: α3 (190 kDa), β3 (145 kDa), and γ2 (155 kDa), and the digested ~100 kDa band that appeared upon treatment of Ln-332 with matriptase produced 19 different peptides (gray) that were clearly identical to amino acid sequences of Ln-332 β3 chain.

Migration of DU145 Cells Is Enhanced on Ln-332 Cleaved by Matriptase

Since previous studies have shown that cleavage of the Ln-332 β3 chain by other proteases leads to changes in cell migration [9,32], we also investigated the effect of the cleavage of Ln-332 β3 chain by matriptase on prostate cancer cell migration in this study. First, we examined motility of DU145 cells using modified Boyden chambers treated with either untreated Ln-332, matriptase-cleaved Ln-332, PBS, or a mixture containing Ln-332, matriptase, and KD1. The number of cells that passed through filters after 5 hr were then manually counted under a microscope. As shown in Figure 3, cells seeded in chambers coated with matriptase-cleaved Ln-332 migrated significantly more (~1.6-fold) than cells on uncleaved Ln-332. In addition, cells in chambers treated with the Ln-332, matriptase, and KD1 inhibitor mixture migrated significantly less than cells on matriptase-cleaved Ln-332, which was similar to migration levels on untreated Ln-332.

Fig. 2. Matriptase cleaves the Ln-332 β3 chain. A: A known matriptase inhibitor, KD1, was added to the Ln-332 and matriptase mixture. Purified rat Ln-332 (0.2 μM) alone revealed four bands, representing its respective chains (lane 1). Ln-332 (0.2 μM) treated with matriptase (6 μM) again revealed the ~100 kDa cleavage product (lane 2). The mixture also containing KD1 lacked the ~100 kDa cleavage product (lanes 3 and 4). B: A Western blot was performed using a pAb against the C-terminal sequence of Ln-332 β3 chain. The antibody reacted with both the full-length β3 chain in Ln-332 alone and the ~100 kDa matriptase-cleaved fragment. C: Mass spectrometry was performed to analyze the contents of the protein bands of Ln-332 and the ~100 kDa product from Ln-332 treated with matriptase. This analysis revealed that the Ln-332 protein band contained its three chains: α3 (190 kDa), β3 (145 kDa), and γ2 (155 kDa), and the digested ~100 kDa band that appeared upon treatment of Ln-332 with matriptase produced 19 different peptides (gray) that were clearly identical to amino acid sequences of Ln-332 β3 chain.
Since our experiments indicated that DU145 cell migration was increased on matriptase-cleaved Ln-332, we decided to also examine migration of LNCaP prostate cancer cells that stably overexpress matriptase (LNCaP-mt). Migration of these overexpressor cells was compared to that of wild-type LNCaP cells (LNCaP-wt) first using modified Boyden chamber assays. Prior to performing assays, we verified LNCaP cell expression by RT-PCR (data not shown) and Western blot (Fig. 4A); these results indicated that LNCaP-mt cells expressed \(\approx 2\) times more matriptase than LNCaP-wt cells. For Boyden chamber assays, inserts were pretreated with either Ln-332 or PBS, and cells were allowed to migrate for 24 hr prior to counting the number of cells that passed through filters. As expected, LNCaP-mt cells exhibited significantly more migration than LNCaP-wt cells on Ln-332 (\(~3\)-fold; Fig. 4B,C). In contrast, both cell types migrated minimally on PBS-treated Transwells.

**Matriptase-Overexpression Enhances LNCaP Cell Migration on Ln-332**

In order to acquire more detailed information about the motility of LNCaP cells, we also performed single-cell motility assays using high content microscopy. This technique involves tracking individual cell movement over time using video microscopy. In contrast to static Boyden chamber assays, this approach allows inspection of dynamic cell movement in real-time. Further, single-cell level parameters can help to model and predict population level behavior.

Time-lapse movies of LNCaP motility can be found in Supplementary Materials. In line with previous Boyden chamber results, the LNCaP-mt population moved significantly faster than LNCaP-wt cells using this technique (Fig. 5A). In addition, LNCaP-mt cells also moved in a more directed manner than LNCaP-wt cells, leading to significantly increased directionality ratios for this cell type (Fig. 5B). This ratio represents the *linear* distance a cell travels during an assay (d) versus the *total* distance traveled by that cell (t), which essentially captures cell persistence, or the tendency of a cell to continue moving in a particular direction without turning [57].

We also created Windrose plots by overlaying single cell tracks onto a single origin \((0,0)\), in order to qualitatively assess the persistence for each cell type (Fig. 5C). Twelve-hour trajectories of LNCaP-wt (gray) and LNCaP-mt (black) cells are shown, which indicate that matriptase overexpressing cells are generally more persistent and travel farther than wt cells. The concentric circles superimposed on the plots indicate the root mean squared displacement (MSD) for each cell population after 12 hr. This value was
obtained by first calculating the MSD for each cell population, using Equation (1), where \( \mathbf{r}(t) \) is the position vector of the cell after time \( t \), \( \mathbf{r}(0) \) is the position at the beginning, and \( \langle \cdot \rangle \) denotes the average over the entire cell population, followed by taking the square root of the MSD.

\[
\text{MSD} = \langle (\mathbf{r}(t) - \mathbf{r}(0))^2 \rangle \quad \text{RMSD} = \sqrt{\text{MSD}} \quad (1)
\]

Fig. 4. Matriptase-overexpression enhances LNCaP cell migration on Ln-332. A: Matriptase expression of both types of LNCaP cells was confirmed by Western blot. Band intensity for each sample was normalized to its corresponding \( \beta \)-actin control. These results indicated that LNCaP-mt cells express \( \sim 2 \) times more matriptase than LNCaP-wt cells. B: Boyden chamber assay was used to examine cell migration of both LNCaP-wt and LNCaP-mt cell types. Transwells were coated with either Ln-332 (10 \( \mu \)g/ml) or PBS, cells were allowed to migrate for 24 hr, and cells that migrated across the filter were fixed, stained, and counted manually under a microscope. C: LNCaP-mt cells exhibited significantly more migration than LNCaP-wt cells on Ln-332 (\( \sim 3 \)-fold; \( N = 3 \), in duplicate; \( P < 0.001 \)). In contrast, both cell types migrated minimally on PBS-treated inserts.

Fig. 5. Matriptase-overexpression enhances cell speed and directional persistence. A: Single-cell motility assays were performed using high content microscopy. Individual cell speed was measured using Metamorph software. The LNCaP-mt cell population moved significantly faster than LNCaP-wt cells using this technique (\( N = 32 \) and 35 cells, respectively; \( P = 0.006 \)). Box and whisker plots show the population mean (bold horizontal line), 25th and 75th quartiles (box), and 95% confidence intervals (whiskers) overlaid on raw single-cell data (scatter). B: LNCaP-mt cells were also found to move in a more directed manner than LNCaP-wt cells, leading to significantly increased directionality ratios for this cell type, which represents the linear distance a cell travels during an assay (d) versus the total distance traveled by that cell (t). C: Windrose plots were made to qualitatively examine the persistence for these cell types; these plots overlay all cell tracks (from \( x \) and \( y \) coordinates) starting with a common origin (0,0). Twelve-hour trajectories of LNCaP-wt (gray) and LNCaP-mt (black) cells are shown, which indicate that, in general, the matriptase overexpressing cells are more persistent and travel further than wt cells. The circles superimposed on the Windrose plots indicate the root mean squared dispersal for each cell type.
In summary, these results show that the LNCaP-mt cell population covered a greater area than the LNCaP-wt population. Taken together, all single-cell motility parameters indicate that matriptase overexpressing cells exhibit a different migration phenotype than wild-type cells.

DISCUSSION

In this report, we studied two important steps of the metastatic cascade: breach of BL and cell migration [58]. Specifically, we have focused on investigating the interactions and effects of two potentially important players in prostate cancer progression: Ln-332, an essential BL component, and matriptase, 1 of 20 members of an emerging TTSP family [59,60]. The role of Ln-332 in prostate cancer has been of special interest because, in contrast to other tumors where Ln-332 is overexpressed, Ln-332 is instead reduced or lost in this type of cancer [14–16]. In contrast, matriptase is reportedly overexpressed in many cancers, including prostate, and its expression has been shown to correlate with disease progression [45,47–49]. Intuitively, this inverse expression of Ln-332 and matriptase in prostate cancer suggests a potential interaction between these molecules during disease progression.

It has previously been shown that overexpression of matriptase in mouse epidermis induces spontaneous skin lesions in the absence of genetic alteration and independent of carcinogen exposure [44]. Furthermore, the authors showed that epidermal hyperproliferation and matriptase-induced tumors were abolished by coexpression of HAI-1. However, it remains to be determined how matriptase leads to neoplastic progression in such mouse models [44]. Matriptase is reportedly overexpressed in a wide variety of epithelial tumors, including breast, cervix, esophagus, liver, mesothelium, prostate, and colorectal cancers [36,38,61–69]. Interestingly, in the case of prostate cancer there is also correlation between matriptase expression and tumor grade [36–38]. The mechanism of action of matriptase remains to be determined, however, the possibilities include activation of growth factors, receptors, proteases, and the processing of ECM components. Identification of new substrates of matriptase and definition of their biological function(s) upon cleavage contribute in unraveling the mechanistic action of matriptase in cancer.

Our results show, for the first time, that Ln-332 is proteolytically processed by matriptase in vitro. Mass spectrometry and Western blotting analyses indicated that catalytically active recombinant matriptase cleaves the β3 chain of Ln-332 and produces a novel ~100 kDa fragment (Figs. 1 and 2). Our previous work has shown that Ln-332 is also a substrate of another TTSP, hepsin [32]. Interestingly, both matriptase and hepsin cleave at the N-terminal of the Ln-332 β3 chain [32]. Proteolysis has been shown to be an important step in both physiological and pathological conditions [6,70]. One of the most widely reported effects of cleavage of Ln-332 has been increased cell migration [9,13]. It has been reported that MMPs generate a domain, DIII, from Ln-332 and binding of this domain to EGFR stimulates mitogen-activated protein kinase signaling, MMP-2 gene expression, and cell migration [20]. Also, it has been shown that processing of γ2 chain may influence Ln-332 turnover in BL and affect epithelial morphogenesis [23]. Although there are other protease systems that cleave Ln-332, we propose that the effects of cell surface proteases, including the family of TTSPs, have more physiological consequences than other protease systems due to their physiological positioning on the cell surface [55]. This locale gives such cell surface proteases access to ECM components, giving them an added advantage over other protease systems. In future studies, it would be worth screening whether all identified TTSPs possess the ability to cleave Ln-332, particularly at the β3 chain. In addition, it would be interesting to determine whether the cleavage site is conserved across species.

Our previous studies revealed that hepsin cleavage of Ln-332 increased cell migration [32], therefore we also investigated this function in this report. Modified Boyden chamber assays revealed that the human prostate cancer cell line, DU145, migrated significantly more on Ln-332 cleaved by matriptase than on intact Ln-332 substrate (Fig. 3). In line with this result, LNCaP prostate cancer cells overexpressing matriptase also migrated significantly more on Ln-332 than wild-type cells with a lower level of matriptase in Boyden chambers (Fig. 4). LNCaP cell motility was also analyzed at the single-cell level using high content microscopy for more in-depth study of cell dynamics. Interestingly, we found that single cells from both cell types exhibited variable speeds significantly different than the average for each population (Fig. 5), highlighting the importance of single-cell techniques for examining in-depth behavior. Furthermore, this analysis revealed that matriptase overexpressing cells migrated faster and more persistently than their corresponding control cells.

Cell migration studies are largely conducted at the population level (e.g., Boyden chamber), whereby a cell population is represented by an average measurement and some range of error. It is thought that in vivo single cell migration is required for metastasis through blood, whereas cohesive migration is required for lymphatic metastasis [71]. In this study, we investigated cell motility both at the population level and at the single-cell level using time-lapse video microscopy for slightly
different purposes. Boyden chamber assays captured an end-point population level behavior of cells seeded at a high density. In contrast, single cell assays were performed at a lower cell density, and data were collected dynamically and assessed at both the single-cell and population levels. Recently, it was reported that two different phenotypic outcomes for cell migration may occur in vitro, dependent upon whether an investigator uses single-cell or cohesive migration strategies [72]. In other words, studying migration using these two different approaches investigates two different questions. In our case, change(s) in the migratory phenotype due to cleavage of Ln-332 by matriptase produced a similar result using both approaches. Intuitively, the increased number of LNCaP-mt cells that crossed through Boyden chambers can be explained by the single cell results, which show that matriptase overexpressing cells have increased speed and persistence on Ln-332. Persistence in cell motion can be defined as the property by which a cell continues to migrate in one direction without much deviation (before changing its path) [57]. Studies have shown that cancer cell migration is directionally persistent, as in the cases of highly invasive cancers like neuroepithelial tumors [73] or epithelial cell over-expressing HER2 [74]. In addition, genetic modifications of biological molecules in a cell line have previously been shown to change the intrinsic pattern of cell migration [57]. However, to our knowledge, our report is the first study to show that overexpression of a cell surface protease leads to increased cell speed and persistence of cells plated on its substrates. Further analysis of single-cell assays must be performed to better understand whether the parameters obtained at this level can help in predicting their population level behavior.

Due to the limited availability of human Ln-332, most of the studies in this field (including ours) have been carried out using rat Ln-332. However, due to the functional interchangeability of ECM components across mammalian species, we are confident that one would see similar processing of human Ln-332 by matriptase. Interestingly, both hepsin and matriptase cleave the N-terminal of the Ln-332 β3 chain. Therefore, it is possible that cleavage of Ln-332 by these two TTSs affects the interaction of Ln-332 with collagen VII, which in turn may effect hemidesmosome formation and tumor invasion, which has been previously reported [75]. Future studies should also be carried out to investigate this possibility.

We realize that the enzyme/substrate ratio used in this study is quite high. However, our rationale for choosing these ratios are based on the fact that Ln-332 is a bulky molecule with a molecular weight of 490 kDa, whereas the enzyme used in this study, the protease domain of matriptase, is only 27.5 kDa. For these reasons, many other previous studies using MMPs have also been performed at higher ratios in vitro [19,21,25,27]. As mentioned above, under physiological conditions, the TTSPs, including matriptase, are in close vicinity of the BM, thus they are also close to the substrate Ln-332.

CONCLUSIONS

In summary, we show that matriptase cleaves Ln-332 and this cleavage event results in increased migration of prostate cancer cells. LNCaP cells over-expressing matriptase migrated faster and in a more directionally persistent manner on Ln-332. Based on our findings, we propose that proteolytic processing of Ln-332 could be a possible mechanistic role for matriptase in prostate cancer progression via altered migration parameters and subsequent BL transgression. Therefore, identification of Ln-332 as one of the substrates for matriptase takes us one step closer to unraveling the mechanism(s) of action of matriptase.

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