Detection of Early Prostate Cancer Using a Hepsin-Targeted Imaging Agent

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Abstract

Early detection and diagnosis of prostate cancer is key to designing effective treatment strategies. Microarrays have resulted in the discovery of hepsin (HPN) as a biomarker for detection of prostate cancer. In this study, we explore the development of HPN imaging probes for detection of prostate cancer. We used phage display to isolate HPN binding peptides with 190 ± 2.2 nmol/L affinity in monomeric form and high specificity. The identified peptides were able to detect human prostate cancer on tissue microarrays and in cell-based assays. HPN-targeted imaging agents were synthesized by conjugating multiple peptides to fluorescent nanoparticles to further improve avidity through multivalency and to improve pharmacokinetics. When injected into mouse xenograft models, HPN-targeted nanoparticles bound specifically to HPN-expressing LNCaP xenografts compared with non–HPN-expressing PC3 xenografts. HPN imaging may provide a new method for detection of prostate cancer. [Cancer Res 2008;68(7):2286–91]

Introduction

Prostate cancer is the most commonly diagnosed nonskin cancer in American men and the second leading cause of cancer-related death in this group (1). Local therapy (surgery or radiation) can be curative, and surgery is proved to save lives when compared with watchful waiting (2). However, the efficacy of local therapy is predicated on early detection; prostate cancer recurs in up to 40% of men after local therapy, presumably because it had spread before diagnosis (3). Moreover, imaging tools to accurately define tumor volume are lacking. Clearly, to improve current prostate cancer cure rates and decrease treatment-related morbidity by selecting appropriate patients for active surveillance, tools to accurately identify small amounts of localized disease and quantify tumor volume are needed.

The foundation of prostate cancer screening is serum prostate-specific antigen (PSA). This test is limited by the fact that PSA is produced both by prostate cancer and by the normal prostate. Thus, not surprisingly, serum PSA screening for early detection of prostate cancer is neither sensitive nor specific and leads to (a) unnecessary biopsies in men with benign prostatic hypertrophy (BPH) and high circulating PSA and (b) missed prostate cancer in men with smaller prostate glands and low circulating PSA (4–6). Novel prostate cancer biomarkers without these significant flaws are needed.

An important step in the discovery of novel biomarkers is a molecular understanding of prostate cancer development and progression. Several published, microarray analyses have revealed the presence of a cluster of genes up-regulated only in prostate cancer tumor cells (7). One gene, hepsin (HPN), was consistently up-regulated and had one of the highest differences in expression ratio between normal/BPH and prostate cancer. Specifically, analyses of nine published datasets (7–15), comprising ~500 patients revealed that HPN was collectively up-regulated in prostate cancer.

HPN (EC 3.4.21.106) is a type II transmembrane serine protease that is normally expressed during development but has absent or low levels of expression in normal prostate or BPH (7–15). Importantly, HPN is expressed both in the precursor lesion of prostate cancer, high-grade prostate intraepithelial neoplasia (HG-PIN), and hormone-refractory metastatic tumors (7). Its cell surface expression and malignant cell localization thus make it an ideal candidate for the development of prostate cancer–targeted imaging agents. Unfortunately, to date, there are no commercially available reagents with sufficiently high affinity, such as peptides or small molecules. Therefore, we used iterative phage display selection to identify novel high-affinity binding peptides selective and specific for HPN. The identified peptides were able to detect human prostate cancer on tissue microarrays. Using the developed agent, we were able to visualize 4.6-mm prostate cancer by tomographic optical imaging.

Materials and Methods

Materials. RPMI medium without phenol red, 1 mol/L HEPES solution, sodium pyruvate solution, sodium bicarbonate solution, Dulbecco’s PBS with Ca++ and Mg++, and HBSS were purchased from Biowhitaker Bioproducts. FCS was purchased from Cellgro. Lipofectin was obtained from Invitrogen. All other chemicals were of the highest quality grade available from Fisher Scientific or Sigma Chemical Co.

Cell lines. PC3, LNCaP, and DU145 human prostate cancer cells were obtained from American Type Tissue Culture Collection. The PC3 and DU145 (HPN negative; ref. 16) and LNCaP (HPN positive; ref. 16) cell lines were maintained in RPMI without phenol red medium supplemented with 10% FCS, 2 mmol/L L-glutamine, and 1 mmol/L sodium pyruvate at 37°C in 5% CO2. HPN stably transfected PC3 cells (HPN-PC3) were maintained in RPMI without phenol red supplement with 10% FCS, 2 mmol/L glutamine, 1 mmol/L sodium pyruvate, and 500 μg/mL G418 at 37°C in 5% CO2. For routine maintenance, cells were passaged by trypsinization immediately upon reaching confluence.
Transfections. The entire HPN encoding sequence fused to a FLAG tag was cloned into the vector PC-DNA3.1 (gift from Dr. Dhanasekaran and Dr. Chinnaiyan). The plasmid was grown in Escherichia coli (MAX Efficiency DH5α Competent Cells, Invitrogen) and isolated with an endotoxin-free preparation kit (Endo-Free Plasmid Maxikit, Qiagen). Transfection into PC3 cells was performed by overnight incubation of endotoxin-free preparation kit (Endo-Free Plasmid Maxikit, Qiagen) medium, the cells were transferred into selection medium (700 g/mL of 0.5 mol/L Tris-HCl (pH 9.0)). To deplete phage that bound to identical markers present on both HPNPC3 and PC3 cells, the phage pool isolated after one round of selection was subtracted by three rounds of successive incubation at 37°C for 30 min with confluent monolayers of PC3 cells that are negative for HPN. The phage that were internalized by HPNPC3 cells, but not PC3 cells, were amplified by E. coli, titered, and subjected to three additional rounds of positive selection, and individual clones were selected for ELISA assay and sequencing (Fig. 1A).

Phage display selection. Phage selection and negative depletion were performed using previously published protocols (17). Briefly, 10^10 plaque-forming units (pfu) of phage, displaying random seven amino acid peptides (7PhID; New England Biolabs), were incubated with HPNPC3 cells at 37°C for 1 h to allow time for HPN-mediated internalization. Extracellular restricted phages were removed with 0.2 mol/L glycyne (pH 2.2; 3 × 8 min). Internalized phages were recovered by lysis with 0.1% triethanolamine (Sigma Chemical) in PBS (pH 7.4; 4 min, room temperature). Extracts were neutralized with 500 μL of 0.5 mol/L tris-HCl (pH 9.0). To deplete phage that bound to identical markers present on both HPNPC3 and PC3 cells, the phage pool isolated after one round of selection was subtracted by three rounds of successive incubation at 37°C for 30 min with confluent monolayers of PC3 cells that are negative for HPN. The phage that were internalized by HPNPC3 cells, but not PC3 cells, were amplified by E. coli, titered, and subjected to three additional rounds of positive selection, and individual clones were selected for ELISA assay and sequencing (Fig. 1A).

ELISA. Confluent monolayers of HPNPC3 or PC3 cells were incubated at room temperature with individual phage clones (10^9 pfu, 1 h), washed with PBS containing 0.1% Tween 20, incubated with biotinylated anti-M13 antibody (1:40, 1 h), detected with streptavidin–horseradish peroxidase (1:500), developed with tetramethyl benzidine and absorbance at 650 was determined (Emax, Molecular Devices). Heat maps are a graphical representation of the ELISA data with green representing highest values, black representing mean values, and red representing low values. Affinity refers to the absorbance of clones on HPNPC3 cells, and specificity is the ratio of absorbance of clones on HPNPC3 and PC3 cells (Fig. 1B).

Peptide synthesis. Peptides (HPN targeted and scrambled; Table 1) were purchased from Tufts peptide core facility with a GS/GSK/FTIC/C linker for conjugation of the peptide to a model fluorescent nanoparticle, cross-linked iron oxide (CLIO), which has been previously used as a targeting platform (18). Briefly, succinimidyl iodoacetic acid was reacted with aminated iron oxide (CLIO), which has been previously used as a targeting platform (18). After coupling, the iron oxide nanoparticles were purified again using size exclusion chromatography, and the ratio of peptides/nanoparticle was quantified by absorbance spectroscopy. A typical synthesis results in 11 peptides/CLIO.

Fluorescence confocal microscopy and flow cytometry. Phage were FITC labeled as previously described (19). Confluent HPNPC3 cells or PC3 cells were incubated with 10^9 pfu of FITC labeled phage for 1 h at 37°C, washed 3× with PBS, analyzed via flow cytometry (Fig. 1C; 10,000 cells per sample) on a Becton Dickinson FACsCalibur, and then visualized by fluorescence microscopy (Fig. 1D; Nikon 80i Eclipse equipped with 1152 Photometrics Cascade CCD, Nikon). HPNPC3, LNCaP, PC3, and DU145 cells were incubated with 1 μmol/L of IPL-F for 1 h at 37°C, washed, and then analyzed via flow cytometry. Mean fluorescence from 30,000 cells was plotted to describe relative IPL-F uptake (Fig. 2A).

Competition experiments. HPNPC3 cells were incubated with IPL-F and either anti-HPN antibody or vehicle for 1 h at 37°C, washed, detached, and then analyzed via flow cytometry (Becton Dickinson FACsCalibur; Fig. 2B). Additionally, IPL-D and IPL-F or ScP were incubated with HPNPC3 cells for 1 h at 37°C and washed, and then phage were eluted and titered to determine the extent of competition (Fig. 2C).

Tissue microarray. Binding of HPN-specific peptides to ex vivo human tissues was tested using frozen tissue sections and tissue microarrays (Fig. 3 and Supplementary Fig. S1). The prostate samples used for the study

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Table 1. Description of agents used in this study

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Sequence</th>
<th>Label</th>
<th>Use</th>
</tr>
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<tbody>
<tr>
<td>IPL-D</td>
<td>Plasmid</td>
<td>FITC</td>
<td>Microscopy, competition, FACS</td>
</tr>
<tr>
<td>IPL-F</td>
<td>Peptide</td>
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</tr>
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<td>Peptide</td>
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<tr>
<td>IPL-NP</td>
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<tr>
<td>NP</td>
<td>Nanoparticle</td>
<td>AF750(NP)</td>
<td>Imaging</td>
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were part of the prostatectomy cohort obtained from the BWH DFHCC prostate cancer Specialized Programs of Research Excellence tissue bank. A multitumor frozen array was constructed using a manual tissue arrayer (Beecher Instruments) with prior institutional review board approval. The array consisted of benign prostate samples ($n = 6$) and localized prostate cancer ($n = 5$). Four of the tumor samples had Gleason grades of 6 with one sample having a Gleason grade of 8. Each sample was represented by two cores in the array for a total of 22 tissue sections. During tissue sectioning, samples were placed in dry ice to prevent melting. Dry ice was held against the needle before and after coring and while dispensing the core to minimize melting. Sections (5-μm) were used for in situ analysis of the peptides. The sections were fixed in ice-cold acetone and blocked with 5% bovine serum albumin (BSA), 30 min at room temperature followed by 10% normal goat serum, 30 min at room temperature and with serum-free protein block (DakoCytomation) for 10 min at room temperature. Following this, the sections were washed thrice with calcium magnesium–free PBS and incubated with 0.1 μmol/L IFL-F or 0.1 μmol/L IPL-NP in PBS containing 0.1% BSA for 2 h at 37°C, washed 3× with PBS, mounted with prolong gold antifade reagent containing 4',6-diamidino-2-phenylindole (DAPI; Invitrogen). The binding was visualized via fluorescence microscopy (Olympus) BX51 fluorescence microscope equipped with a charge-coupled device camera, and the CytoVision fluorescence in situ hybridization imaging and capturing software (Applied Imaging). The specificity of binding was confirmed by two pathologists (S.P. and M.A.R.).

**Specificity of IPL-NP.** HPN-PC3 and PC3 cells were incubated with 1 μmol/L of IFL-F or IPL-NP for 1 h at 37°C, washed, and then analyzed via flow cytometry. Mean fluorescence from 30,000 cells was plotted to describe relative IFL-F uptake (Fig. 4A).

**In vivo optical imaging experiments.** Male nude mice ages 7 to 9 weeks were purchased from the MGH breeding colony (C0X7) and maintained in approved pathogen-free institutional housing facilities. All experiments were performed according to institutional guidelines. In all experiments, probe was given i.v. at 20 mg Fe/kg body weight. For the in vivo specificity of IPL-NP, optical imaging of nude mice ($n = 7$) bearing s.c. implanted tumors (diameter, 5-6 mm) derived from the immortalized human prostate cancer cell lines, the HPN-negative PC3 (Fig. 4B, left flank) and HPN-positive LNCaP (Fig. 4B, right flank) was performed. We chose LNCaP cells because they express HPN, IFL-F binds to them, and HPN-PC3 cells fail to form tumors in immunocompromised mice (data not shown; ref. 20). Mice were coimplanted with tail vein with IFL-NP and HPN (see Table 1 for description) then imaged via fluorescence-mediated tomography (FMT) 24 h after injection (Fig. 4B), and the results quantitated in Fig. 4C. Image acquisition variables, analysis, and quantization were similar to what was recently described (21). For both half-life studies and tumor accumulation time-course studies, nude mice ($n = 5$) were s.c. implanted with LNCaP cells on each flank (Fig. 5A and B). The mice were imaged via FMT for IPL-NP tumoral accumulation, and blood was drawn at 0, 1, 3, 6, 24, and 48 h post-probe administration. Blood was analyzed for IPL-NP presence by fluorescence reflectance imaging, which quantitates the amount of fluorochrome in the sample (Bonsai, Siemens Medical). Background fluorescence of blood was analyzed using noninjected control animals ($n = 3$). To determine the size limit of tumor detection, mice bearing different sizes of LNCaP-derived tumors were injected with probe then imaged via FMT 3 h postinjection (Fig. 5C).

**Results**

**Identification of peptides.** To facilitate the phage display selection, we used HPN-negative PC3 cells and stably transfected them to perform iterative positive and negative selections. Two transfected clones in particular had high levels of HPN mRNA expression by real-time quantitative PCR when compared with parental PC3 cells (clone 27, 1,960-fold difference; clone 30, 967-fold difference). A total of 30 individual phage were randomly identified and selected for further analysis. Two of the 30 clones had identical sequences with each other, and three other clones shared homology to form a consensus sequence, IPLXVPL (Fig. 1A). The two identical phage clones were shown to have nearly identical binding patterns as seen by heatmaps of ELISA analysis results (Fig. 1B, closed arrows). Additionally, peptides with similarity to the consensus sequence had the same binding profile (peptides 7, 21, 25). To confirm the ELISA results, phage corresponding to the sequence IPLXVPL were labeled with FITC (IFL-Φ) incubated with either HPN-PC3 or PC3 cells and analyzed via flow cytometry (Fig. 1C) and fluorescence microscopy (Fig. 1D). Using IFL-Φ, we were able image and label 95.2% of HPN-PC3 cells 12-fold better than PC3 cells. In further validation experiments, IFL-F bound to HPN-PC3
cells and LNCaP cells, which express HPN (Fig. 2A), whereas PC3 cells and DU145 cells, which do not express HPN, had background levels of IFL-F binding (Fig. 2A). In addition, IFL-F were able to compete with IFL-F for binding to HPN-PC3 cells with an IC50 of 1.4 + 1.7 μmol/L, whereas ScP was unable to compete at the concentrations examined (Fig. 2B). Preincubation of HPN-PC3 cells with anti-HPN antibody abrogated IFL-F binding by 84.4% (Fig. 2B). Finally, synthetic IPLVPL peptide (IFL-F) had an affinity of 190 + 2.2 nmol/L (mean ± SD) for HPN-PC3 cells, whereas scrambled control peptide, ScP, exhibited negligible binding (Fig. 2C). Taken together, these experiments show specific binding of the identified peptide to the prostate cancer biomarker, HPN.

HPN-targeted peptides and nanoparticles are able to identify human prostate cancer. Because IPL-F, IPL-NP, and IFL-F were all able to detect human prostate cancer in vitro, we next determined their specificity on human tissues samples. The binding of IFL-F was first tested on frozen tissue sections. The peptide was seen to bind specifically to the tumor glands, with adjacent benign glands showing no staining (Supplementary Fig. S1). The results were further validated on a tissue microarray. Frozen tissue microarray consisting of 22 prostate samples (two cores per case, six normal prostates, and five prostate cancers) was constructed and used for in situ fluorescent histochemical analysis. IFL-F and IPL-NP were able to bind selectively to prostate cancer cells (Fig. 3A and C), staining 100% of the tumor cores and 0% of the control benign cores (Fig. 3B and D). Staining was localized to the prostate tumor cores with the stromal cells showing basal staining. Control NP did not show any binding (data not shown).

In vivo imaging of HPN-expressing tumors. Conjugation of IFL-F to a model nanoparticle (IPL-NP) increased the fluorescent signal via FACs by >10-fold when compared with peptide alone, presumably due to multivalency effects (Fig. 4A). IPL-NP retained its specificity for HPN-expressing cells, as binding to PC3 cells was 80-fold less than binding to HPN-PC3 cells (Fig. 4A). To determine whether we could detect HPN-expressing tumors in vivo, we used a xenograft model of prostate cancer with a PC3 cell–derived tumor on the left flank and LNCaP (HPN positive) cell–derived tumor on the right flank (Fig. 4B). After i.v. coinjection of IPL-NP and NP (untargeted control nanoparticle), FMT images were obtained (Fig. 4B). HPN-positive tumors accumulated 65 + 3.1 nmol/L IPL-NP, whereas HPN-negative tumors showed lower amounts (23 + 1.6 nmol/L; P = 0.0675, Student’s t test). The differences in tumoral accumulation were not due to differences in bulk tumor properties, such as angiogenesis or tumor size, since untargeted NP had a statistically identical accumulation between the two tumors (Fig. 4C). Moreover, the peptide sequence imparted in vivo HPN targeting to the nanoparticle because LNCap tumors had 21 + 8.1 nmol/L of NP, which is indistinguishable from NP accumulation in PC3 derived tumors. As part of the in vivo characterization of IPL-NP, we determined the blood half-life of IPL-NP to be 7.4 hours (R2, 0.99; Fig. 5A). The time course of tumoral accumulation was assessed showing maximal signal at 24 hours with signal persisting past 48 hours (Fig. 5B). Finally, an important clinical question especially for early detection of tumors is determining the smallest size of tumors that can be detected. Using FMT imaging, we were able to detect 4.6-mm diameter prostate cancer (Fig. 5C). The correlation of tumoral accumulation of IPL-NP with tumor volume was linear with an R2 value of 0.89.

Discussion

Improving prostate cancer detection will ultimately lead to improving prostate cancer cure rates. Moreover, accurately quantifying prostate cancer burden could dramatically improve the ability to select appropriate candidates for active surveillance. Currently, prostate cancer detection relies on serum PSA and the digital rectal exam (22). Due to the fact that the normal prostate produces PSA and the high rates of BPH in the patient population

Figure 3. HPN-targeted peptides and nanoparticles are able to identify human prostate cancer. IFL-F or IPL-NP was incubated with tissue microarrays consisting of six normal and five cancer human radical prostatectomy specimens. Results shown are representative images. A and C, binding of IFL-F/IPL-NP to cancer glands. B and D, binding to benign glands. Original magnification of A and B, 20×; original magnification of C and D, 60×. Note staining in the tumor cells and the absence/weak staining in benign tissues and stroma.
at risk for prostate cancer, serum PSA is neither a sensitive nor specific biomarker (23). In fact, it has been argued that in the era of PSA screening, serum PSA is much more highly correlated with prostate volume than with the presence of prostate cancer (24). Whereas several variations of serum PSA may improve its characteristics slightly (free PSA, PSA density), overall it is a relatively poor screening test (25).

The goal of this project was to develop targeted imaging agents against an emerging, potential new prostate cancer biomarker to improve prostate cancer detection. We choose HPN as a candidate target because it is expressed at high levels by prostate cancer cells (and HG-PIN) and not at all or at low levels in the nonmalignant prostate (7–15). The fact that HPN is a cell surface protein and is absent or at low levels in BPH made it a particularly attractive candidate for further study. Additionally, HPN expression in other cancers, including ovarian, breast, and renal, has been documented (26–28). Using an iterative phage display selection approach, we identified a novel peptide (IPL) with both high affinity and high selectivity for HPN. This peptide, when tested on human tissue sections, was seen to bind specifically to the tumor glands (characterized by small glands and large nuclei), with the adjacent benign glands (large glands with smaller nuclei) showing little or no staining. The stromal compartment showed minimal staining.

In our view, the ideal prostate cancer screening modality incorporates imaging, because imaging is noninvasive, repeatable, and provides both binary information (cancer present versus cancer absent) and anatomic information useful in making treatment decisions. Therefore, we have developed a noninvasive approach for imaging HPN expression and prostate cancer. Incorporating a COOH terminal extension onto IPL allowed fluorescein attachment

Figure 4. In vivo imaging of prostate cancer. A, HPN-PC3 (black and green histograms) or PC3 (red histogram) cells were incubated with IPL-F (black histogram) or IPL-NP (green histogram) and then analyzed via flow cytometry. B and C, mice bearing tumors derived from PC-3 (left flank) or LNCaP (right flank) were coinjected with IPL-NP (green columns) and NP (red columns) and then imaged (B), and accumulation was quantified via FMT 24 h postinjection (C).

Figure 5. In vivo characterization of IPL-NP. A, mice bearing tumors derived from LNCaP cells were injected with IPL-NP (20 mg/kg Fe), and then blood was analyzed for agent presence at indicated time points postinjection. B, time course of IPL-NP accumulation into LNCaP-derived tumors. Mice bearing LNCaP-derived tumors were injected with IPL-NP (20 mg/kg Fe) and then imaged via FMT at 0, 1, 3, 6, 24, and 48 h postinjection. C, mice bearing tumors of different volumes derived from LNCaP cells were injected with identical doses of IPL-NP (20 mg/kg Fe) and then imaged via FMT.
and nanoparticle conjugation, ideal for increasing the circulation time of the agent. As expected, nanoparticle conjugation (at 11 peptides to one nanoparticle) improved avidity, as shown by a >10-fold increase in fluorescent signal compared with peptide alone. Importantly, the fluorescent IPL-NP was able to distinguish between an HPN-expressing LNCaP xenograft and a non–HPN-expressing PC-3 xenograft in vivo. The agent seemed to be selective for human prostate cancer samples tested and was absent in the normal prostate sections.

In conclusion, we have identified and developed a novel imaging agent (IPL-NP), which binds HPN with high affinity. We have shown, with both an in vivo model and with in situ histochemical analysis on patient tissues, its potential as an imaging agent for prostate cancer. Our next step is to investigate the suitability of IPL-based peptides for human use. We envision using similar nanoparticle platforms as those used in clinical trials, as the nanoparticles used in this study have similar properties to those used in the clinic (29).

Such an HPN-targeted imaging agent would be useful in all stages of prostate cancer, from biopsy targeting (prediagnosis) to the delineation of nonlocalized disease to identification of disease recurrence posttherapy, and would make a significant effect on the diagnosis and treatment of this lethal disease.

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References

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