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TITLE: The Role of Stromally Produced Cathepsin D in Promoting Prostate Tumorigenesis

PRINCIPAL INVESTIGATOR: Freddie Pruitt

CONTRACTING ORGANIZATION: Vanderbilt University
Nashville, TN. 37240-0001

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**ABSTRACT**

Stromal-epithelial interactions are important in development and cancer of the prostate. Estrogen receptor functions as a transcription factor to regulate gene expression. One such ER-regulated gene is the protease cathepsin D (CathD). *This proposal has two immediate objectives. The first is to determine how overexpression of cyclin D1 (CD1) in the stroma induces the upregulation of the estrogen regulated gene CathD. The second objective is to determine how overexpression of CathD in the stroma can contribute to tumorigenesis in the epithelium.* Results show 1) CD1 interacts with the ERα to modify the expression of estrogen regulated genes like CathD in prostate fibroblasts. 2) ER signaling in the stroma contributing to CAFs induced tumorigenesis in adjacent epithelium. 3) Stromal specific overexpression of CathD promotes prostate tumorigenesis through activation of TGFβ signaling pathways.

**SUBJECT TERMS**

Cathepsin D, Cyclin D1, Estrogen Receptor, hormonal carcinogenesis, stroma, prostate.
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Introduction

Stromal-epithelial interactions are important in development and cancer of the prostate. Estrogen receptor functions as a transcription factor to regulate gene expression. One such ER-regulated gene is the protease cathepsin D (CathD). This project had two immediate objectives. The first was to determine how overexpression of cyclin D1 (CD1) in the stroma induces the upregulation of the estrogen regulated gene CathD. The second objective is to determine how overexpression of CathD in the stroma can contribute to tumorigenesis in the epithelium. We believe that changes in the stroma result in alterations in stromal-to-epithelial paracrine signaling. This altered environment promotes the initiation and progression of tumorigenesis.
Keywords

Cathepsin D, Cylin D1, Estrogen Receptor, Hormonal Carcinogenesis, Prostate, Stroma
Overall Project Summary

The original stated goal in the approved statement of work for Task 1 was to determine if CD1 overexpression modifies estrogen regulated genes through interaction with endogenous ERα in prostate fibroblasts. To accomplish this task we developed a benign human prostate stromal cell line BHPṛS to co-express CD1 and ERα to use as tool for examining the interaction of CD1 with ERα on the CathD promoter. In figure 1A, Co-immunoprecipitation experiments were performed to determine if CD1 could interact with ERα in prostate stromal cells. Co-overexpressing BHPṛS$^{CD1-ERα}$ cells demonstrate CD1 and ERα can interact when ectopically expressed. Overexpression of CD1 in BHPṛS alone demonstrated interaction with endogenous ERα in prostate stromal cells. We next examined if CD1 overexpression modifies ERα transcriptional activity. In figure 1B, luciferase activity assays were performed with the use of estrogen responsive element fused with the luciferase gene. These experiments were performed in presence/absence of β-estradiol. In comparison to the empty vector control BHPṛS$^{EV}$, CD1 overexpression induced increased ER transcriptional activity in the absence of hormone. This indicates that CD1 can induce the transcriptional activity of the ER with out the ligand bound in prostate stromal cells. This finding is in agreement with studies of ER and CD1 in breast cancer (1). Co-overexpression of ERα and CD1 in BHPṛS cells induced greater luciferase expression compared to CD1 overexpression alone, both in the presence and absence of hormone. These data show that CD1 can interact with ERα to drive transcriptional activity of the estrogen receptor on non-chromosomal DNA. To determine if the CD1-ERα interaction binds chromosomal DNA we performed chromatin immunoprecipitation (ChIP) experiments with the BHPṛS$^{CD1}$ and BHPṛS$^{CD1-ERα}$ cell lines. Overexpression of CD1 in BHPṛS showed greater than 11 fold recruitment over the IgG control of CD1 to the estrogen receptor element (ERE) in the CathD gene in the absence of hormone. In the presence of hormone CD1 recruitment to the ERE in the CathD gene was only increased 8 fold over the IgG control.
We have previously shown that CD1 overexpression in benign prostate fibroblast cells (NPF CD1) produces a phenotype similar cancer associated fibroblasts (2). A comparison of gene expression profiles from NPF CD1 and CAFs cells identified CathD to be overexpressed in both cell types in comparison to NPFs. Task 1 from the approved statement of work sought to determine if CD1 overexpression modifies estrogen regulated genes through interaction with endogenous ERα in prostate fibroblasts. Our experimental approach showed that CD1 overexpression leads to aberrant ERα activity on the CathD gene.

Our original stated goal in the approved statement of work for Task 2 was to determine if inhibition of the ERα in signaling in the stroma abrogates the ability of CAF and/or NPF cyclin D1 to induce tumorigenesis. Our rational for this task is based on the understanding of the hormone levels in aging men. Levels of circulating testosterone in younger men are greater in comparison levels of estrogen. After 30 years of age, the hormone ratios in men start to change, with older men having lower levels of circulating testosterone and greater levels of estrogen(3). In figure 2, we examined the expression of ERα in human prostate clinical samples using a tissue microarray which contained 30 cases of adenocarcinoma, 5 cases of normal prostate tissue, and 5 cases of normal prostate tissue adjacent to malignant tissue (NAT). The tissue array contained duplicate cores per case.
Quantification of nuclear ERα in stromal regions was significantly greater in malignant prostate tissue in comparison to normal prostate tissue. In figure 3A, and 3B, gene expression analysis of ERα and aromatase in primary fibroblasts isolated from NPF and CAFs showed significantly greater expression in CAFs in comparison to NPFs with a greater than 5 fold difference. Due to our observations of increased ERα expression in CAFs and in malignant human tissue, we questioned the role of the ER in the cancer associated stroma’s ability to promote transformation in tissue recombination experiments.

Knockout animal models have been used to examine the role of the estrogen receptor in prostate. Recombination experiments with ERβ knockout mesenchyme with wild type epithelium produced squamous metaplasia (SQM) when supplemented with a synthetic estrogen, however, SQM was not observed in ERα knockout mesenchyme was combined with wild type epithelium, indicating the importance of ERα in the stroma (4). In human PCa tissue, the expression of the ERα is up regulated in the stroma in comparison to benign prostate stroma (5) The co-administration of Testosterone (T) with Estrogen (E) stimulates cancer progression and malignant transformation in tissue recombination experiments of rat UGM (rUGM) and BPH-1, however, the single administration of T alone induces benign growth (6,7). In the majority of models for hormonal carcinogenesis, estrogen along with the actions of the ER are required for maximal carcinogenic response to androgens.

To accomplish our approved second task we stated CAF and NPFCD1 cells would be engineered with shRNA specific to ERα. We attempted to transduce our different fibroblast cells with the shRNA constructs to make stable cell lines, however, CAFs and NPFCD1 cells under went replicative senescence. A downside to working with non-immortalized cells, is the limitation to the number of cell cycle replications. To overcome this
pitfall and still address the second task we decided to use a pharmacologic approach and inhibit the ER in signaling with the use of Tamoxifen. In figure 4, we recombined CAFs with the initiated but non-transformed prostate epithelial cell line BPH-1 or NPF with BPH-1 and performed xenograft experiments in mice supplemented with T, E, T + E, T + Tamoxifen, and E + Tamoxifen for a period of 8 weeks. Recombinations of NPF + BPH-1 in mice supplemented with T produced benign growths with no malignant conversions. CAF + BPH-1 in mice supplemented with T induced a malignant conversions denoted by the BPH-1 cells invading into the mouse kidney. Recombinations of NPF+ BPH-1 in mice supplemented with T + E resulted in histologies resembling CAF + BPH-1 recombinations. The addition of estrogen in combination with T, drove a malignant conversion in the adjacent epithelium. This result is similar to published findings of tissue recombinations experiments of rUGM with BPH-1.

In figure 5 we quantitated the distance the transformed BPH-1 cells invaded into the kidney of the mouse. A comparison of BPH-1 cell invasion into the kidney in recombinations with CAFs shows a significant decrease in mice treated with tamoxifen. These results suggest that the activation of the ER in the stroma plays a critical role in prostate tumor progression. From our knowledge of the biosynthesis of sex steroid hormones, we know that testosterone can be directly converted into estrogen by the aromatase enzyme. The co-administration of Testosterone (T) with Estrogen (E) stimulates cancer progression and malignant transformation in tissue recombination experiments of rat UGM (rUGM) and BPH-1, however, the single administration of T alone induces benign growth (5,6). In the majority of models for hormonal carcinogenesis, estrogen along with the actions of the ER are required for maximal carcinogenic response to androgens. Recombinations of NPF + BPH-1 in mice supplemented with T produced benign growths with no malignant conversions. CAF + BPH-1
in mice supplemented with T induced a malignant conversions denoted by the BPH-1 cells invading into the mouse kidney. Recombinations of NPF + BPH-1 in mice supplemented with T + E resulted in histologies resembling CAF + BPH-1 recombinations. The addition of estrogen in combination with T, drove a malignant conversion in the adjacent epithelium. This result is similar to published findings of tissue recombinations experiments of rUGM with BPH-1.

Since we observed increased expression of ERα in the stroma of malignant prostate tissue, we examined the consequences of ERα modulation in stromal cells. To address this question we engineered BHPPrS cells to stably overexpress ERα (BHPPrSERα). As a control, we also overexpressed ERβ in BHPPrS (BHPPrSERβ). We prepared tissue recombinations using the prostate epithelial cell line BPH-1 as a reporter with the BHPPrSERα and BHPPrSERβ stromal lines and grafted under the renal capsule in castrated SCID mice supplemented with a combination of 17β-estradiol and testosterone (T), estrogen (E), and testosterone + estrogen (T+E). As a control we also used rUGM recombined with BPH-1 cells and grafted under the renal capsule in castrated mice supplemented with a combination of 17β-estradiol and testosterone pellets, testosterone alone, or no treatment (NT). As a control we also used rUGM recombined with BPH-1 cells and grafted under the renal capsule in castrated mice supplemented with a combination of 17β-estradiol and testosterone pellets, testosterone alone, or no treatment (NT).

<table>
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<th>Treatments</th>
<th>rUGM</th>
<th>EV</th>
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<tr>
<td>NT</td>
<td>No Growth</td>
<td>No Growth</td>
<td>No Growth</td>
<td>No Growth</td>
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<tr>
<td>T</td>
<td>Small Benign Growth</td>
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<td>T + E</td>
<td>Large Benign Tumors</td>
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Table 1. Results of various recombinations of BPH-1 cells with rUGM, BHPPrSEV (EV), BHPPrSERα (ERα), and BHPPrSERβ (ERβ), supplemented with the different hormones testosterone (T), estrogen (E), and testosterone + estrogen (T+E).

Significance determined by ANOVA, p-value ≤ 0.05.

Our study highlights human ERα expression in the cancer associated stroma. The overexpression of the ERα in benign human prostate stromal cells promotes the expression of markers associated with the reactive stromal phenotype. We see that overexpression of ERα specifically in the stroma is sufficient to drive tumorigenesis and induce the malignant conversion of initiated, but non-transformed prostate epithelial cells. We also show CAFs express relatively higher levels of ERα and the enzyme aromatase in comparison to NAFs. We believe the increased expression of aromatase aids in producing a local estrogen rich environment.
promoting the development of prostate tumorigenesis. Lastly this second task identifies stromal ERα as a target for therapeutic intervention by showing pharmacologic inhibition of the ER in the cancer associated stroma inhibits malignant transformation in the adjacent epithelium.

Our final approved task was to determine how stromal production of CathD promotes tumorigenesis. The overexpression of CathD in neoplastic cells and neoplastic associated connective tissue was described as long as 25 years ago, and is reported to play several roles in cancer progression (8-11). To accomplish this task we engineered BHPs cells to overexpress CathD (BHPsCathD) to perform in vitro and in vivo experiments. Sub-renal capsule xenograft experiments were performed using the BHPsCathD recombined with BPH-1 cells. After a period of 8 weeks, overexpression of CathD in the stroma induced a malignant transformation in the adjacent epithelium along with significantly larger tumors in comparison to recombinations with the empty vector control BHPs cells. The reverse experiment was performed using BPH-1 cells made to overexpress CathD (BPH-1CathD) and recombined with parental BHPs cells and grafted under the kidney capsule. After a period of 8 weeks, epithelial overexpression of CathD failed to induce significant differences in growth or a malignant transformation. Our model identifies that stromal specific expression of CathD plays a role in promoting tumorigenesis. Our model is further supported by our findings of CathD knockdown expression in CAFs abolishes the ability to induce a malignant transformation in adjacent epithelium (12).

To delve deeper into understanding the role of stroma specific expression of CathD promoting tumorigenesis, we examined growth factor signaling pathways in the BPH-1 cells treated with conditioned media from BHPsCathD. Conditioned media collected from BHPsCathD and BHPsEV was added to monolayer cultures of BPH-1 cells prior to isolation of protein lysates. Western blot analysis was performed on BPH-1 cell lysates for changes in the growth factor signaling pathways MAPK, and Akt/PKB. No differences were observed in the phosphorylation states of MAP kinases p38 of ERK1/2, however, we did observe increased p-Akt levels in BPH-1 cells supplemented with BHPsCathD conditioned medium.

Our laboratory has previously published a study identifying a possible mechanism for CAFs to induce tumorigenesis in the adjacent epithelium. The study identified two molecules, transforming growth factor-beta (TGF-β) and stromal cell derived factor-1 (SDF-1) as being overexpressed in CAF cells. These factors were

![Figure 7. Stromal specific expression of CathD promotes malignant transformation. Tissue recombination experiments of A) BHPs overexpressing CathD recombined with BPH-1 cells, B) BPH-1 cells overexpressing CathD recombined with parental BHPs cells. Green hashmarks delineate graft-kidney interface. Red asterisk denotes enlarged inlay showing SV40 expressing BPH-1 cells invading into mouse kidney.](image-url)
acting in a paracrine manor on the epithelial cells resulting in Akt hyperactivation \((13)\). We questioned whether the increased levels of p-Akt seen in our experiments with the conditioned media isolated from BHPrSCathD cells was related to TGF-\(\beta\) activity. It has been previously shown in \textit{in vitro} experiments that CathD can liberate TGF-\(\beta\) from the latency inhibitor complex, allowing for activation of the TGF\(\beta\) receptor (TGFBR) complex \((14)\). In vivo experiments using the CathD prozyme, showed CathD can be fully activated in the extracellular environment and cleave substrates in the mouse prostate \((15)\). We examined our tissue recombinations of BHPrSCathD with BPH-1 cells for increased TGF-\(\beta\) signaling. Immunohistochemical staining for p-SMAD2/3, the immediate downstream substrate of TGFBR activation, showed significantly greater p-SMAD2/3 staining in CathD overexpressing recombinants. We also examined differences in the TGF-\(\beta\) responsive gene collagen type 4 \(\alpha\)2 (ColIV\(\alpha\)2). Staining for showed increased deposition of ColIV\(\alpha\)2 in recombinations overexpressing CathD. Masson’s trichrome staining of tissue xenografts from the CathD overexpressing stromal cells also revealed increased production of collagen in comparison to recombinations with the EV control stromal cells. Collectively we concluded that stromal derived CathD was promoting tumorigenesis through the activation of TGF-\(\beta\) signaling pathways.

The second line item under the approved third task was to determine if the conversion of pro-CathD to the active protease is dependent on the presence of glycosaminoglycans (GAGs) on the surface of PCa cells. To accomplish this task we treated a series of PCa cell lines with heparinase to remove heparin sulfate containing proteoglycans from the cell surface prior to the addition of pro-CathD. The heparin sulfate proteoglycans have been previously shown to convert the CathD zymogen to the active state \((14)\). Completion of the CathD activity assay after treatment with heparin lyase did not show any inhibition in the conversion of pro-CathD to the active protease. We also performed this experiment with 5-(N-ethyl-N-isopropyl)-amiloride (EIPA), a specific inhibitor for Na\(+\)/H\(\(+\)\) antiporters. It has been shown that human PCa and PCa cell lines have lower extracellular pH due to increased proton pump expression \((16)\). We hypothesized that CathD activity was dependent on the activity of proton pumps present on the surface of PCa cells. PCa cell lines treated with EIPA show drastic decreases in extracellular CathD activity similar to levels when of cell lines treated with the CathD inhibitor pepstatin A. Our results show that extracellular CathD activity is not dependent on heperan sulfate containing proteoglycans, however is dependent on the extracellular pH.
Figure 9. Stromal expression of CathD leads to activation of TGF-β. Overexpression of CathD in BHPrS (E) results in malignant transformation (D) through increased TGF-β signaling leading to increased p-SMAD 2/3 activation (K), and downstream TGF-β responsive genes like collagen type IVα2 (L). Masson’s trichrome staining reveals a reactive stromal phenotype, denoted by increased collagen deposition in BHPrSCathD recombinations (J).
Figure 10. Extracellular CathD activity is dependent on proton pump exchangers. Deactivation of proton exchangers on benign and malignant prostate cell lines with EIPA inhibits CathD proteolytic cleavage on fluorescent substrate.
Key Research Accomplishments

- Prostate stromal cell lines were generated and characterized.
- ERα is overexpressed in the stroma of malignant human prostate tissue in comparison to benign.
- ERα and aromatase expression is upregulated in patient derived cancer associated fibroblasts.
- Pharmacologic inhibition of the ER prevents malignant conversion of adjacent prostate epithelial cells in vivo.
- Overexpression of ERα in benign prostate stromal cells produces robust tumorigenesis and malignant conversion of adjacent epithelial cells in vivo

Conclusions

The aims executed in this grant provided advances in the field of PCa biology by 1) identifying expansion of ERα positive cells in PCa reactive stroma and in patient derived primary cancer associated fibroblasts cells. 2) Showing pharmacological inhibition of the ERα abrogates epithelial malignant conversion by the cancer associated stroma. 3) Showing that ER responsive genes, i.e. CathD, are expressed in the reactive stroma of malignant prostate tissue and promotes prostate tumorigenesis. We have also identified a possible mechanism to explain how stromal derived CathD overexpression can promote tumorigenesis through TGF-β signaling. These data further demonstrate stromal ERα role in prostate tumorigenesis and highlight ERα as a therapeutic target for chemoprevention. Significant progress has been made towards achieving the stated goals considering the technical limitations experienced in the second aim. This work has produced one publication.

Publications

Pruitt, FL., He, Y., Franco, OE., Jiang, M., Cates, JM., Hayward, SW. Cathepsin D acts as an essential mediator to promote malignancy of benign prostatic epithelium. The Prostate. 73(5):476-88, April, 2013. PMID: 22996917
Abstracts

2011 Annual Meeting for the Society of Basic Urologic Research. Atlanta, GA.
The interaction of Cyclin D1 with ERα in stromal fibroblast promotes prostate tumorigenesis.

Stromal-epithelial interactions are important in the development of the prostate, and have also been shown to be important in the development of prostate cancer. Stromal changes have been shown to be powerful prognostic indicators of prostate cancer progression and of patient death. Expression of estrogen receptor-alpha (ERα) in the prostate is localized to the stroma and becomes elevated during the progression of PCa. The ER functions as a transcription factor and regulates the expression of a number of different genes. One such ER-regulated gene is the aspartic endopeptidase cathepsin D (CathD). In various malignancies e.g. breast, and colon cancers, CathD is overexpressed and hypersecreted. Our laboratory has previously demonstrated that several molecules found to be aberrantly expressed in cancer associated fibroblasts (CAFs) (including cyclin D1 [CD1], TGFβ and SDF-1) induce tumorigenesis and malignant transformation in tissue recombination experiments. The cell cycle regulator CD1 and the ERα are known to interact and can induce estrogenic gene transcription in breast cancer studies. We investigated whether CD1 can interact with ERα to modify stromally produced CathD expression. We also wanted to determine whether CathD expression affected prostate tumorigenesis. Gene reporter assay experiments using CD1 overexpressing benign human prostate stromal (BHPrSCD1) cells show increased ER transcriptional activity in the absence of estradiol. Co-immunoprecipitation experiments show CD1 interacts with endogenous ERα in BHPrS cells and increases CathD expression and secretion. A comparison of benign human prostate tissue and prostate tumor revealed increased stromal staining of ERα, CD1 and CathD in tumor bearing prostate samples. Recombination experiments using normal prostate fibroblast (NPF) with BPH-1 reporter epithelial cells supplemented with estradiol and testosterone induces a malignant transformation with strong CathD staining. This malignant transformation is inhibited in mice supplemented with tamoxifen. Knockdown of CathD expression in CD1 overexpressing fibroblast (NPFCD1) or CAF abrogates the tumorigenic response of the epithelial reporter cell line BPH-1. Our preliminary findings suggest that CathD is overexpressed and hypersecreted through an interaction with ERα and CD1. The overexpression of CD1 and CathD appears to play an important role in prostate tumorigenesis and may represent a possible therapeutic target.

*2012 Annual Meeting for the Society of Basic Urologic Research. Las Vegas, NV.

Stromal specific cathepsin D overexpression promotes prostate tumorigenesis through activation of TGF-β signaling pathways in adjacent epithelium.

Stromal-epithelial interactions are important in both development and prostate cancer. Stromal changes have been shown to be powerful prognostic indicators of prostate cancer progression and of patient death helping to define lethal versus indolent phenotypes. The specific molecular underpinnings of these interactions are incompletely understood. Our laboratory has previously demonstrated that several molecules found to be aberrantly expressed in cancer associated fibroblasts (CAFs) (including cyclin D1 [CD1], TGFβ) induce tumorigenesis and malignant transformation in xenograft experiments. The cell cycle regulator CD1 and the estrogen receptor are known to interact and can induce estrogenic gene transcription. Cathepsin D (CathD) is an estrogen regulated aspartic endopeptidase, known to be involved in a number of physiological processes as well as in the regulation of apoptosis. In various malignancies e.g. breast, CathD expression is associated with a poor prognosis and increased likelihood for the development of metastasis. There are limited data defining CathD’s function in prostate cancer and in disease progression. In the present study, we highlight CathD as a
mediator of cancer associated stromal promotion of prostate tumorigenesis. An examination of human prostate tissue revealed significantly increased stromal staining of CathD in malignant prostate tissue in comparison to benign prostate tissue. High stromal expression of CathD in CAFs affects the expression of cell cycle regulators in adjacent epithelium. Knockdown of CathD expression in CAFs inhibits malignant transformation in tissue recombination experiments. Stromal specific overexpression of CathD in benign prostate stromal cells induced malignancy in adjacent epithelium through increased TGFβ signaling and responsive gene expression. Whereas epithelial cell specific overexpression of CathD produced benign structures in tissue recombination experiments. The proteolytic function of stromal derived CathD is dependent on the activity of hydrogen-proton pump activity on the surface of prostate epithelial cell lines. The study presented here indicates that CathD is not only an important mediator of stroma-epithelial cross talk, but also an essential component in promotion of tumorigenesis \textit{in vivo}.

\textbf{Inventions, Patents and Licenses}

The work performed for this award did not result in any patentable inventions or any licenses.

\textbf{Reportable Outcomes}

This award generated a 2013 publication in the peer reviewed journal “\textit{The Prostate}”.

\textbf{Other Achievements}

Two human benign prostate stromal cell lines (BHPrS) were generated that stably overexpress ERα, and ERβ. These cell lines are very useful tools for understanding the roles of these sex hormone receptors in the context of prostate biology. This award supported my studies for the completion of my doctoral degree in Cancer Biology from Vanderbilt University Medical center in 2013.
References

The Prostate

Cathepsin D Acts as an Essential Mediator to Promote Malignancy of Benign Prostatic Epithelium

Freddie L. Pruitt,1 Yue He,1 Omar E. Franco,2 Ming Jiang,2 Justin M. Cates,3 and Simon W. Hayward1,2,4*

1Department of Cancer Biology, Vanderbilt University Medical Center, Nashville, Tennessee
2Department of Urologic Surgery, Vanderbilt University Medical Center, Nashville, Tennessee
3Department of Pathology, Vanderbilt University Medical Center, Nashville, Tennessee
4Department of Vanderbilt-Ingram Cancer Center, Vanderbilt University Medical Center, Nashville, Tennessee

BACKGROUND. Stromal–epithelial interactions are important in both development and prostate cancer. Stromal changes have been shown to be powerful prognostic indicators of prostate cancer progression and of patient death helping to define lethal versus indolent phenotypes. The specific molecular underpinnings of these interactions are incompletely understood. We investigated whether stromal cathepsin D (CathD) overexpression affects prostate tumorigenesis through a paracrine mechanism.

METHODS. Normal prostate fibroblasts (NPF) were retrovirally transduced to overexpress cyclin D1 (CD1) and were designated NPFCD1. Cathepsin D expression was knocked down using shRNA in cancer associated fibroblasts (CAF) and NPFCD1. We analyzed these stromal cell lines using immunohistochemistry, Western blot, and tissue recombination.

RESULTS. An examination of human prostate tissue revealed significantly increased stromal staining of CathD in malignant prostate tissue. Overexpression of CD1 in normal prostate fibroblasts (NPFCD1) produced a phenotype similar to, but more moderate than, CAF in a tissue recombination model. Knockdown studies revealed that CathD is required for NPFCD1 motility and invasive growth in vitro. BPH-1 cell proliferation was found to be induced when cultured with NPFCD1 conditioned medium, this effect was inhibited when CathD was knocked down in NPFCD1 cells. Overexpression of CathD in prostate stromal cells induced malignancy in adjacent epithelium, and this transformation was inhibited when stromal CathD expression was knocked down in CAF.

CONCLUSIONS. The study presented here demonstrates increased CathD expression is seen in human CAF. The upregulation of CD1 results in concomitant increases in CathD expression. Elevated CathD expression in the stroma contributes to tumor promotion.

KEY WORDS: cathepsin D; stromal–epithelial interactions; prostate cancer

INTRODUCTION

Historically, the field of cancer biology has primarily been focused on studying the malignant tumor epithelium [1]. The emergence of the field of tumor microenvironment is providing some much needed insight into how non-malignant cells associated with cancer (cancer associated stroma) can promote or suppress tumorigenesis. The stromal phenotype has been shown to be a powerful prognostic indicator of cancer progression and of patient death underlining the...
importance of local stromal cells in defining lethal versus indolent phenotypes [2].

Stromal–epithelial interactions are important in both the development of the prostate, and in prostate cancer (PCa) [3–5]. During carcinoma evolution, the stroma cells adjacent to the pre-malignant or malignant epithelium experience phenotypic alterations that have been shown to enhance the invasive potential of the epithelial tumor [6–8]. These stromal–epithelial interactions are mediated, in large part, by paracrine signaling between epithelial tumor cells and neighboring stromal fibroblasts [8]. We have previously published on several molecules found to be aberrantly expressed in cancer-associated stroma that induce tumorigenesis and malignant conversion [7–13]. These intercellular interactions are clearly complex and there are likely a number of molecular routes which can either promote or suppress tumor-inducing activity. One purpose of pursuing these studies is to start to determine the identity of pathways which are either sufficient or necessary to induce transformation and to examine how such pathways might interact. We showed that the up-regulation of a cell cycle regulator known as cyclin D1 (CD1) in normal prostate fibroblasts mimics aspects of the phenotype of malignant conversion seen in cancer associated stroma. In addition to receiving signals from malignant epithelial cells, the stromal fibroblasts stimulate tumorigenesis by releasing factors that act on adjacent epithelial tumor cells or exchange enzymes that modify local microenvironment promoting the proliferation and survival of the neoplastic cells [7,8,14,15].

One mechanism by which modifications to the local tumor microenvironment are accomplished is via the actions of several different families of proteases produced by either the tumor or the stroma [16]. These enzyme families include matrix metalloproteases (MMP), cysteine, and serine proteases, which have been shown to play a role in the degradation of the basement matrix, promotion of angiogenesis, and the liberation of growth factors to stimulate tumor cell growth [17,18].

Cathepsin D (CathD) is a ubiquitous lysosomal aspartic endoproteinase. CathD has been shown to be involved in a number of physiological processes, playing a critical role in barrier function, regulation of apoptosis, and epithelial differentiation [19–21]. In cancer, CathD is overexpressed and hypersecreted in various malignancies including PCa [22,23]. In breast cancer, CathD expression is associated with a poor prognosis and increased likelihood for the development of metastasis [24]. Experimental evidence has shown CathD can stimulate the proliferation of PCa cell lines [25].

There are limited data defining CathD’s function in prostate cancer progression. Some studies have concluded that CathD is overexpressed in the epithelium and stroma of PCa, and may promote proliferation [25,26]. Other studies have concluded that CathD produced by PCa may be inhibiting tumor growth [27,28]. In the present study, we highlight CathD as a mediator of cancer associated stromal promotion of prostate tumorigenesis.

METHODS

Cells

BPH-1 (a non-tumorigenic human prostatic epithelial cell), and its tumorigenic derivatives BPH\textsuperscript{ECAFTD} were isolated from our own stocks [29,30]. CAF cells were isolated from human prostate tumor samples and their activity validated in a tissue recombination model. The technique for the isolation of CAF is described in Olumi et al. [4] which also describes a bioassay which was used to confirm the tumor-inducing activity of the CAF used in the present study. NPF\textsuperscript{CD1} cells, which we have shown in the past to overexpress CathD, were generated as previously described [9]. Benign human prostate stromal cells (BHPrS1) were isolated from a prostate surgical sample and immortalized with hTERT as previously described [10]. Cells were maintained in RPMI 1640 (Gibco, Carlsbad, CA) with 1% antibiotic/antimycotic (Life Technologies, Grand Island, NY), and 5% Cosmic Calf Serum (CCS-HyClone, Logan, UT).

Generating Genetically Modified Cell Lines

The pSuper.Retro-control (PSR-OligoEngine, Seattle, WA) and pSuper.Retro-CD1 shRNA (PSR-CD1sh) were kindly provided by Drs. Rene Bernards and Daniel Peepier from the Netherlands Cancer Institute. The two plasmids were engineered into CAF by retroviral transduction as previously described [9]. Positive transduced cells were selected for resistance to puromycin (5 μg/ml) to generate two cell strains (CAF\textsuperscript{cram} and CAF\textsuperscript{CD1sh})\textsuperscript{11}. The pSilencer 2.1-CathD1shRNA vector was kindly provided by Dr. Daniel E. Johnson from the University of Pittsburgh Cancer Institute. PSR-cathepsin Dsh was generated by removing the CathD1sh coding sequence from pSilencer 2.1-CathD1shRNA with HindIII and BamH1 and ligated into the PSR construct. The PSR and PSR-CathDsh were engineered into CAF by retroviral infection as described previously [9]. The positive transduced cells were selected for resistance to puromycin (5 μg/ml) to generate the cell line (CAF CathD\textsuperscript{sh}). BPH-1\textsuperscript{NPF}, BPH-1\textsuperscript{ECAFTD}, and BPH-1\textsuperscript{ECAFTD}\textsuperscript{NPFCD1} cells were re-isolated from resulting growths as previously described [9].
BHPrS1 cells were engineered to overexpress CathD by lentiviral transduction (Genecopoeia, Inc., Rockville, MD). Viral supernatant was generated, centrifuged at 3,000 rpm for 5 min and passed through a 0.45 μm filter before frozen at −80°C until used. Polybrene (Sigma–Aldrich, St. Louis, MO) was added to the viral suspension at 5 μg/ml to increase the efficiency of the transduction. GFP-expressing cells were selected by fluorescence-activated cell sorting (FACS) to establish the BHPrScathD and BHPrSeV as an empty vector control.

**Western Blotting Analysis**

Cell lysates were prepared and Western blotting was performed as previously described [31]. Membranes were incubated with mouse primary antibody to PTEN (1:1,000, Santa Cruz Biotechnology, Santa Cruz, CA), Cdk2 (1:1,000, Santa Cruz), Cdk4 (1:1,000, Santa Cruz), cyclin E (1:1,000, Santa Cruz), CD1 (1:1,000, BD Biosciences Pharmingen, San Jose, CA), β-actin (1:5,000, Sigma) or CathD (1:1,000, Cell Signaling, Denvers, MA) overnight and washed with PBS–TWEEN 20 for 1 hr, and incubated with horseradish-Peroxidase linked anti-mouse secondary antibody (Amersham Biosciences, Piscataway, NJ, 1:10,000 dilution) for 1 hr. Bound antibodies were visualized using enhanced chemiluminescence Western blotting detection reagents (Amersham Biosciences).

**Tissue Recombination and Xenografting**

Rat urogenital mesenchyme (rUGM) was obtained from 18-day embryonic fetuses (plug date denoted as day 0). Urogenital sinuses were dissected from fetuses and separated into epithelial and mesenchymal components by tryptic digestion, as described previously [32]. BPH-1 + rUGM, BPH-1 + NPF, BPH-1 + NPFCD1, BPH-1 + CAF, BPH-1 + CAFpCathD, BPH-1 + CAFpCathDsh, BPH-1 + BHPrScathD, and BPH-1 + BHPrSeV tissue recombinants were made as previously described [33]. 1.0 × 10^5 epithelial cells and 2.5 × 10^5 stromal cells combined in type I rat tail collagen were used to make the recombinants. After overnight incubation, the tissue recombinants were grafted under the kidney capsule of adult male severe combined immunodeficient (SCID) mice (Harlan, Indianapolis, IN) supplemented with 25 mg testosterone pellets (PCCA, Houston, TX). All the experiments were repeated six times. Mice were sacrificed at 8 weeks and grafts were harvested, fixed, and paraffin embedded. Graft dimensions were measured using the formula: volume = width × length × depth × π/6 as described previously [10].

**Wound Healing Assays**

Confluent monolayers of NPF and NPFCD1 cells were grown in 6-well plates. Confluent cell monolayers were wounded by scratching with a pipette tip. Specific points on the wounds were identified and marked. These open areas were then inspected microscopically over time as the cells migrated in and filled the damaged area. Wounds were imaged at 0, 3, 6, and 8 hr post wounding and the cell migration rate into the wound was calculated. Experiments were performed in triplicate.

**Outgrowth Assay**

1.0 × 10^5 NPF, NPFCD1, NPFCD1-CathD control or NPFCD1-CathDsh were resuspended at 4°C in Matrigel (0.2 ml, 10 mg/ml; Becton and Dickinson), and overlaid to a previously solidified layer of Matrigel in 24-well plates. The top Matrigel layer was solidified at 37°C for 30 min and covered with culture medium containing 10% FCS (0.5 ml).

**Conditioned Medium**

NPF or NPFCD1 were seeded with 5% FCS in RPMI 1640 at a density of 5.0 × 10^5 per 75-cm² flask, allowed to grow, and attached overnight. Confluent cultures of NPF or NPFCD1 were rinsed twice in PBS and incubated for 3 days in RPMI + 0.5% FCS. The medium was collected, centrifuged, passed through a 0.45-μm filter (Millipore), and stored at −80°C for later use. Conditioned medium was thawed and diluted 1:1 with fresh DMEM + 0.5% FCS before use. BPH-1 cells were seeded at 2.0 × 10^4 per well in 6-well plates in conditioned medium. The cultures were incubated for 3 days and the total number of cells was determined by direct counting in a hemacytometer.

**Human Prostate and Prostate Tissue Microarray**

Human prostate tissue array (PR806) was obtained from US Biomax, Inc. The array contained duplicates from 30 cases of adenocarcinoma ranging in Gleason scores and 10 cases of normal prostate tissue. Normal human prostate tissue was also obtained from the Vanderbilt University Medical Center Department of Pathology.

**Histochemical and Immunohistochemical Staining**

Masson’s trichrome stain was performed as previously described using Diagnostics Accustain Masson trichrome stain kit (Sigma), Bouin’s solution (Sigma) and Weigert’s Iron Hematoxylin set (Sigma) [10]. Immunohistochemical staining was performed following a protocol that was described previously [31].
Tissue slides were then incubated with the primary antibody against CD1 (1:200, BD Biosciences Phar-mingen), CathD (1:200, Santa Cruz), p-SMAD2/3 (1:400, Santa Cruz), Col4A2 (1:200, Santa Cruz). The polyclonal rabbit or mouse immunoglobulins/bioti-nylated anti-mouse secondary antibody (DAKO, Carpentaria, CA) was incubated for 60 min after the slides were washed with PBS buffer for 1 hr. After washing the slides in PBS extensively, slides were incu-bated in ABC-HRP complex (Vector Laboratories) for 30 min. Bound antibodies were then visualized by incubation with 3,3’-diaminobenzidine tetrahydro-chloride (liquid DAB, DAKO). Slides were then rinsed extensively in tap water, counterstained with hematoxylin, and mounted.

**Immunofluorescence**

For histological analysis, 5 μm tissue sections were dewaxed, and the antigen was unmasked by heating samples in unmasking solution (Vector Laboratories). Slides were blocked in 12% BSA in PBS for 30 min at room temperature before incubating with primary antibodies against CathD and GFP (1:200, Santa Cruz). After 1 hr washing in PBS buffer, slides were incubated with secondary antibodies (1:200; AlexaFlour 488 anti-Rabbit IgG and AlexaFlour 594 anti-mouse IgG2a) for 30 min at room temperature. Slides were incubated in Hoechst 33258 (4 mg/L) for 5 min. Tissue sections were washed for 30 min in PBS, mounted, and visualized.

**Quantitative Image Analysis**

Immunostained slides were analyzed using the Ariol SL-50 automated slide scanner (Applied Imaging, San Jose, CA) to quantitate the amount of staining for CathD in the stroma of benign and malignant human prostate tissue sections. Positive staining was calculated by applying two thresholds, with one recognizing weaker brown-positive cells, and another recognizing stronger brown-positive cells. The inten-sity of the stain was calculated by masking out all non-stromal areas from the tissue section and calculating the integrated optical density of brown within the remaining area. This value was divided by the area in pixels of the brown mask to calculate the average intensity of the tissue section.

**Statistical Analysis**

Data from in vitro and in vivo are presented as the mean ± standard deviation (SD). The data was ana-lyzed using GraphPad PRISM software (La Jolla, CA). P-values less than 0.05 were considered statistically significant. Quantitated intensity of CathD expression in human prostate samples were compared with analysis of variance followed by post hoc analysis of signif-icant means by Mann–Whitney’s U-test was used in comparison to normal to tumor tissue. Post hoc analysis of significant means by Dunn’s multiple comparison test was used for the comparisons of normal tissue with low grade and high grade malignant prostate tissue. P-values less than 0.05 were consid-ered statistically significant.

**RESULTS**

**CathD Expression is Upregulated in Prostate Clinical Samples, and CathD Is Overexpressed in the Stroma of Tumorigenic Tissue Recombinants**

We examined the expression patterns of CathD in human prostate clinical samples using a tissue microarray containing 30 cases of adenocarcinoma, and 10 cases of normal prostate tissue. The tissue microarray contained duplicate cores per case. Quantification of positive CathD staining in stromal regions of prostate tissue showed significantly greater areas of CathD expression in tumor tissue in comparison to normal prostate tissue. When tumor tissue was stratified between low and high grades, a significant difference was only observed in high grade tumors compared with normal tissue with a non-significant elevation of expression in low grade tumors (Fig. 1). A similar trend with no significant difference was also observed in comparisons of low and high grade tumors. It was noteworthy that the expression of CathD apparently corresponded to areas which stained a light red color in the adjacent trichrome-stained sections. This likely indicates the presence of myofibroblastic cells in this area, which would correspond to the source of our experimental CAF. Examination of CathD expression in tissue recombinations of BPH-1 + NPF, BPH-1 + rUGM, BPH-1 + NPF\(^{CD1}\), and BPH-1 + CAF was performed by IHC. The recombinations of BPH-1 + NPF and BPH-1 + UGM isolated after 8 weeks of growth produced small growths overall which displayed solid epithelial cord structures surrounded by a muscular stroma. IHC staining displayed minimal expression of CathD in the stroma with some epitheli-al expression seen in the BPH-1 + NPF recombinants (Fig. 2A1 and A2). In marked contrast, recombinations of BPH-1 + NPF\(^{CD1}\) and BPH-1 + CAF isolated after 8 weeks produced poorly differentiated carcinoma along with areas of squamous metaplasia similar to previously published results (Fig. 2A3 and A4) [9]. Recombinations of BPH-1 + NPF\(^{CD1}\) and BPH-1 + CAF displayed strong CathD staining in the stroma and epithelium (Fig. 2A3 and A4). These results are consistent with the observations of CathD overexpression in the stroma of human PCa clinical tissues.
These data raised the question of whether the upregulation of CathD protein is a passive result of prostatic tumorigenesis or plays an active role as a paracrine mediator required to induce a malignant transformation in the adjacent prostatic epithelium.

**Stromal Expression of CD1 Affects Cell Cycle Regulators in Adjacent Epithelium**

Fluorescence-activated cell sorting of BPH-1 cells isolated from xenografts of BPH-1 + NPF, BPH-1 + CAF, and BPH-1 + NPF<sup>CD1</sup>. The resulting cells were designated BPH-1<sup>NPF</sup>, BPH-1<sup>CAF</sup>, and BPH-1<sup>NPF-CD1</sup> respectively. Densitometric analysis of band intensities from Western blots revealed that overexpression of CD1 in the local stromal cells increased the expression of the cell cycle related proteins CD1, cyclin dependent kinases-6 (CDK6) and 2 (CDK2), and CathD in BPH-1<sup>NPF-CD1</sup> over BPH-1<sup>NPF</sup> (Fig. 2B), consistent with increasing proliferative activity in these cells. Similar results were observed in BPH-1<sup>CAF</sup>. No changes in the expression CDK4, cyclin E were observed.

**CathD Is a Critical Component in NPF<sup>CD1</sup> Motility and 3D Outgrowth**

We have previously reported the abilities of NPF<sup>CD1</sup> and CAF to induce tumorigenesis in tissue recombination experiments [4,9]. In order to further investigate mechanisms underlying this malignant transformation we characterized the effects of knocking down CathD expression in NPF<sup>CD1</sup>. NPF<sup>CD1</sup> displayed enhanced motility in wound healing assays compared with control NPFs (Fig. 3A). The enhanced motility displayed by NPF<sup>CD1</sup> was significantly abrogated when CathD expression was knocked down, with the use of CathD specific shRNA (NPF<sup>CD1</sup>-CathD<sub>sh</sub>) (P-value < 0.005). These findings demonstrate that CathD plays a role in NPF<sup>CD1</sup> migration in vitro. Western blot analysis was performed to confirm the knockdown of CathD expression in CD1 overexpressing fibroblast (Fig. 3D). Overexpression of CD1 in NPF results in increased CathD expression as previously published [9]. Stable expression of stable hairpin RNA (shRNA) specific for CathD results in 66% knockdown in CathD expression. Expression of non-specific shRNA in NPF<sup>CD1</sup> does not alter CathD expression.

To further characterize the requirement for CathD expression in NPF<sup>CD1</sup>, we examined fibroblast outgrowth in 3D matrices. As shown in Figure 3B, overexpression of CD1 promoted outgrowth of normal prostatic fibroblasts embedded into Matrigel. After 14 days of culture, NPF<sup>CD1</sup> cells had adopted a stellate morphology and formed invasive colonies with protrusions sprouting into the surrounding matrix (Fig. 3B2). In contrast, normal prostatic fibroblasts presented a well-delineated spherical appearance of quiescent and/or dying cells and grew poorly, neither invading nor forming protrusions to the surrounding matrix.
matrix (Fig. 3B1). NPFCD1-CathDsh cultured in 3D matrix failed to form invasive colonies that protruded into the surrounding matrix (Fig. 3B3), unlike NPFCD1-control, which retained the ability to form invasive growth feature (Fig. 3B4). These data strongly imply a role for CathD as a factor in promoting the invasive growth of NPFCD1 cells in vitro.

**CathD Is a Paracrine Mediator of Neoplastic Epithelial Cell Growth In Vitro**

To investigate the role of CathD as a paracrine mediator of prostate epithelial cell growth, we generated conditioned media from NPF and NPFCD1 cells, and measured BPH-1 cell numbers after growth for 3 days.
Fig. 3. Cathepsin D is a critical mediator between BPH-1 cells and NPF\textsuperscript{CD\textscript{1}} in vitro. A: Wound healing assay. Confluent monolayers of NPF, NPF\textsuperscript{CD\textscript{1}}, NPF\textsuperscript{CD\textscript{1}}-control, and NPF\textsuperscript{CD\textscript{1}}-CathD \textsuperscript{sh} were scratched with a pipette tip. Bar graphs represent the mean ± SD of rate of wound closure over 8 hr period. Significance determined by ANOVA, p-value ≤ 0.005, n = 3. B: 3D outgrowth assays. NPF cell lines were embedded in matrigel and cultured for 14 days, scale bar = 100 μm. Images taken at 10× (3B, b). C: Evaluation of CathD as a paracrine mediator of growth. BPH-1 cells were treated with conditioned media collected from NPF cell lines for 3 days. Cell numbers were quantitated by direct counting, graphical representation of the mean ± SD of the experiment is shown, significance determined by ANOVA, p-value ≤ 0.005, n = 3. D: Western blot confirming knockdown of CathD in NPF, NPF\textsuperscript{CD\textscript{1}}, NPF\textsuperscript{CD\textscript{1}}-control, and NPF\textsuperscript{CD\textscript{1}}-CathD \textsuperscript{sh} cells (Top). Densitometric analysis of band intensities performed to determine knockdown efficiency (Bottom).
in the conditioned medium. Conditioned medium from NPFCD1 increased the proliferation of BPH-1 cells by 1.7-fold, when compared with medium conditioned by parental NPF (Fig. 3C). The pro-mitogenic effects from NPFCD1 conditioned medium were abrogated when CathD expression was knocked down in NPFCD1-CathDsh (P-value ≤ 0.005). These results suggest that a significant component of NPFCD1 proliferative influence toward epithelium is mediated through secreted CathD.

CathD Is an Essential Mediator of CAF Induced Tumorigenicity In Vivo

To elucidate the role of CD1 and CathD in CAF’s ability to induce tumorigenesis of BPH-1, we took a knockdown expression approach. CAF were engineered to express shRNA vectors specific for either CD1 or CathD. Western blotting was used to assess knockdown efficiency. CD1 expression was knocked down 50% and CathD expression was knocked down 95% in CAF cells (Fig. 4C). Based on gross morphology it was found that BPH-1 + CAFcyclinD1sh and BPH-1 + CAF-CathDsh recombinants formed significantly smaller grafts compared with BPH-1 + CAF grafts (p-value ≤ 0.05) (Fig. 4A and C). Histologically, BPH-1 + CAF recombinants formed adenomas in comparison to the empty vector construct (Fig. 5I).

To further clarify the role of stromal derived CathD in promoting tumorigenesis we engineered BHPrS, a benign human prostate stromal cell line, to over-express CathD (BHPrS-CathD) by lentiviral transduction. In comparison to recombinations of BPH-1 + BHPrS EV, BPH-1 + BHPrS-CathD recombinants exhibited a malignant transformation. Based on the H&E staining, recombinations of BPH-1 + BHPrS EV exhibited thick stromal regions delineating BPH-1 cells from the kidney interface (Fig. 5A). The opposite was observed in recombinations of BPH-1 + BHPrS-CathD, where BPH-1 cells are directly adjacent to kidney interface (Fig. 5D). IHC staining for CathD indicates strong stromal expression of CathD in the recombinations of BPH-1 + BHPrS-CathD (Fig. 5E). IF staining for GFP positive stromal cells (red) and CathD expression (green) show strong stromal specific expression of CathD seen in the yellow overlay (Fig. 5F).

Masson’s trichrome staining was performed on tissue sections from recombinations of BPH-1 + BHPrS EV and BPH-1 + BHPrS-CathD (Fig. 5G and J). Heavy analine blue stains indicated increased deposition of newly synthesized collagen fibrils in the CathD overexpressing recombinations (Fig. 5J). IHC staining was performed to examine the phosphorylated-SMAD2/3 (p-SMAD2/3), a surrogate reporter of transforming growth factor-beta (TGF-β) activity (Fig. 5H and K). CathD overexpressing recombinations (Fig. 5K) shows increased nuclear p-SMAD2/3 staining. Quantitation of p-SMAD2/3 positive cells indicated a significant difference in the CathD overexpressing recombinations. Type IVα2 collagen (Col.IVα2) is a known TGF-β responsive gene. We performed IHC staining for Col.IVα2 and the CathD overexpressing recombinations (Fig. 5L) displayed strong expression for Col.IVα2, in comparison to recombinations with the empty vector construct (Fig. 5I).

DISCUSSION

Tumor stroma has been implicated in the regulation of cell growth, determining metastatic potential, and impacting the outcome of therapy. Stromal–epithelial interactions in cancer been have implicated as promoting several malignancies including prostate, breast, colon, and pancreatic cancers [1,4]. The stroma is often radically changed around malignant tumors and such changes both predict prognosis and may actually contribute to disease progression [1,4]. We have previously examined the role of several molecules found to be aberrantly expressed in cancer associated stroma that induce tumorigenesis and malignant conversion [8]. We reported that CD1-overexpressing BPH-1 cells are non-tumorigenic in the presence of rUGM in tissue recombination experiments, but in contrast, the overexpression of CD1 in prostate fibroblasts induces a strong tumorigenic response in the non-malignant but genetically initiated BPH-1 cells [9]. The tumor-promoting abilities of NPFCD1 produce changes very similar to published descriptions of tissue recombinations of CAF with BPH-1 [4]. A comparison of the genetic profiles from CAF and NPFCD1 identified CathD as being upregulated sevenfold in comparison to NPF [9]. From this finding we hypothesized that CathD may be a mediator of stromal–epithelial interactions contributing to prostate tumorigenesis.

CathD overexpression in neoplastic cells and neoplasia-associated connective tissue was described as long as 24 years ago, and is reported to play several roles in cancer progression [34–37]. Cathepsins have recently been shown to be upregulated in a pancreatic tumor model and also contribute to invasive breast tumor growth [38–40]. We previously reported that CathD is upregulated in both NPFCD1 cells (which mimic CAF) as well in CAF. Microarray analysis
revealed a sevenfold increase in CathD resulting from CD1 expression in NPF. NPF<sup>CD1</sup> cells display increased motility in comparison to control NPFs in a wound healing assay which was shown to depend upon the expression of CathD [9]. Here we show that the ability of NPF<sup>CD1</sup> to survive and invade into 3D matrices was also dependent on CathD. The overexpression of CD1 in NPF produced invasive colonies with protrusions sprouting into the surrounding matrix. This invasive growth was inhibited in NPF<sup>CD1</sup> cells when CathD expression was knocked down. This finding is supported by similar results from Laurent-Matha [22] where CathD was critical for outgrowth of human fibroblast in 3D matrices. Our findings were consistent with a model in which CD1-induced overexpression of CathD resulted in increased fibroblast motility and invasion.

An immunohistochemical examination of clinical specimens revealed low levels of expression of CathD in normal prostate stromal tissue. Malignant areas

**Fig. 4.** Cyclin D1 and Cathepsin D are required for CAF induced tumorigenicity in vivo. A: Gross morphology of 2 month grafts of BPH-1 + CA<sup>PSR</sup>, BPH-1 + CAF<sup>CDish</sup>, and BPH-1 + CAF<sup>CashDish</sup>. Scale bar equal to 5 cm. B: H&E staining of BPH-1 + CA<sup>PSR</sup>, CAF<sup>CDish</sup>, or CAF<sup>CashDish</sup> recombinants formed adenosquamous carcinoma as previously described. Scale bar equal to 50 μm. C: Western blot confirming knockdown of CD1 and CathD in CAFs (left). Knockdown efficiency determined by performing densitometric analysis of Western blot represented by bar graph (middle). Quantiation of tumor volume of 2 month grafts of BPH-1 + CA<sup>PSR</sup>, BPH-1 + CAF<sup>CDish</sup>, and BPH-1 + CAF<sup>CashDish</sup> (right), graphical representation of the mean ± SD of the grafts is shown in n = 6. Significance determined by ANOVA, P-value ≤ 0.05.
showed prominent stromal expression of CathD, with the significantly greater stromal CathD expression in high grade tumor samples. IHC analysis of CathD expression in tissue recombination of BPH-1 + CAF and NPF<sup>CD1</sup> also revealed strong stromal staining in comparison to recombination of BPH-1 + NPF and BPH-1 + rUGM. CathD expression in human prostate cancer stroma correlates with shorter survival and recurrence-free periods [41]. Our experimental data establishing a link between the overexpression of CD1 with the up-regulation of CathD in prostate CAF, coupled with the similar findings in human disease, indicates a strong association between cell cycle regulation and protease expression in prostate tumorgenesis. The cell cycle regulator CD1 and the estrogen receptor alpha (ERα) are known to interact and can induce estrogenic gene transcription [42]. This suggests the possibility that the overexpression of CathD in PCa associated stroma is due to the interaction of ERα with CD1.

Further investigation into the role of CathD in the tumor microenvironment showed that CathD expression is necessary for NPF<sup>CD1</sup> cells to promote epithelial growth under in vitro conditions. The pro-mitogenic effect of NPF<sup>CD1</sup> conditioned medium on BPH-1 cells was inhibited when CathD expression was knocked-down in NPF<sup>CD1</sup>. This result mirrors published findings showing that CathD is mitogenic to PCa cell lines [25]. These data do not, of course, imply that CathD is a direct mitogen, merely that its presence results in a mitogenic environment. Given the possibility that this protease may activate latent growth factors associated with extracellular proteoglycans, an indirect mechanism is not only possible but likely.

To further pursue an underlying mechanism, we engineered BHPrS cells to overexpress CathD and combined these BHPrS<sup>CathD</sup> cells with BPH-1 and performed renal grafting experiments. Recombinations of BPH-1 cells with BHPrS resulted in benign solid...
epithelial cords similar to recombinations of NPF with BPH-1 cells [10]. However, the overexpression of CathD in BHPtS in recombination experiments with BPH-1 cells induced a malignant transformation with invasion into the mouse kidney. This is consistent with our findings with the CathD knockdown approach in experiments with the CAFs. A feature of the prostate tumor microenvironment in human disease is the expansion of myofibroblast-like cells with increased deposition of extracellular matrix proteins [18]. Masson’s trichrome staining of tissue xenografts from the CathD overexpressing stromal cells revealed increased production of collagen in tissue recombinations with the EV control stromal cells. These staining patterns were similar to previous publications that pointed towards altered TGF-β signaling. We have previously shown that the overexpression of TGF-β in BHPtS cells resulted in the development of poorly differentiated adenocarcinoma with increased deposition of collagen in tissue recombinations [10]. TGF-β is expressed by most cultured cells in an inactive form due to binding with latent complex, and activation requires the proteolytic degradation of this complex. CathD derived from fibroblast conditioned media has been shown to liberate active TGF-β from the latent complex [43]. Investigation of altered TGF-β signaling in our model revealed increased p-SMAD2/3 staining, a surrogate marker for TGF-β response, in the CathD overexpressing recombinations. Examination of Col.IVα2, a direct TGF-β responsive gene, expression in the tissue xenografts revealed increased staining for Col.IVα2. Collectively, the differences in stromal composition observed from trichrome staining can be linked to increased TGF-β signaling and responsive gene expression as a result of stromal derived CathD. The overexpression of CathD in the stroma resulted in a somewhat minor, all though sufficient, malignant transformation of initiated epithelial cells similar to the tumor inductive properties of CAF.

We previously demonstrated that NPFCD1 cells and CAF elicited permanent malignant transformation of BPH-1 cells [9,30,44]. Data from IHC of clinical tissue showed increased CathD in the stroma adjacent to malignant regions of the prostate. To address the contribution of CathD in CAF-induced tumorigenesis of BPH-1, we engineered CathD knock-down CAF. The ability of CAF to induce tumorigenesis in BPH-1 recombinations was abolished when CathD expression was knocked-down. Similar results were observed when CD1 expression was knocked down in CAF. These data indicated that CathD is not only an important mediator of stroma-epithelial cross talk in vitro, but also an essential component in promotion of tumorigenesis in vivo, at least in this model.

In summary, the study presented here demonstrates that CathD can play a role as a paracrine mediator contributing to prostate tumorigenesis. The identification of key players, such as CathD, that participate in the promotion of the tumor microenvironment contributes to our understanding of the molecular mechanisms underlying this process and may prove to be valuable for the development of novel anti-cancer therapies. Current anti-cancer therapies target the malignant epithelial cells, which progressively acquire genetic alterations during the progression of the disease [45–47]. The biggest obstacle facing clinicians treating people with cancer in general is the toxicity of treatments combined with the development of resistance to therapy. The tumor microenvironment has been shown to be more genetically stable and therefore less likely to develop resistance to novel anti-cancer therapeutics [48,49]. Since tumor promotion by the microenvironment is a function of many different signaling molecules, it should be possible to develop therapeutic strategies which appropriately modify several pathways simultaneously rather than simply attempting to totally block a single signal. This is likely to be both more effective and better tolerated, since the normal biological effects of the molecules concerned will be less affected. Further investigation is needed to explain in detail how CathD is acting. A better understanding of the complexities of CathD in the tumor microenvironment may provide targets for suppressing lethal PCA phenotypes.

REFERENCES


The Prostate
The Prostate


