ANTIBODIES TO PROSTATE-SPECIFIC MEMBRANE ANTIGEN

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Assignee: Sloan-Kettering Institute for Cancer Research, New York, NY (US)

Notice: This patent is subject to a terminal disclaimer.

Appl. No.: 12/283,636 Filed: Sep. 12, 2008

United States

Reissued Patent

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31 Claims, 48 Drawing Sheets

ABSTRACT

This invention provides purified antibodies to the outer membrane domain of prostate-specific membrane (PSM) antigen, compositions of matter comprising PSM antigen antibodies conjugated to a radioisotope or a toxin, and a method of imaging prostate cancer by using PSM antigen antibodies.


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Response to Aug. 17, 2007 Communication Pursuant to Rule 57(1) filed Apr. 28, 2008 in connection with European Patent No. 668777. Communication of a Notice of Opposition mailed Jul. 18, 2007 in connection with related European Application No. 94900538.3, now European Patent No. 668777 granted Oct. 11, 2006, as well as (A) opposition documents filed by BZL Biologies ILC including (1) a Notice of Opposition Form 2300 filed by BZL Biologies ILC.; and (2) Statement of Grounds for Opposition and copies of references cited in the Grounds for Opposition (i) EP 0668777; (ii) WO94/ 095820; (iii) Holmes et al. 2001, Expert Opinion on Investigational Drugs, 10(3):544-519; and (iv) Schulke et al., 2003, PNAS, 100(22):12590-12595 and (B) opposition documents filed by Northwest Bio Therapeutics Inc. including a Notice of Opposition Form 2300 from Northwest Bio Therapeutics Inc.; and (2) Statement of Grounds for Opposition including a list of documents D1 through D36 (D1 through D25, D27, D28, D30-D32 and D36 of which are of record in the subject application. 


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Declaration of Dr. Mohan Srinivasan filed Apr. 24, 2008 in connection with Opposition to European Patent No. 1210374.

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Written Submission of PSMA Development Company LLC filed Nov. 29, 2008 in connection with Opposition to European Patent No. 0956506.

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Table Summarizing Potential Therapeutic Indications and Development Stages for Medarex Strategic Assets.


U.S. Appl. No. 08/466,381, filed Jun. 6, 1995, Israeli et al.

* cited by examiner
FIGURE 1

1 - anti- EGFr PoAB RK-2
2 - Cyt-358 MoAB/RAM
3 - RAM

M.K. 1 2 3

200KD -
100.5KD -
72KD -
43KD -
28.5KD -
FIGURE 14-1

Done on sequence PMSANTIGEN.
Total number of residues is: 750.
Analysis done on the complete sequence.

In Helical (H) conformation [DC = -75 CNAT ] : 264 AA => 35.2%
In Extended (E) conformation [DC = -88 CNAT ] : 309 AA => 41.2%
In Turn (T) conformation [DC = 0 CNAT ] :  76 AA => 10.1%
In Coil (C) conformation [DC = 0 CNAT ] : 101 AA => 13.4%

-----------------------------------
Sequence shown with conformation codes.
-----------------------------------
Consecutive stretch of 5 or more residues in a given conformation are overlined.

```
  1  H H H H H H H H H H H H E E E T T E E E E E E E E E 
  31 E E E E E E E H H H H H C C C C C C T H H H H H H H H H 
  61 H H H H H H H H H H H H E E E T T E E E E C C C C C C H H H 
```
FIGURE 14-4

Semi-graphical output.

Symbols used in the semi-graphical representation:

- Helical conformation: X
- Turn conformation: >
- Coiled conformation: *

MWLHTEGAVATARRPPWRQGALAGYLVFLGFGFLPFGWFTKSSNEAT

XXXXXXX---<>---XXX

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<td>F</td>
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<td>V</td>
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<td>KVP</td>
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<td>V</td>
<td>P</td>
<td>V</td>
<td>G</td>
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<td>F</td>
<td>G</td>
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<td>H</td>
<td>T</td>
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<td>L</td>
<td>G</td>
<td>P</td>
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</tr>
<tr>
<td>4</td>
<td>G</td>
<td>I</td>
<td>G</td>
<td>G</td>
<td>W</td>
</tr>
<tr>
<td>5</td>
<td>A</td>
<td>N</td>
<td>A</td>
<td>E</td>
<td>G</td>
</tr>
</tbody>
</table>
FIGURE 15B

******************************************************************************
* PREDICTION OF ANTIGENIC DETERMINANTS *
******************************************************************************

Done on sequence PMSANTIGEN.
Total number of residues is: 750.
Analysis done on the complete sequence.

The method used is that of Hopp and Woods.
The averaging group length is: 6 amino acids.
--> This is the value recommended by the authors <--

----------------------------------------

The three highest points of hydrophilicity are:

( 1) Ah= 1.62 : From 63 to 68 : Asp-Glu-Leu-Lys-Ala-Glu
( 2) Ah= 1.57 : From 132 to 137 : Asn-Glu-Asp-Gly-Asn-Glu
( 3) Ah= 1.55 : From 482 to 487 : Lys-Ser-Pro-Asp-Glu-Gly

Ah stands for: Average hydrophilicity.

Note that, on a group of control proteins, only the highest point was in 100% of the cases assigned to a known antigenic group. The second and third point gave a proportion of 33% of incorrect predictions.
The best scores are:

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<th>Gene</th>
<th>Sequence Description</th>
<th>Initn</th>
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<th>Opt</th>
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<td>CHKTFER</td>
<td>G. gallus mRNA for transferrin receptor</td>
<td>203</td>
<td>120</td>
<td>321</td>
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<tr>
<td>RATTRFR</td>
<td>Rat transferrin receptor mRNA, 3' end.</td>
<td>164</td>
<td>164</td>
<td>311</td>
</tr>
<tr>
<td>HUMTFRR</td>
<td>Human transferrin receptor mRNA, complete cd</td>
<td>145</td>
<td>145</td>
<td>266</td>
</tr>
</tbody>
</table>

CHKTFER  G. gallus mRNA for transferrin receptor  203  120  321
51.9% identity in 717 nt overlap

```
1020 1030 1040 1050 1060 1070
pmsgen TGTTCAAGCGTGAAATATACTCCTAATCTGAATGTGCAGGAGACCCTCTCACACCAGTTA
          ::::  :::::::::  :::::::::  :
CHKTFE   TACACTTTATCCATCCATTGGACATGCCCCACCTTGGAAACTGGAGACCCTTTACACCACCCAGGCTT
         990 1000 1010 1020 1030 1040

1080 1090 1100 1110 1120 1130
pmsgen CCGAGCAAATGAATATGCTTATAGGGCTGGAATTGCAGGCTTGTGGCTTCCAAATAT
          ::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  ::::
CHKTFE   CCCCTTCGTTCAACCAACCCCA---GTTTCCACCCAGTTGAATCTTCCAGGACTACCCCACAT
         1050 1060 1070 1080 1090 1100

1140 1150 1160 1170 1180 1190
pmsgen TTCTGTTCTGGTAATCTATCTATGCAATGCAAGAGCCTTGAGAAATAATGGTTGGCTC
          ::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  :::::::::  ::::
CHKTFE   TGCTGTTGCAACCACATTCTAGCATGCAGCAGCAGGCTTCTCAGGCAAATGGATGGAGA
         1110 1120 1130 1140 1150 1160
```
FIGURE 16-5

RATTTRFR  Rat transferrin receptor mRNA, 3' end.  164  164  311
55.5% identity in 560 nt overlap

1210  1220  1230  1240  1250
pmsgen  CCACCAGATAGCAGCTGGAGAGGAGTCTCAAAAGTGCCCTACAATGTGGGACCTGGCTT-

1260  1270  1280  1290  1300  1310
pmsgen  -TACTGGAAACTTTTCACACAAAGTGAAGATGCACATC-CACTCT-ACCAATG---

1210  1220  1230  1240  1250  1260  1270  1280  1290  1300  1310
RATTTRFR  TGCAGAAAAGCTATTCAAAAAACATGGAAAGGAACTTGCTCTCTCTATGTGGAAATATAGATTC
610  620  630  640  650  660

1210  1220  1230  1240  1250  1260  1270  1280  1290  1300  1310
pmsgen  -TACTGGAAACTTTTCACACAAAGTGAAGATGCACATC-CACTCT-ACCAATG---

1210  1220  1230  1240  1250  1260  1270  1280  1290  1300  1310
RATTTRFR  CTCATGTAAGCTGGAACTTTTCACAGAAATCTAAAATGTGGAAGCTACTGTGAACAATGTACT
670  680  690  700  710  720
FIGURE 16-8

GCTTTGAAGGCAAAATCTCTTTAT-GAA------AGTTGGAATTTAAAAAGTCCTTCCAG

---TTGATGGAAATATCTATATCGAAACAGTAATTGGATTAGCAAAATGGAGGAACTTT

AGTTCAGTGGCATGCGCAGGATAAGCAAAATTGGGATCTGGAAATGATTTTGGAGGTGTTCT

CTTTGGCAATGCTGATTCCTTTCTTCTGATTTACAGAATCCGACCAGTTTCTCTTCTTCT

1730  1740  1750  1760  1770
1140  1150  1160  1170  1180  1190
1780  1790  1800  1810  1820  1830
1200  1210  1220  1230  1240  1250
FIGURE 16-9

HUMTFRR  Human transferrin receptor mRNA, complete cd 145 145 266
54.3% identity in 464 nt overlap

1230 1240 1250 1260 1270
pmsgen AGGAAGTCTCTCAAAGTGCCCTACAATGTGGACCTGGCTTTTAC-TGGAAACTTTTCTACAC

1140 1150 1160 1170 1180 1190
HUMTFRR TATGGAAGGAGACTGTCCCTCTGACTGGAAGAACAGACTCTACATGTAAGGATGGTAAACCTC

1280 1290 1300 1310 1320 1330
pmsgen AAAAAAGTCAAGATGCACATC-CACTCT-ACCAATG-------AAGTGACAAGAATTTACAA

1200 1210 1220 1230 1240 1250
HUMTFRR AGAAAAGCAAGAATGTGAAGCTCACTGTGAACATGTGCTGAAAGAGATAAAAATCTTAA

1340 1350 1360 1370 1380 1390
pmsgen TGTGATAGGTACTCTCAAGAGGACGTGGAAACAGAACAGATATGTCATTCTGGAGGCTCA

1260 1270 1280 1290 1300 1310
HUMTFRR CATCTTTGGAGTTATTTAAGGCTTTGTAAGAAACCAGATCACTATGTTGAGTTGGGCCC

1400 1410 1420 1430 1440 1450
pmsgen CCGGCACTCATGCTTTTGTTGGTATGTACCCCTACAGAGT-GAGACAGCTGTTGTTCATG

1320 1330 1340 1350 1360 1370
HUMTFRR GAGAGATGCATGGGCCCCTGGAGCTGAAATC-CGGTGATAAGCAGACCTCTCCTATTTGA
FIGURE 22

298 bp

260 bp
### FIGURE 23

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FIGURE 26
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1 ANTIBODIES TO PROSTATE-SPECIFIC MEMBRANE ANTIGEN

Matter enclosed in heavy brackets [ ] appears in the original patent but forms no part of this reissue specification; matter printed in italics indicates the additions made by reissue.

This application is a continuation application of U.S. Ser. No. 08/403,803, filed (Mar. 17, 1995, now abandoned) and a continuation of PCT International Application No. PCT/US93/10624, filed Nov. 5, 1993, which is a continuation-in-part of U.S. Ser. No. 07/973,337, filed Nov. 5, 1992, now abandoned the contents of which are hereby incorporated by reference.

This invention disclosed herein was made in part with Government support under NIH Grants No. DK47650 and CA58192 from the Department of Health and Human Services. Accordingly, the U.S. Government has certain rights in this invention.

BACKGROUND OF THE INVENTION

Throughout this application various references are referred to within parentheses. Disclosures of these publications in their entirety are hereby incorporated by reference into this application to more fully describe the state of the art to which this invention pertains. Full bibliographic citation for these references may be found at the end of each series of experiments.

Prostate cancer is among the most significant medical problems in the United States, as the disease is now the most common malignancy diagnosed in American males. In 1992 there were over 132,000 new cases of prostate cancer detected with over 36,000 deaths attributable to the disease, representing a 17.3% increase over 4 years (2). Five year survival rates for patients with prostate cancer range from 88% for those with localized disease to 29% for those with metastatic disease. Therefore, the number of cases appears to result from an increase in disease awareness as well as the widespread use of clinical markers such as the secreted proteins prostate-specific antigen (PSA) and prostate acid phosphatase (PAP) (37).

The prostate gland is a site of significant pathology affected by conditions such as benign growth (BPH), neoplasms (prostatic cancer) and infection (prostatitis). Prostate cancer represents the second leading cause of death from cancer in men (1). However, prostatic cancer is the leading site for cancer development in men. The difference between these two facts relates to prostatic cancer occurring with increasing frequency as men age, especially in the ages beyond 60 at a time when death from other factors often intervenes. Also, the spectrum of biologic aggressiveness of prostatic cancer is great, so that in some men following detection the tumor remains a latent histologic tumor and does not become clinically significant, whereas in other it progresses rapidly, metastasizes and kills the man in a relatively short 2-5 year period (1, 3).

In prostate cancer cells, two specific proteins that are made in very high concentrations are prostatic acid phosphatase (PAP) and prostate specific antigen (PSA) (4, 5, 6). These proteins have been characterized and have been used to follow response to therapy. With the development of cancer, the normal architecture of the gland becomes altered, including loss of the normal duct structure for the removal of secretions and thus the secretions reach the serum. Indeed measurement of serum PSA is suggested as a potential screening method for prostatic cancer. Indeed, the relative amount of PSA and/or PAP in the cancer reduces as compared to normal or benign tissue.

PAP was one of the earliest serum markers for detecting metastatic spread (4). PAP hydrolyzes tyrosine phosphate and has a broad substrate specificity. Tyrosine phosphorylation is often increased with oncogenic transformation. It has been hypothesized that during neoplastic transformation there is less phosphatase activity available to inactivate proteins that are activated by phosphorylation on tyrosine residues. In some instances, insertion of phosphatases that have tyrosine phosphatase activity has reversed the malignant phenotype.

PSA is a protease and it is not readily appreciated how loss of its activity correlates with cancer development (5, 6). The proteolytic activity of PSA is inhibited by zinc. Zinc concentrations are high in the normal prostate and reduced in prostatic cancer. Possibly the loss of zinc allows for increased proteolytic activity by PSA. As proteases are involved in metastasis and some proteases stimulate mitotic activity, the potentially increased activity of PSA could be hypothesized to play a role in the tumors metastases and spread (7).

Both PSA and PAP are found in prostatic secretions. Both appear to be dependent on the presence of androgens for their production and are substantially reduced following androgen deprivation.

Prostate-specific membrane antigen (PSM) which appears to be localized to the prostatic membrane has been identified. This antigen was identified as the result of generating monoclonal antibodies to a prostatic cancer cell, LNCaP (8).

Dr. Horoszewicz established a cell line designated LNCaP from the lymph node of a hormone refractory, heavily pre-treated patient (9). This line was found to have an aneuploid human male karyotype. It maintained prostatic differentiation functionality in that it produced both PSA and PAP. It possessed an androgen receptor of high affinity and specificity. Mice were immunized with LNCaP cells and hybridomas were derived from sensitized animals. A monoclonal antibody was derived and was designated 7E11-C5 (8). The antibody staining was consistent with a membrane location and isolated fractions of LNCaP cell membranes exhibited a strongly positive reaction with immunoblotting and ELISA techniques. This antibody did not inhibit or enhance the growth of LNCaP cells in vitro or in vivo. The antibody to this antigen was remarkably specific to prostatic epithelial cells, as no reactivity was observed in any other component. Immunohistochemical staining of cancerous epithelial cells was more intense than that of normal or benign epithelial cells.

Dr. Horoszewicz also reported detection of immunoreactive material using 7E11-C5 in serum of prostatic cancer patients (8). The immunoreactivity was detectable in nearly 60% of patients with stage D-2 disease and in a slightly lower percentage of patients with earlier stage disease, but the numbers of patients in the latter group are small. Patients with benign prostatic hyperplasia (BPH) were negative. Patients with no apparent disease were negative, but 50-60% of patients in remission yet with active stable disease or with progression demonstrated positive serum reactivity. Patients with non prostatic tumors did not show immunoreactivity with 7E11-C5.

The 7E11-C5 monoclonal antibody is currently in clinical trials. The aldehyde groups of the antibody were oxidized and the linker-chelator glycol-tyrosyl- (n, e-diethylaminoethylurea-peracetic acid)-lysine (GTY-DTPA) was coupled to the reactive aldehydes of the heavy chain (10). The resulting antibody was designated CYT-356. Immunohistochemical
staining patterns were similar except that the CYT-356 modified antibody stained skeletal muscle. The comparison of CYT-356 with 7E11-CS monoclonal antibody suggested both had binding to type 2 muscle fibers. The reason for the discrepancy with the earlier study, which reported skeletal muscle to be negative, was suggested to be due to differences in tissue fixation techniques. Still, the most intense and definite reaction was observed with prostatic epithelial cells, especially cancerous cells. Reactivity with mouse skeletal muscle was detected with immunohistochemistry but not in imaging studies. The Indium-111-labeled antibody localized to LNCaP tumors grown in nude mice with an uptake of nearly 30% of the injected dose per gram tumor at four days. In vivo, no selective retention of the antibody was observed in antigen negative tumors such as PC-3 and DU-145, or by skeletal muscle.

Very little was known about the PSM antigen. An effort at purification and characterization has been described at meetings by Dr. George Wright and colleagues (11, 12). These investigators have shown that following electrophoresis on acrylamide gels and Western blotting, the PSM antigen maintains a molecular weight of 100 kilodaltons (kd). Chemical and enzymatic treatment showed that both the peptide and carbohydrate moieties of the PSM antigen are required for recognition by the 7E11-C5 monoclonal antibody. Competitive binding studies with specific lectins suggested that gal-NAc is the dominant carbohydrate of the antigenic epitope.

The 100 kd glycoprotein unique to prostate cells and tissues was purified and characterized. The protein was digested proteolytically with trypsin and nine peptide fragments were sequenced. Using the technique of degenerate PCR (polymerase chain reaction), the full-length 2.65 kilobase (kb) cDNA coding for this antigen was cloned. Preliminary results have revealed that this antigen is highly expressed in prostate cancer tissues, including bone and lymph node metastases (13). The entire DNA sequence for the cDNA as well as the predicted amino acid sequence for the antigen was determined. Further characterization of the PSM antigen is presently underway in the applicants' laboratory including: analysis of PSM gene expression in a wide variety of tissues, transfected or of the PSM gene into cells not expressing the antigen, chromosome localization of the PSM gene, cloning of the genomic PSM gene with analysis of the PSM promoter and generation of polyclonal and monoclonal antibodies against highly antigenic peptide domains of the PSM antigen, and identification of any endogenous PSM binding molecules (ligands).

Currently, LNCaP cells provide the best in vitro model system to study human prostate cancer, since they produce all three prostate bio-markers: PSA, PAP and PSM. The cells possess an aneuploid male karyotype with a Y chromosome, express a high affinity androgen receptor, and are hormonally responsive to both testosterone and DHT. Because PSM appears to be a transmembrane glycoprotein, it is considered an attractive target for both antibody-directed imaging and targeting of prostatic tumor deposits (38). We have demonstrated expression of PSM protein in LNCAP cell membranes and in PC-3 cells transfected with PSM cDNA and also the characterization of PSM mRNA expression in human tissues, and in response to steroid hormones.

BRIEF DESCRIPTION OF FIGURES

FIG. 1: Signal in lane 2 represent the 100 kDa PSM antigen. The EGFR was used as the positive control and is shown in lane 1. Incubation with rabbit antimouse (RAM) antibody alone served as negative control and is shown in lane 3.

FIG. 2 A-D: Upper two photos show LNCaP cytospins staining positively for PSM antigen. Lower left in DU-145 and lower right is PC-3 cytospin, both negative for PSM antigen expression.

FIG. 3 A-D: Upper two panels are human prostate sections (BPH) staining positively for PSM antigen. The lower two panels show invasive prostatic carcinoma human sections staining positively for expression of the PSM antigen.

FIG. 4: 100 kD PSM antigen following immunoprecipitation of 35S-Methionine labelled LNCaP cells with Cryt-356 antibody.

FIG. 5: 3% agarose gels stained with Ethidium bromide revealing PCR products obtained using the degenerate PSM antigen primers. The arrow points to sample IN-20, which is a 1.1 kb PCR product which we later confirmed to be a partial cDNA coding for the PSM gene.

FIG. 6 A-B: 2% agarose gels of plasmid DNA resulting from TA cloning of PCR products. Inserts are excised from the PCR II vector (Invitrogen Corp.) by digestion with EcoRI. 1.1 kb PSM gene partial cDNA product is shown in lane 3 of gel 1.

FIG. 7: Autoradiogram showing size of cDNA represented in applicants' LNCaP library using M-MLV reverse transcriptase.

FIG. 8: Restriction analysis of full-length clones of PSM gene obtained after screening cDNA library. Samples have been cut with Not I and Sal I restriction enzymes to liberate the insert.

FIG. 9: Plasmid Southern autoradiogram of full length PSM gene clones. Size is approximately 2.7 kb.

FIG. 10: Northern blot revealing PSM expression limited to LNCaP prostate cancer line and H26 Ras-transfected LNCaP cell line. PC-3, DU-145, T-24, SKRC-27, HEILA, MCF-7, HL-60, and others were all negative.

FIG. 11: Autoradiogram of Northern analysis revealing expression of 2.8 kb PSM message unique to the LNCaP cell line (lane 1), and absent from the DU-145 (lane 2) and PC-3 cell lines (lane 3). RNA size ladder is shown on the left (kb), and 28S and 18S ribosomal RNA bands are indicated on the right.

FIG. 12 A-B: Results of PCR of human prostate tissues using PSM gene primers. Lanes are numbered from left to right. Lane 1, LNCaP; Lane 2, H26; Lane 3, DU-145; Lane 4, Normal Prostate; Lane 5, BPH; Lane 6, Prostate Cancer; Lane 7, BPH; Lane 8, Normal; Lane 9, BPH; Lane 10, BPH; Lane 11, BPH; Lane 12, Normal; Lane 13, Normal; Lane 14, Cancer; Lane 15, Cancer; Lane 16, Cancer, Lane 17, Normal; Lane 18, Cancer; Lane 19, IN-20 Control; Lane 20, PSM cDNA

FIG. 13: Isoelectric point of PSM antigen (non-glycosylated)

FIG. 14: 1-8 Secondary structure of antigen (panels 14-4 to 14-H; SEQ ID NO:2)


FIG. 16: 1-11 Homology of PSM antigen (SEQ ID NO:1) with chicken (SEQ ID NO:27), rat (SEQ ID NO:28) and human (SEQ ID NO:29) transferrin receptor sequence.

FIG. 17A-C: Immunohistochemical detection of PSM antigen expression in prostate cell lines. Top panel reveals uniformly high level of expression in LNCaP cells; middle panel and lower panel are DU-145 and PC-3 cells respectively, both negative.

FIG. 18: Autoradiogram of protein gel revealing products of PSM coupled in vitro transcription/translation. Non-glycosylated PSM polypeptide is seen at 84 kDa (lane 1) and...
PSM glycoprotein synthesized following the addition of microsomes is seen at 100 kDa (lane 2).

FIG. 19: Western Blot analysis detecting PSM expression in transfected non-PSM expressing PC-3 cells. 100 kDa PSM glycoprotein species is clearly seen in LNCaP membranes (lane 1), LNCaP crude lysate (lane 2), and PSM-transfected PC-3 cells (lane 4), but is undetectable in native PC-3 cells (lane 3).

FIG. 20: Autoradiogram of ribonuclease protection gel assay for PSM mRNA expression in normal human tissues. Radiolabeled 1 kb DNA ladder (Gibco-BRL) is shown in lane 1. Undigested probe is 400 nucleotides (lane 2), predicted protected PSM band is 350 nucleotides, and tRNA control is shown (lane 3). A strong signal is seen in human prostate (lane 11), with very faint, but detectable signals seen in human brain (lane 4) and human salivary gland (lane 12).

FIG. 21: Autoradiogram of ribonuclease protection gel assay for PSM mRNA expression in LNCaP tumors grown in nude mice, and in human prostatic tissues. 32P-labeled 1 kb DNA ladder is shown in lane 1. 298 nucleotide undigested probe is shown (lane 2), and tRNA control is shown (lane 3). PSM mRNA expression is clearly detectable in LNCaP cells (lane 4), orthotopically grown LNCaP tumors in nude mice with and without matrigel (lanes 5 and 6), and subcutaneously implanted and grown LNCaP tumors in nude mice (lane 7). PSM mRNA expression is also seen in normal human prostate (lane 8), and in a moderately differentiated human prostatic adenocarcinoma (lane 10). Very faint expression is seen in a sample of human prostate tissue with benign hyperplasia (lane 9).

FIG. 22: Ribonuclease protection assay for PSM expression in LNCaP cells treated with physiologic doses of various steroids for 24 hours. 32P-labeled DNA ladder is shown in lane 1. 298 nucleotide undigested probe is shown (lane 2), and tRNA control is shown (lane 3). PSM mRNA expression is highest in untreated LNCaP cells in charcoal-stripped media (lane 4). Applicant see significantly diminished PSM expression in LNCaP cells treated with DHT (lane 5) Testosterone (lane 6), Estradiol (lane 7), and Progestrone (lane 8), with little response to Dexamethasone (lane 9).

FIG. 23: Data illustrating results of PSM DNA and RNA presence in transfected Dunning cell lines employing Southern and Northern blotting techniques.

FIG. 24-A-B Figure A indicates the power of cytokine transfect. Cells to teach unmodified cells. Administration was directed to the parental flank or prostate cells. The results indicate the microenvironment considerations. Figure B indicates actual potency at a particular site. The tumor was implanted in prostate cells and treated with immune cells at two different sites.

FIG. 25-A-B Relates potency of cytokines in inhibiting growth of primary tumors. Animals administered un-modified parental tumor cells and administered as a vaccine transfectected cells. Following prostatectomy of rodent tumor results in survival increase.

FIG. 26: PCR amplification with nested primers improved our level of detection of prostatic cells from approximately one prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using either PSHA.

FIG. 27: PCR amplification with nested primers improved our level of detection of prostatic cells from approximately one prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using PSHA-derived primers.

FIG. 28: A representative ethidium stained gel photograph for PSM-PCR. Samples run in lane A represent PCR products generated from the outer primers and samples in lanes labeled B are products of inner primer pairs.

FIG. 29: PSM Southern blot autoradiograph. The sensitivity of the Southern blot analysis exceeded that of ethidium staining, as can be seen in several samples where the outer product is not visible on FIG. 3 A-D, but is detectable by Southern blotting as shown in FIG. 4.

FIG. 30: Characteristics of the 16 patients analyzed with respect to their clinical stage, treatment, serum PSA and PAP values, and results of assay.

SUMMARY OF THE INVENTION

This invention provides an isolated mammalian nucleic acid molecule encoding a mammalian prostate-specific membrane (PSM) antigen. The isolated mammalian nucleic acid may be DNA, cDNA or RNA.

This invention also provides nucleic acid molecule comprising a nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence included within the sequence of a nucleic acid molecule encoding the PSM antigen. The nucleic acid molecule may either be DNA or RNA.

This invention provides nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule which is complementary to the nucleic acid molecule encoding a mammalian prostate-specific membrane antigen.

This invention further provides a method of detecting expression of the PSM antigen which comprises obtaining total mRNA from the cell and contacting the mRNA so obtained with a labeled PSM antigen specific nucleic acid molecule under hybridizing conditions, determining the presence of mRNA hybridized to the probe, and thereby detecting the expression of the PSM antigen by the cell. The PSM antigen in tissue sections may be similarly detected.

This invention provides isolated nucleic acid molecule of PSM antigen operatively linked to a promoter of RNA transcription. This invention further provides a vector which comprises an isolated mammalian nucleic acid molecule of PSM antigen.

This invention further provides a host vector system for the production of a polypeptide having the biological activity of a mammalian PSM antigen which comprises the vector comprising the mammalian nucleic acid molecule encoding a mammalian PSM antigen and a suitable host. The suitable host for the expression of PSM antigen may be a bacterial cell, insect cell, or mammalian cell.

This invention also provides a method of producing a polypeptide having the biological activity of a mammalian PSM antigen which comprises growing the host cell of vector system having a vector comprising the isolated mammalian nucleic acid molecule encoding a mammalian PSM antigen and a suitable host under suitable conditions permitting production of the polypeptide and recovery of the polypeptide so produced.

This invention provides a method for determining whether a ligand can bind to a mammalian PSM antigen which comprises contacting a mammalian cell having an isolated mammalian DNA molecule encoding a mammalian PSM antigen with the ligand under conditions permitting binding of ligands to the mammalian PSM antigen, and determining whether the ligand binds to a mammalian PSM antigen. This invention further provides ligands which bind to PSM antigen.

This invention provides purified mammalian PSM antigen. This invention also provides a polypeptide encoded by the isolated mammalian nucleic acid molecule encoding a mamm-
malian PSM antigen. This invention further provides a method to identify and purify ligands of mammalian PSM antigen.

This invention further provides a method to produce both polyclonal and monoclonal antibody using purified PSM antigens or polypeptides encoded by an isolated mammalian nucleic acid molecule encoding a mammalian PSM antigen.

This invention provides polyclonal and monoclonal antibody most likely but not limited to directed either to peptide Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. 35), or Asp-Glu-Gly Asp-Glu-Asn-Glu (SEQ ID No. 36) or Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. 37) of the PSM antigen.

This invention provides a therapeutic agent comprising an antibody directed against a mammalian PSM antigen and a cytotoxic agent conjugated thereto.

This invention also provides a method of imaging prostate cancer in human patients which comprises administering to the patient at least one antibody directed against PSM antigen, capable of binding to the cell surface of the prostate cancer cell and labeled with an imaging agent under conditions so as to form a complex between the monoclonal antibody and the cell surface PSM antigen. This invention further provides a composition comprising an effective imaging amount of the antibody directed against PSM antigen and a pharmaceutically acceptable carrier.

This invention further provides a method of imaging prostate cancer in human patients which comprises administering to the patient multiple antibodies directed towards different PSM epitopes.

The invention also provides a method of imaging prostate cancer in human patients which comprises administering to the patient at least one ligand, capable of binding to the cell surface of the prostate cancer cell and labelled with an imaging agent under conditions so as to form a complex between the ligand and the cell surface PSM antigen. This invention further provides a composition comprising an effective imaging amount of PSM antigen and a pharmaceutically acceptable carrier.

This invention provides an immunosassay for measuring the amount of the PSM antigen in a biological sample, e.g. serum, comprising steps of a) contacting the biological sample with at least one PSM antibody to form a complex with said antibody and the PSM antigen, and b) measuring the amount of PSM antigen in said biological sample by measuring the amount of said complex.

This invention also provides an immunosassay for measuring the amount of the PSM antigen in a biological sample comprising steps of a) contacting the biological sample with at least one PSM ligand to form a complex with said ligand and the PSM antigen, and b) measuring the amount of the PSM antigen in said biological sample by measuring the amount of said complex.

This invention provides a method to purify mammalian PSM antigen comprising steps of a) coupling the antibody directed against PSM antigen to a solid matrix; b) incubating the coupled antibody of a) with a cell lysate containing PSM antigen under the condition permitting binding of the antibody and PSM antigen c) washing the coupled solid matrix to eliminate impurities and d) eluting the PSM antigen from the bound antibody.

This invention further provides transgenic nonhuman mammals which comprises an isolated nucleic acid molecule of PSM antigen. This invention also provides a transgenic nonhuman mammal whose genome comprises antisense DNA complementary to DNA encoding a mammalian PSM antigen so placed as to be transcribed into antisense mRNA complementary to mRNA encoding the PSM antigen and which hybridizes to mRNA encoding the PSM antigen thereby reducing its translation.

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells comprising introducing a DNA molecule encoding a prostate specific membrane antigen operatively linked to a regulatory element into a tumor cell of a subject, in a way that expression of the prostate specific membrane antigen is under the control of the regulatory element, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells.

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells, comprising introducing a DNA molecule encoding a prostate specific membrane antigen operatively linked to a regulatory element coupled with a therapeutic DNA into a tumor cell of a subject, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells.

This invention provides a method of detecting hematogenous micrometastatic tumor cells of a subject, comprising (A) performing nested polymerase chain reaction (PCR) on blood, bone marrow, or lymph node samples of the subject using the prostate specific membrane antigen primers, and (B) verifying micrometastases by DNA sequencing and Southern analysis, thereby detecting hematogenous micrometastatic tumor cells of the subject.

This invention provides a method of abrogating the mitogenic response due to transferrin, comprising introducing a DNA molecule encoding prostate specific membrane antigen operatively linked to a regulatory element into a tumor cell, the expression of which gene is directly associated with a defined pathological effect within a multicellular organism, thereby abrogating mitogen response due to transferrin.

DETAILED DESCRIPTION OF THE INVENTION

Throughout this application, references to specific nucleotides are to nucleotides present on the coding strand of the nucleic acid. The following standard abbreviations are used throughout the specification to indicate specific nucleotides:

C = cytosine
A = adenosine
T = thymidine
G = guanosine

A “gene” means a nucleic acid molecule, the sequence of which includes all the information required for the normal regulated production of a particular protein, including the structural coding sequence, promoters and enhancers.

This invention provides an isolated mammalian nucleic acid encoding a mammalian prostate-specific membrane (PSM) antigen.

This invention further provides an isolated mammalian DNA molecule of an isolated mammalian nucleic acid molecule encoding a mammalian prostate-specific membrane antigen. This invention also provides an isolated mammalian cDNA molecule encoding a mammalian prostate-specific membrane antigen. This invention provides an isolated mammalian RNA molecule encoding a mammalian prostate-specific membrane antigen.
In the preferred embodiment of this invention, the isolated nucleic sequence is cDNA from human as shown in sequence ID number 1. This human sequence was submitted to GenBank (Los Alamos National Laboratory, Los Alamos, N.M.) with Accession Number, M99487 and the description as PSM, Homo sapiens, 2653 base-pairs. This invention also encompasses DNAs and cDNAs which encode amino acid sequences which differ from those of PSM antigen, but which should not produce phenotypic changes. Alternatively, this invention also encompasses DNAs and cDNAs which hybridize to the DNA and cDNA of the subject invention. Hybridization methods are well known to those of skill in the art.

The DNA molecules of the subject invention also include DNA molecules coding for polypeptide analogs, fragments or derivatives of antigenic polypeptides which differ from naturally occurring forms in terms of the identity or location of one or more amino acid residues (deletion analogs containing less than all of the residues specified for the protein, substitution analogs wherein one or more residues specified are replaced by other residues and addition analogs where in one or more amino acid residues is added to a terminal or medial portion of the polypeptides) and which share some or all properties of naturally-occurring forms. These molecules include: the incorporation of codons “preferred” for expression by selected non-mammalian hosts; the provision of sites for cleavage by restriction endonuclease enzymes; and the provision of additional initial, terminal or intermediate DNA sequences that facilitate construction of readily expressed vectors.

The DNA molecules described and claimed herein are useful for the information which they provide concerning the amino acid sequence of the polypeptide and as products for the large scale synthesis of the polypeptide by a variety of recombinant techniques. The molecule is useful for generating new cloning and expression vectors, transformed and transfected prokaryotic and eukaryotic host cells, and new and useful methods for cultured growth of such host cells capable of expression of the polypeptide and related products.

Moreover, the isolated mammalian nucleic acid molecules encoding a mammalian prostate-specific membrane antigen are useful for the development of probes to study the tumorigenesis of prostate cancer.

This invention also provides nucleic acid molecules of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule encoding the prostate-specific membrane antigen.

This nucleic acid molecule produced can either be DNA or RNA. As used herein, the phrase “specifically hybridizing” means the ability of a nucleic acid molecule to recognize a nucleic acid sequence complementary to its own and to form double-helical segments through hydrogen bonding between complementary base pairs.

This nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule encoding the prostate-specific membrane antigen can be used as a probe. Nucleic acid probe technology is well known to those skilled in the art who will readily appreciate that such probes may vary greatly in length and may be labeled with a detectable label, such as a radioisotope or fluorescent dye, to facilitate detection of the probe. DNA probe molecules may be produced by insertion of a DNA molecule which encodes PSM antigen into suitable vectors, such as plasmids or bacteriophages, followed by transforming into suitable bacterial host cells, replication in the transformed bacterial host cells and harvesting of the DNA probes, using methods well known in the art. Alternatively, probes may be generated chemically from DNA synthesizers.

RNA probes may be generated by inserting the PSM antigen molecule downstream of a bacteriophage promoter such as T3, T7 or SP6. Large amounts of RNA probe may be produced by incubating the labeled nucleotides with the linearized PSM antigen fragment where it contains an upstream promoter in the presence of the appropriate RNA polymerase.

This invention also provides a nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule which is complementary to the mammalian nucleic acid molecule encoding a mammalian prostate-specific membrane antigen. This molecule may either be a DNA or RNA molecule.

The current invention further provides a method of detecting the expression of a mammalian PSM antigen expression in a cell which comprises obtaining total mRNA from the cell, contacting the mRNA so obtained with a labelled nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of the nucleic acid molecule encoding a mammalian PSM antigen under hybridizing conditions, determining the presence of mRNA hybridized to the molecule and thereby detecting the expression of the mammalian prostate-specific membrane antigen in the cell. The nucleic acid molecules synthesized above may be used to detect expression of a PSM antigen by detecting the presence of mRNA coding for the PSM antigen. Total mRNA from the cell may be isolated by many procedures well known to a person of ordinary skill in the art. The hybridizing conditions of the labelled nucleic acid molecules may be determined by routine experimentation well known in the art. The presence of mRNA hybridized to the probe may be determined by gel electrophoresis or other methods known in the art. By measuring the amount of the hybrid made, the expression of the PSM antigen by the cell can be determined. The labelling may be radioactive. For an example, one or more radioactive nucleotides can be incorporated in the nucleic acid when it is made.

In one embodiment of this invention, nucleic acids are extracted by precipitation from lysed cells and the mRNA is isolated from the extract using an oligo-dT column which binds the poly-A tails of the mRNA molecules (13). The mRNA is then exposed to radioactively labelled probe on a nitrocellulose membrane, and the probe hybridizes to and thereby labels complementary mRNA sequences. Binding may be detected by luminescence autoradiography or scintillation counting. However, other methods for performing these steps are well known to those skilled in the art, and the discussion above is merely an example.

This invention further provides another method to detect expression of a PSM antigen in tissue sections which comprises contacting the tissue sections with a labelled nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of nucleic acid molecules encoding a mammalian PSM antigen under hybridizing conditions, determining the presence of mRNA hybridized to the molecule and thereby detecting the expression of the mammalian PSM antigen in tissue sections. The probes are also useful for in-situ hybridization or in order to locate tissues which express this gene, or for other hybridization assays for the presence of this gene or its mRNA in various biological tissues. The in-situ hybridization using a labelled nucleic acid molecule is well known in the art. Essentially, tissue sections are incubated with the labelled nucleic acid molecule to allow the hybridization to occur. The molecule will carry a marker for the detection because it is "labelled", the amount of the
hybrid will be determined based on the detection of the amount of the marker and so will the expression of PSM antigen.

This invention further provides isolated PSM antigen nucleic acid molecule operatively linked to a promoter of RNA transcription. The isolated PSM antigen sequence can be linked to vector systems. Various vectors including plasmid vectors, cosmid vectors, bacteriophage vectors and other viruses are well known to ordinary skilled practitioners. This invention further provides a vector which comprises the isolated nucleic acid molecule encoding for the PSM antigen.

As an example to obtain these vectors, insert and vector DNA can both be exposed to a restriction enzyme to create complementary ends on both molecules which base pair with each other and are then ligated together with DNA ligase. Alternatively, linkers can be ligated to the insert DNA which correspond to a restriction site in the vector DNA, which is then digested with the restriction enzyme which cuts at that site. Other means are also available and known to an ordinary skilled practitioner.

In an embodiment, the PSM sequence is cloned in the Not I/Sal I site of pSPORT/Vector (Gibco-BRL). This plasmid, p55A-PSM, was deposited on Aug. 14, 1992 with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Md. 20852, U.S.A, under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganism for the Purposes of Patent Procedure. Plasmid, p55A-PSM, was accorded ATCC Accession Number 75294.

This invention further provides a host vector system for the production of a polypeptide having the biological activity of the prostate-specific membrane antigen. These vectors may be transformed into a suitable host cell to form a host cell vector system for the production of a polypeptide having the biological activity of PSM antigen.

Regulatory elements required for expression include promoter sequences to bind RNA polymerase and transcription initiation sequences for ribosome binding. For example, a bacterial expression vector includes a promoter such as the lac promoter and for transcription initiation the Shine-Dalgarno sequence and the start codon AUG (14). Similarly, a eukaryotic expression vector includes a heterologous or homologous promoter for RNA polymerase II, a downstream polyadenylation signal, the start codon AUG, and a termination codon for detachment of the ribosome. Such vectors may be obtained commercially or assembled from the sequences described by methods well known in the art, for example the methods described above for constructing vectors in general. Expression vectors are useful to produce cells that express the PSM antigen.

This invention further provides an isolated DNA or cDNA molecule described hereinabove wherein the host cell is selected from the group consisting of bacterial cells (such as E. coli), yeast cells, fungal cells, insect cells and animal cells. Suitable animal cells include, but are not limited to Vero cells, HeLa cells, Cos cells, CVI cells and various primary mammalian cells.

This invention further provides a method of producing a polypeptide having the biological activity of the prostate-specific membrane antigen which comprising growing host cells of a vector system containing the PSM antigen sequence under suitable conditions permitting production of the polypeptide and recovering the polypeptide so produced.

This invention provides a mammalian cell comprising a DNA molecule encoding a mammalian PSM antigen and the regulatory elements necessary for expression of the DNA in the mammalian cell so located relative to the DNA encoding the mammalian PSM antigen as to permit expression thereof.

Numerous mammalian cells may be used as hosts, including, but not limited to, the mouse fibroblast cell NIH3T3, CHO cells, HeLa cells, ltk cells, Cos cells, etc. Expression plasmids such as that described supra may be used to transfect mammalian cells by methods well known in the art such as calcium phosphate precipitation, electroporation or DNA encoding the mammalian PSM antigen may be otherwise introduced into mammalian cells, e.g., by microinjection, to obtain mammalian cells which comprise DNA, e.g., cDNA or a plasmid, encoding a mammalian PSM antigen.

This invention provides a method for determining whether a ligand can bind to a mammalian prostate-specific membrane antigen which comprises contacting a mammalian cell comprising an isolated DNA molecule encoding a mammalian prostate-specific membrane antigen with the ligand under conditions permitting binding of ligands to the mammalian prostate-specific membrane antigen, and thereby determining whether the ligand binds to a mammalian prostate-specific membrane antigen.

This invention further provides ligands bound to the mammalian PSM antigen.

This invention also provides a therapeutic agent comprising a ligand identified by the above-described method and a cytotoxic agent conjugated thereto. The cytotoxic agent may either be a radioisotope or a toxin. Examples of radioisotopes or toxins are well known to one of ordinary skill in the art.

This invention also provides a method of imaging prostate cancer in human patients which comprises administering to the patients at least one ligand identified by the above-described method, capable of binding to the cell surface of the prostate cancer cell and labelled with an imaging agent under conditions permitting formation of a complex between the ligand and the cell surface PSM antigen. This invention further provides a composition comprising an effective imaging agent of the PSM antigen ligand and a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers are well known to one of ordinary skill in the art. For example, such a pharmaceutically acceptable carrier can be physiological saline.

Also provided by this invention is a purified mammalian PSM antigen. As used herein, the term “purified prostate-specific membrane antigen” shall mean isolated naturally-occurring prostate-specific membrane antigen or protein (purified from nature or manufactured such that the primary, secondary and tertiary conformation, and posttranslational modifications are identical to naturally-occurring material) as well as non-naturally-occurring polypeptides having a primary structural conformation (i.e., continuous sequence of amino acid residues). Such polypeptides include derivatives and analogs.

This invention further provides a polypeptide encoded by the isolated mammalian nucleic acid sequence of PSM antigen.

It is believed that there may be natural ligand interacting with the PSM antigen. This invention provides a method to identify such natural ligand or other ligand which can bind to the PSM antigen. A method to identify the ligand comprises a) coupling the purified mammalian PSM antigen to a solid matrix, b) incubating the coupled purified mammalian PSM protein with the potential ligands under the conditions permitting binding of ligands and the purified PSM antigen; c) washing the ligand and coupled purified mammalian PSM antigen complex formed in b) to eliminate the nonspecific
binding and impurities and finally d) eluting the ligand from the bound purified mammalian PSM antigen. The techniques of coupling proteins to a solid matrix are well known in the art. Potential ligands may either be deduced from the structure of mammalian PSM or by other empirical experiments known by ordinary skilled practitioners. The conditions for binding may also easily be determined and protocols for carrying out such experimentation have long been well documented (15). The ligand-PSM antigen complex will be washed. Finally, the bound ligand will be eluted and characterized. Standard ligands characterization techniques are well known in the art.

The above method may also be used to purify ligands from any biological source. For purification of natural ligands in the cell, cell lysates, serum or other biological samples will be used to incubate with the mammalian PSM antigen bound on a matrix. Specific natural ligand will then be identified and purified as above described.

With the protein sequence information, antigenic areas may be identified and antibodies directed against these areas may be generated and targeted to the prostate cancer for imaging the cancer or therapies.

This invention provides an antibody directed against the amino acid sequence of a mammalian PSM antigen.

This invention provides a method to select specific regions on the PSM antigen to generate antibodies. The protein sequence may be determined from the PSM DNA sequence. Amino acid sequences may be analyzed by methods well known to those skilled in the art to determine whether they produce hydrophobic or hydrophilic regions in the proteins which they build. In the case of cell membrane proteins, hydrophobic regions are well known to form the part of the protein that is inserted into the lipid bilayer of the cell membrane, while hydrophilic regions are located on the cell surface, in an aqueous environment. Usually, the hydrophilic regions will be more immunogenic than the hydrophobic regions. Therefore the hydrophilic amino acid sequences may be selected and used to generate antibodies specific to mammalian PSM antigen. For example, hydrophilic sequences of the human PSM antigen shown in hydrophilicity plot of FIG. 16 may be easily selected. The selected peptides may be prepared using commercially available machines. As an alternative, DNA, such as a cDNA or a fragment thereof, may be cloned and expressed and the resulting polypeptide recovered and used as an immunogen.

Polyclonal antibodies against these peptides may be produced by immunizing animals using the selected peptides. Monoclonal antibodies are prepared using hybridoma technology by fusing antibody producing B cells from immunized animals with myeloma cells and selecting the resulting hybridoma cell line producing the desired antibody. Alternatively, monoclonal antibodies may be produced by in vitro techniques known to a person of ordinary skill in the art. These antibodies are useful to detect the expression of mammalian PSM antigen in living animals, in humans, or in biological tissues or fluids isolated from animals or humans.

In one embodiment, peptides Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. 35), Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. 36) and Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. 37) of human PSM antigen are selected.

This invention further provides polyvalent and monoclonal antibody(ies) against peptides Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. 35), Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. 36) and Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. 37).

This invention provides a therapeutic agent comprising antibodies or ligand(s) directed against PSM antigen and a cytotoxic agent conjugated thereto or antibodies linked enzymes which activate prodrug to kill the tumor. The cytotoxic agent may either be a radioisotope or toxin.

This invention provides a method of imaging prostate cancer in human patients which comprises administering to the patient the monoclonal antibody directed against the peptide of the mammalian PSM antigen capable of binding to the cell surface of the prostate cancer cell and labeled with an imaging agent under conditions permitting formation of a complex between the monoclonal antibody and the cell surface prostate-specific membrane antigen. The imaging agent is a radioisotope such as Indium.

This invention further provides a prostate cancer specific imaging agent comprising the antibody directed against PSM antigen and a radioisotope conjugated thereto.

This invention also provides a composition comprising an effective imaging amount of the antibody directed against the PSM antigen and a pharmaceutically acceptable carrier. The methods to determine effective imaging amounts are well known to a skilled practitioner. One method is by titration using different amounts of the antibody.

This invention further provides an immunoassay for measuring the amount of the prostate-specific membrane antigen in a biological sample comprising steps of a) contacting the biological sample with at least one antibody directed against the PSM antigen to form a complex with said antibody and the prostate-specific membrane antigen, and b) measuring the amount of the prostate-specific membrane antigen in said biological sample by measuring the amount of said complex. One example of the biological sample is a serum sample.

This invention provides a method to purify mammalian prostate-specific membrane antigen comprising steps of a) coupling the antibody directed against the PSM antigen to a solid matrix; b) incubating the coupled antibody of a) with lysate containing prostate-specific membrane antigen under the condition which the antibody and prostate membrane specific can bind; c) washing the solid matrix to eliminate impurities and d) eluting the prostate-specific membrane antigen from the coupled antibody.

This invention also provides a transgenic nonhuman mammal which comprises the isolated nucleic acid molecule encoding a mammalian PSM antigen. This invention further provides a transgenic nonhuman mammal whose genome comprises antisense DNA complementary to DNA encoding a mammalian prostate-specific membrane antigen so placed as to be transcribed into antisense mRNA complementary to mRNA encoding the prostate-specific membrane antigen and which hybridizes to mRNA encoding the prostate specific antigen thereby reducing its translation.

Animal model systems which elucidate the physiological and behavioral roles of mammalian PSM antigen are produced by creating transgenic animals in which the expression of the PSM antigen is either increased or decreased, or the amino acid sequence of the expressed PSM antigen is altered, by a variety of techniques. Examples of these techniques include, but are not limited to: 1) Insertion of normal or mutant versions of DNA encoding a mammalian PSM antigen, by microinjection, electroporation, retroviral transfection or other means well known to those skilled in the art, into appropriate fertilized embryos in order to produce a transgenic animal (16) or 2) Homologous recombination (17) of mutant or normal, human or animal versions of these genes with the native gene locus in transgenic animals to alter the regulation of expression or the structure of these PSM antigen sequences. The technique of homologous recombination is well known in the art. It replaces the native gene with the inserted gene and so is useful for producing an animal that cannot express native PSM antigen but does express, for
example, an inserted mutant PSM antigen, which has replaced the native PSM antigen in the animal's genome by recombination, resulting in underexpression of the transporter. Microinjection adds genes to the genome, but does not remove them, and so is useful for producing an animal which expresses its own and added PSM antigens, resulting in overexpression of the PSM antigens.

One means available for producing a transgenic animal, with a mouse as an example, is as follows: Female mice are mated, and the resulting fertilized eggs are dissected out of their oviducts. The eggs are stored in an appropriate medium such as M2 medium (16). DNA or cDNA encoding a mammalian PSM antigen is purified from a vector by methods well known in the art. Inducible promoters may be fused with the coding region of the DNA to provide an experimental means to regulate expression of the trans-gene. Alternatively or in addition, tissue specific regulatory elements may be fused with the coding region to permit tissue-specific expression of the trans-gene. The DNA, in an appropriately buffered solution, is put into a microinjection needle (which may be made from capillary tubing using a pipet puller) and the egg to be injected is put in a depression slide. The needle is inserted into the pronucleus of the egg, and the DNA solution is injected. The injected egg is then transferred into the oviduct of a pseudopregnant mouse (a mouse stimulated by the appropriate hormones to maintain pregnancy but which is not actually pregnant), where it proceeds to the uterus, implants, and develops to term. As noted above, microinjection is not the only method for inserting DNA into the egg cell, and is used here only for exemplary purposes.

Another use of the PSM antigen sequence is to isolate homologous gene or genes in different mammals. The gene or genes can be isolated by low stringency screening of either cDNA or genomic libraries of different mammals using probes from PSM sequence. The positive clones identified will be further analyzed by DNA sequencing techniques which are well known to an ordinary person skilled in the art. For example, the detection of members of the protein serine kinase family by homology probing (18).

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells comprising introducing a DNA molecule encoding a prostate specific membrane antigen operatively linked to a 5' regulatory element into a tumor cell of a subject, in a way that expression of the prostate specific membrane antigen is under the control of the regulatory element, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells. The subject may be a mammal or more specifically a human.

In one embodiment, the DNA molecule encoding prostate specific membrane antigen operatively linked to a 5' regulatory element forms part of a transfer vector which is inserted into a cell or organism. In addition the vector is capable of replication and expression of prostate specific membrane antigen. The DNA molecule encoding prostate specific membrane antigen can be integrated into a genome of a eukaryotic or prokaryotic cell or in a host cell containing and/or expressing a prostate specific membrane antigen.

Further, the DNA molecule encoding prostate specific membrane antigen may be introduced by a bacterial, viral, fungal, animal, or liposomal delivery vehicle. Other means are also available and known to an ordinary skilled practitioner.

Further, the DNA molecule encoding a prostate specific membrane antigen operatively linked to a promoter or enhancer. A number of viral vectors have been described including those made from various promoters and other regulatory elements derived from virus sources. Promoters consist of short arrays of nucleic acid sequences that interact specifically with cellular proteins involved in transcription. The combination of different recognition sequences and the cellular concentration of the cognate transcription factors determines the efficiency with which a gene is transcribed in a particular cell type.

Examples of suitable promoters include a viral promoter. Viral promoters include: adenovirus promoter, an simian virus 40 (SV40) promoter, a cytomegalovirus (CMV) promoter, a mouse mammary tumor virus (MMTV) promoter, a Malony murine leukemia virus promoter, a murine sarcoma virus promoter, and a Rous sarcoma virus promoter.

Further, another suitable promoter is a heat shock promoter. Additionally, a suitable promoter is a bacteriophage promoter. Examples of suitable bacteriophage promoters include but not limited to, a T7 promoter, a T3 promoter, an SP6 promoter, a lambda promoter, a baculovirus promoter.

Also suitable as a promoter is an animal cell promoter such as an interferon promoter, a metallothionein promoter, an immunoglobulin promoter. A fungal promoter is also a suitable promoter. Examples of fungal promoters include but are not limited to, an ADC1 promoter, an ARG promoter, an ADH1 promoter, a CYC1 promoter, a CUP promoter, an ENO1 promoter, a GAL promoter, a PRO promoter, a PGK promoter, a GAPDH promoter, a mating type factor promoter. Further, plant cell promoters and insect cell promoters are also suitable for the methods described herein.

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells, comprising introducing a DNA molecule encoding a prostate specific membrane antigen operatively linked to a 5' regulatory element coupled with a therapeutic DNA into a tumor cell of a subject, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells. The subject may be a mammal or more specifically a human.

Further, the therapeutic DNA which is coupled to the DNA molecule encoding a prostate specific membrane antigen operatively linked to a 5' regulatory element into a tumor cell may code for a cytokine, viral antigen, or a pro-drug activating enzyme. Other means are also available and known to an ordinary skilled practitioner.

The cytokine used may be interleukin-2, interleukin-12, interferon alpha, beta or gamma, granulocyte macrophage—colony stimulating factor, or other immunity factors.

In addition, this invention provides a prostate tumor cell, comprising a DNA molecule isolated from mammalian nucleic acid encoding a mammalian prostate-specific membrane antigen under the control of a prostate specific membrane antigen operatively linked to a 5' regulatory element.

As used herein, DNA molecules include complementary DNA (cDNA), synthetic DNA, and genomic DNA.

This invention provides a therapeutic vaccine for preventing human prostate tumor growth or stimulation of prostate tumor cells in a subject, comprising administering an effective amount to the prostate cell, and a pharmaceutical acceptable carrier, thereby preventing the tumor growth or stimulation of tumor cells in the subject. Other means are also available and known to an ordinary skilled practitioner.

This invention provides a method of detecting hematogenous micrometastatic tumor cells of a subject, comprising (A) performing nested polymerase chain reaction (PCR) on blood, bone marrow or lymph node samples of the subject using the prostate specific membrane antigen primers, and
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(B) verifying micrometastases by DNA sequencing and Southern analysis, thereby detecting hematogenous micrometastatic tumor cells of the subject. The subject may be a mammal or more specifically a human.

The micrometastatic tumor cell may be a prostate cancer and the DNA primers may be derived from prostate specific antigen. Further, the subject may be administered with simultaneously an effective amount of hormones, so as to increase expression of prostate specific membrane antigen.

This invention provides a method of abrogating the mitogenic response due to transferrin, comprising introducing a DNA molecule encoding prostate specific membrane antigen operatively linked to a 5 regulatory element into a tumor cell, the expression of which gene is directly associated with a defined pathological effect within a multicellular organism, thereby abrogating mitogen response due to transferrin. The tumor cell may be a prostate cell.

This invention will be better understood from the Experimental Details which follow. However, one skilled in the art will readily appreciate that the specific methods and results discussed are merely illustrative of the invention as described more fully in the claims which follow thereafter.

EXPERIMENTAL DETAILS

First Series of Experiments

Materials and Methods

The approach for cloning the gene involved purification of the antigen in large quantities by immunoprecipitation, and microsequencing of several internal peptides for use in synthesizing degenerate oligonucleotide primers for subsequent use in the polymerase chain reaction (19, 20). A partial CDNA was amplified as a PCR product and this was used as a homologous probe to clone the full-length cDNA molecule from a LNCaP (Lymph Node Carcinoma of Prostate) cell line cDNA plasmid library (8). Early experiments revealed to us that the CYT-356 antibody (9) was not capable of detecting the antigen produced in bacteria since the epitope was the glycosylated portion of the PSM antigen, and this necessitated our more difficult, yet elaborate approach.

Western Analysis of the PSM Antigen

Membrane proteins were isolated from cells by hypotonic lysis followed by centrifugation over a sucrose density gradient (21). 10-20 µg of LNCaP, DU-145, and PC-3 membrane proteins were electrophoresed through a 10% SDS-PAGE resolving gel with a 4% stacking gel at 9-10 milliamps for 16-18 hours. Proteins were electrophoronted onto PVDF membranes (Millipore Corp.) in transfer buffer (48 mM Tris base, 39 mM Glycine, 20% Methanol) at 25 volts overnight at 4°C. Membranes were blocked in TSB (0.15M NaCl, 0.01M Tris base, 5% BSA) for 30 minutes at room temperature followed by incubation with 10-15 µg/ml of CYT-356 monoclonal antibody (Cytogen Corp.) for 2 hours. Membranes were then incubated with 10-15 µg/ml of rabbit anti-mouse immunoglobulin (Accurate Scientific) for 1 hour at room temperature followed by incubation with 125I-Protein A (Amersham) at 1x10^6 cpm/ml at room temperature. Membranes were then washed and autoradiographed for 12-24 hours at -70°C. (FIG. 1).

Immunochemical Analysis of PSM Antigen Expression

The avidin-biotin method of immunochemical detection was employed to analyze both human tissue sections and cell lines for PSM Antigen expression (22). Cryostat-cut prostate tissue sections (4-6% thick) were fixed in methanol/acetone for 10 minutes. Cell cytopsins were made on glass slides using 50,000 cells/100 µl/slide. Samples were treated with 1% hydrogen peroxide in PBS for 10-15 minutes in order to remove any endogenous peroxidase activity. Tissue sections were washed several times in PBS, and then incubated with the appropriate suppressor serum for 20 minutes. The suppressor serum was drained off and the sections or cells were then incubated with the diluted CYT-356 monoclonal antibody for 1 hour. Samples were then washed with PBS and sequentially incubated with secondary antibodies (horse or goat immunoglobulins, 1:200 dilution for 30 minutes), and with avidin-biotin complexes (1:25 dilution for 30 minutes). DAB was used as a chromogen, followed by hematoxylin counterstaining and mounting. Frozen sections of prostate samples and duplicate cell cytopsins were used as controls for each experiment. As a positive control, the anti-cytokeratin monoclonal antibody CAM 5.2 was used following the same procedure described above. Tissue sections are considered by us to express the PSM antigen if at least 5% of the cells demonstrate immunoreactivity. Our scoring system is as follows: 1<5%; 2<5-15%; 3<20-75%; and 4>75% positive cells. Homogeneity versus heterogeneity was accounted for by evaluating positive and negative cells in 3-5 high power light microscopic fields (400x), recording the percentage of positive cells among 100-500 cells. The intensity of immunostaining is graded on a 1+ to 4+ scale, where 1-represents mild, 2-3+ represents moderate, and 4+ represents intense immunostaining as compared to positive controls.

Immunoprecipitation of the PSM Antigen

80%-confluent LNCaP cells in 100 mm petri dishes were starved in RPMI media without methionine for 2 hours, after which 35S-Methionine was added at 100 µCi/ml and the cells were grown for another 16-18 hours. Cells were then washed and lysed by the addition of 1 ml of lysis buffer (1% Triton X-100, 50 mM Hepes pH 7.5, 10% glycerol, 150 mM MgCl2, 1 mM PMSF, and 1 mM EGTA) with incubation for 20 minutes at 40°C. Lysates were pre-cleared by mixing with Pansorbin® cells (Calbiochem) for 90 minutes at 4°C. Cell lysates were then mixed with Protein A Sepharose® CL-4B beads (Pharmacia®) previously bound with CYT-356 antibody (Cytogen Corp.) and RAM antibody (Accurate Scientific) for 3-4 hours at 4°C. 12 µg of antibody was used per 3 mg of beads per petri dish. Beads were then washed with HNTG buffer (20 mM Hepes pH 7.5, 150 mM NaCl, 0.1% Triton X-100, 10% glycerol, and 2 mM Sodium Orthovana- date), resuspended in sample loading buffer containing mercaptoethanol, denatured at 95°C for 5-10 minutes and run on a 10% SDS-PAGE gel with a 4% stacking gel at 10 milliamps overnight. Gels were stained with Coomassie Blue, destained with acetic acid/methanol, and dried down in a vacuum dryer at 60°C. Gels were then autoradiographed for 16-24 hours at -70°C. (FIG. 2 A-D).

Large-Scale Immunoprecipitation and Peptide Sequencing

The procedure described above for immunoprecipitation was repeated with 8 confluent petri dishes containing approximately 6x10^7 LNCaP cells. The immunoprecipitation product was pooled and loaded into two lanes of a 10% SDS-PAGE gel and electrophoresed at 9-10 milliamps for 16 hours. Proteins were electrophoronted onto Nitrocellulose BA-85 membranes (Schleicher and Schuell®) for 2 hours at 75 volts for 4°C in transfer buffer. Membranes were stained with Ponceau Red to visualize the proteins and the 100 KD protein band was excised, solubilized, and digested proteolytically with trypsin. HPLC was then performed on the digested sample on an Applied Biosystems Model 171C and clear dominant peptide peaks were selected and sequenced by modified Edman degradation on a modified post liquid Applied Biosystems Model 477A Protein/Peptide Microsequencer (23). Sequencing data on all of the peptides is
included within this document. We attempted to sequence the amino-terminus of the PSM antigen by a similar method which involved purifying the antigen by immunoprecipitation and transfer via electro-blotting to a PVDF membrane (Millipore®). Protein was analyzed on an Applied Biosystems Model 477A Protein/Peptide Sequencer and the amino terminus was found to be blocked, and therefore no sequence data could be obtained by this technique.

<table>
<thead>
<tr>
<th>PSM Antigen Peptide Sequences:</th>
</tr>
</thead>
<tbody>
<tr>
<td>2717 SLYES(W)TK (SEQ ID No. 3)</td>
</tr>
<tr>
<td>2722 (S)YPDGNLFG(g)QQR (SEQ ID No. 4)</td>
</tr>
<tr>
<td>2726 FYDPMFK (SEQ ID No. 5)</td>
</tr>
<tr>
<td>2727 INVNGTI(K) (SEQ ID No. 6)</td>
</tr>
<tr>
<td>2734 FLIVXIPILGATGEQNLPGK (SEQ ID No. 7)</td>
</tr>
<tr>
<td>2735 G/P/V/L/YSDPADYPAF/DGVK (SEQ ID No. 8, 9)</td>
</tr>
<tr>
<td>2738 AFDPLGLPDRPFYR (SEQ ID No. 10)</td>
</tr>
<tr>
<td>2746 YAGESPIGDYDALFIESK (SEQ ID No. 11)</td>
</tr>
<tr>
<td>2747 TILFAS(W)/DAEFGQX(q)STE(c)(E).... (SEQ ID No. 12)</td>
</tr>
</tbody>
</table>

Notes: X means that no residue could be identified at this position. Capital denotes identification but with a lower degree of confidence. (Lower case) means residue present but at very low levels... indicates sequence continues but has dropped below detection limit.

All of these peptide sequences were verified to be unique after a complete homology search of the translated Genbank computer database.

Degenere PCR

Sense and anti-sense 5'-unphosphorylated degenerate oligonucleotide primers 17 to 20 nucleotides in length corresponding to portions of the above sequences were synthesized on an Applied Biosystems Model 394A DNA Synthesizer. These primers have degeneracies from 32 to 144. The primers used are shown below. The underlined amino acids in the peptides represent the residues used in primer design.

**Peptide 3:** (SEQ ID No. 5)

PSM Primer “A” T{C or T}—TA(C or T)—GA(C or T)—CCX—ATG—TT (SEQ ID No. 13)

PSM Primer “B” AAC—ATX—GG(A or G)—TC(A or G)—TA(A or G)—AA (SEQ ID No. 14)

Primer A is sense primer and B is anti-sense. Degeneracy is 32-fold.

**Peptide 4:** INYNVGTI(K) (SEQ ID No. 6)

PSM Primer “C” AT{T or C or A}—TA(T or C)—AA(T or C)—GTX—AT(T or C or A)—GG (SEQ ID No. 15)

PSM Primer “D” CC(A or T or G)—ATX13 AC(G or A)—TT(A or G) or —TA(A or G or T)—AT (SEQ ID No. 16)

Primer C is sense primer and D is anti-sense. Degeneracy is 144-fold.

**Peptide 2:** G/P/V/L/YSDPACYFAPF/DGVK (SEQ ID No. 8, 9)

PSM Primer “E” CCX—GCX—GA(T or C)—TA(T or C)—TT(T or C)—CC (SEQ ID No. 17)

PSM Primer “F” GC(G or A)—AA(A or G)—TA(A or G)—TXX—GCX—GG (SEQ ID No. 16)

Primer E is sense primer and F is anti-sense primer. Degeneracy is 128-fold.
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Tubes were overlaid with 60 µl of light mineral oil and amplified for 30 cycles. PCR products were analyzed by electrophoresing 5 µl of each sample on a 2-3% agarose gel followed by staining with Ethidium bromide and photography.

*10x PCR Buffer
166 mM NH₄SO₄
670 mM Tris, pH 8.8
2 mg/ml BSA

Representative photographs displaying PCR products are shown in FIG. 5.

Cloning of PCR Products

In order to further analyze these PCR products, these products were cloned into a suitable plasmid vector using “TA Cloning” (Invitrogen® Corp.). The cloning strategy employed here is to directly ligate PCR products into a plasmid vector possessing overhanging T residues at the insertion site, exploiting the fact that Tag polymerase leaves overhanging A residues at the ends of the PCR products. The ligation mixtures are transformed into competent E.coli cells and resulting colonies are grown up. Plasmid DNA is isolated by the alkaline lysis method (24), and screened by restriction analysis (FIG. 6 A-B).

DNA Sequencing of PCR Products

TA Clones of PCR products were then sequenced by the dideoxy method (25) using Sequenase (U.S. Biochemical). 3-4 µg of each plasmid DNA was denatured with NaOH and ethanol precipitated. Labeling reactions were carried out as per the manufacturer’s recommendations using 32P-ATP, and the reactions were terminated as per the same protocol. Sequencing products were then analyzed on 6% polyacrylamide/7M urea gels using an IBI sequencing apparatus. Gels were run at 120 watts for 2 hours. After electrophoresis, the gels were fixed for 15-20 minutes in 10% methanol/10% acetic acid, transferred onto Whatman 3MM paper and dried down in a BioRad® vacuum dryer at 80°C for 2 hours. Gels were then autoradiographed at room temperature for 16-24 hours. In order to determine whether the PCR products were the correct clones, we analyzed the sequences obtained at the 5' and 3' ends of the molecules looking for the correct primer sequences, as well as adjacent sequences which corresponded to portions of the peptides not used in the design of the primers.

IN-20 was confirmed to be correct and represent a partial cDNA for the PSM gene. In this PCR reaction, I and N primers were used. The DNA sequence we obtained when reading from the 1 primer was:

ACG GAG CAA A12c TTT CAG CTT GCA AAG

(SEQ ID No. 30)

T E O N P O L A X

(SEQ ID No. 31)

The underlined amino acids were the portion of peptide 6 that was used to design this sense primer and the remaining amino acids which agree with those present within our peptide confirm that this end of the molecule represents the correct protein (PSM antigen).

When we analyzed the other end of the molecule by reading from the N primer the sequence was:

CTC TTC GGC ATC CCA GGT TGC ALAA CAA ATT TGT TCT

(SEQ ID No. 32)

Since this represents the anti-sense DNA sequence, we need to show the complementary sense sequence in order to find our peptide.

Sense Sequence:

AGA ACA ATT TTT GCT AGC TGG GAT GCC AAG GAG

(SEQ ID No. 33)

R T I L P A S W D A E B

(SEQ ID No. 34)

The underlined amino acids here represent the portion of peptide 7 used to create primer N. All of the amino acids upstream of this primer are correct in the IN-20 clone, agreeing with the amino acids found in peptide 7. Further DNA sequencing has enabled us to identify the presence of our other PSM peptides within the DNA sequence of our positive clone.

The DNA sequence of this partial cDNA was found to be unique when screened on the Genbank computer database.

cDNA Library Construction and Cloning of Full—Length PSM cDNA

A cDNA library from LNCaP mRNA was constructed using the Superscript® plasmid system (BRL®-Gibco). The library was transformed using competent DH5-α cells and plated onto 100 mm plates containing LB plus 100 µg/ml of Carbenicillin. Plates were grown overnight at 37°C. Colonies were transferred to nitrocellulose filters. Filters were processed and-screened as per Gronstein and Hogness (26), using our 1.1 kb plasmid cDNA homologous probe which was radiolabelled with 32P-dCTP by random priming (27). We obtained eight positive colonies which upon DNA restriction and sequencing analysis proved to represent full-length cDNA molecules coding for the PSM antigen. Shown in FIG. 7 is a autoradiogram showing the size of the cDNA molecules represented in our library and in FIG. 8 restriction analysis of several full-length clones is shown. FIG. 9 is a plasmid Southern analysis of the samples in FIG. 6, showing that they all hybridize to the 1.1 kb partial cDNA probe.

Both the cDNA as well as the antigen have been screened through the Genbank Computer database (Human Genome Project) and have been found to be unique.

Northern Analysis of PSM Gene Expression

Northern analysis (28) of the PSM gene has revealed that expression is limited to the prostate and to prostate carcinoma.

RNA samples (either 10 µg of total RNA or 2 µg of poly A+RNA) were denatured and electrophoresed through 1.1% agarose/formaldehyde gels at 60 milliamps for 6-8 hours. RNA was then transferred to Nytran® nylon membranes (Schleicher and Schuell®) by pressure blotting in 10x SSC with a Posi-blotter (Stratagene®). RNA was cross-linked to the membranes using a Stratalink (Stratagene®) and subsequently baked in a vacuum oven at 80°C for 2 hours. Blots were pre-hybridized at 65°C for 2 hours in prehybridization solution (BRL®) and subsequently hybridized for 16 hours in hybridization buffer (BRL®) containing 1-2x10⁶ cpm/ml of 32P-labelled random-primed cDNA probe. Membranes were washed twice in 1x SSPE/1% SDS and twice in 0.1 x SSPE/1% SDS at 42°C. Membranes were then air-dried and autoradiographed for 12-36 hours at -70°C.
PCR Analysis of PSM Gene Expression in Human Prostate Tissues

PCR was performed on 15 human prostate samples to determine PSM gene expression. Five samples each from normal prostate tissue, benign prostatic hyperplasia, and prostate cancer were used (histology confirmed by MSKCC Pathology Department).

10 μg of total RNA from each sample was reverse transcribed to make cDNA template as previously described in section IV. The primers used corresponded to the 5' and 3' ends of our 1.1 kb partial cDNA, IN-20, and therefore the expected size of the amplified band is 1.1 kb. Since the Tm of our primers is 64°C, we annealed the primers in our PCR at 60°C. We carried out the PCR for 35 cycles using the same conditions previously described in section IV.

LNCaP and H26—Ras transfected LNCaP (29) were included as a positive control and DU-145 as a negative control. 14/15 samples clearly amplified the 1.1 kb band and therefore express the gene.

Experimental Results

The gene which encodes the 100 kd PSM antigen has been identified. The complete cDNA sequence is shown in Sequence ID #1. Underneath that nucleic acid sequence is the predicted translated amino acid sequence. The total number of the amino acids is 750, ID: #2. The hydrophilicity of the predicted protein sequence is shown in FIG. 16. Shown in FIG. 17 are three peptides with the highest point of hydrophilicity. They are: Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. 35); Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. 36) and Lys-Ser-Asp-Pro-Asp-Gly-Glu (SEQ ID No. 37).

By the method of Klein, Kaneshia and Delisi, a specific membrane-spanning domain is identified. The sequence is from the amino acid #19 to amino acid #44: Ala-Gly-Ala-Leu-Val-Leu-Aas-Gly-Gly-Phe-Phe-Leu-Leu-Gly-Phe-Leu (SEQ ID No. 38).

This predicted membrane-spanning domain was computed on PC Gene (computer software program). This data enables prediction of inner and outer membrane domains of the PSM antigen which aids in designing antibodies for uses in targeting and imaging prostate cancer.

When the PSM antigen sequence with other known sequences of the GeneBank were compared, homology between the PSM antigen sequence and the transferrin receptor sequence were found. The data are shown in FIG. 18.

EXPERIMENTAL DISCUSSIONS

Potential Uses for PSM Antigen:

1. Tumor detection:
Microscopic:

Unambiguous tumor designation can be accomplished by use of probes for different antigens. For prostatic cancer, the PSM antigen probe may prove beneficial. Thus PSM could be used for diagnostic purposes and this could be accomplished at the microscopic level using in-situ hybridization using sense (control) and antisense probes derived from the coding region of the cDNA cloned by the applicant. This could be used in assessment of local extraprostatic extension, involvement of lymph node, bone or other metastatic sites. As bone metastasis presents a major problem in prostatic cancer, early detection of metastatic spread is required especially for staging. In some tumors detection of tumor cells in bone marrow portends a grim prognosis and suggests that interventions aimed at metastasis be tried. Detection of PSM antigen expression in bone marrow aspirates or sections may provide such early information. PCR amplification or in-situ hybridization may be used. This could be developed for any possible metastatic region.

2. Antigen site identification

The knowledge of the cDNA for the antigen also provides for the identification of areas that would serve as good antigens for the development of antibodies for use against specific amino acid sequences of the antigen. Such sequences may be at different regions such as outside, membrane or inside of the PSM antigen. The development of these specific antibodies would provide for immunohistochemical identification of the antigen. These derived antibodies could then be developed for use, especially ones that work in paraffin fixed sections as well as frozen section as they have the greatest utility for immunodiagnosis.

3. Restriction fragment length polymorphism and genomic DNA

Restriction fragment length polymorphisms (RFLPS) have proven to be useful in documenting the progression of genetic damage that occurs during tumor initiation and promotion. It may be that RFLP analysis will demonstrate that changes in PSM sequence restriction mapping may provide evidence of predisposition to risk or malignant potential or progression of the prostatic tumor.

Depending on the chromosomal location of the PSM antigen, the PSM antigen gene may serve as a useful chromosome location marker for chromosome analysis.

4. Serum

With the development of antigen specific antibodies, if the antigen or selected antigen fragments appear in the serum they may provide for a serum marker for the presence of metastatic disease and be useful individually or in combination with other prostate specific markers.

5. Imaging

As the cDNA sequence implies that the antigen has the characteristics of a membrane spanning protein with the majority of the protein on the exofacial surface, antibodies, especially monoclonal antibodies to the peptide fragments exposed and specific to the tumor may provide for tumor imaging local extension of metastatic tumor or residual tumor following prostatectomy or irradiation. The knowledge of the coding region permits the generation of monoclonal antibodies and these can be used in combination to provide for maximal imaging purposes. Because the antigen shares a similarity with the transferrin receptor based on cDNA analysis (approximately 54%), it may be that there is a specific normal ligand for this antigen and that identification of the ligand(s) would provide another means of imaging.

6. Isolation of ligands

The PSM antigen can be used to isolate the normal ligand(s) that bind to it. These ligand(s) depending on specificity may be useful for targeting, or their serum levels may be predictive of disease status. If it is found that the normal ligand for PSM is a carrier molecule then it may be that PSM could be used to bind to that ligand for therapy purposes (like an iron chelating substance) to help remove the ligand from the circulation. If the ligand promotes tumor growth or metastasis then providing soluble PSM antigen would remove the ligand from binding the prostate. Knowledge of PSM antigen structure could lend to generation of small fragment that binds ligand which could serve the same purpose.

7. Therapeutic uses

a) Ligands. The knowledge that the cDNA structure of PSM antigen shares structural homology with the transferrin receptor (54% on the nucleic acid level) implies that there may be an endogenous ligand for the receptor that may or may not be transferrin-like. Transferrin is thought to be a ligand
that transports iron into the cell after binding to the transferrin receptor. However, apotransferrin is being reported to be a growth factor for some cells which express the transferrin receptor (30). Whether transferrin is a ligand for this antigen or some other ligand binds to this ligand remains to be determined. If a ligand is identified it may carry a specific substance such as a metal ion (iron or zinc or other) into the tumor and thus serve as a means to deliver toxic substances (radioactive or cytotoxic chemical i.e. toxin like ricin or cytotoxic alkylating agent or cytotoxic prodrug) to the tumor.

The main metastatic site for prostatic tumor is the bone. The bone and bone stroma are rich in transferrin. Recent studies suggest that this microenvironment is what provides the right “soil” for prostatic metastasis in the bone (31). It may be that this also promotes attachment as well, these factors which reduce this ability may diminish prostatic metastasis to the bone and prostatic metastatic growth in the bone.

It was found that the ligand for the new antigen (thought to be an oncogene and marker of malignant phenotype in breast carcinoma) served to induce differentiation of breast cancer cells and thus could serve as a treatment for rather than promoter of the disease. It may be that ligand binding to the right region of PSM whether with natural ligand or with an antibody may serve a similar function.

Antibodies against PSM antigen coupled with a cytotoxic agent will be useful to eliminate prostate cancer cells. Transferrin receptor antibodies with toxin conjugates are cytotoxic to a number of tumor cells as tumor cells tend to express increased levels of transferrin receptor (32). Transferrin receptors take up molecules into the cell by endocytosis. Antibody drug combinations can be toxic. Transferrin linked toxin can be toxic.

b) Antibodies against PSM antigen coupled with a cytotoxic agent will be useful to eliminate prostate cancer cells. The cytotoxic agent may be a radioisootope or toxin as known in ordinary skill of the art. The linkage of the antibody and the toxin or radioisootope can be chemical. Examples of direct linked toxins are doxorubicin, chlorambucil, ricin, pseudomonas exotoxin etc. or a hybrid toxin can be generated 1/2 with specificity for PSM and the other 1/2 with specificity for the toxin. Such a bivalent molecule can serve to bind to the tumor and the other 1/2 to deliver a cytotoxic to the tumor or to bind to and activate a cytotoxic lymphocyte such as binding to the T1, T1 receptor complex. Antibodies of required specificity can also be cloned into T cells and by replacing the immunoglobulin domain of the T cell receptor (TCR) cloning in the desired MAb heavy and light chains; splicing the U4 and U3 gene segments with the constant regions of the α and β TCR chains and transfecting these chimeric Ab/TCR genes in the patients’ T cells, propagating these hybrid cells and infusing them into the patient (33). Specific knowledge of tissue specific antigens for targets and generation of MAb’s specific for such targets will help make this a usable approach. Because the PSM antigen coding region provides knowledge of the entire coding region, it is possible to generate a number of antibodies which could then be used in combination to achieve an additive or synergistic anti-tumor action. The antibodies can be linked to enzymes which can activate non-toxic prodrugs at its site of tumor such as Carboxypeptidase and 4-bis(2 chloroethyl) amino benzoxy-α-glutamatic acid and its active parent drug in mice (34).

It is possible to produce a toxic genetic chimera such as TP-40 a genetic recombinant that possesses the cDNA from TGF-alpha and the toxic portion of pseudomonas exotoxin so the TGF and portion of the hybrid binds the epidermal growth factor receptor (EGFR), and the pseudomonas portion gets uptake into the cell enzymatically and inactivates the ribosome ability to perform protein synthesis resulting in cell death. When we know the ligand for the PSM antigen we can do the same.

In addition, once the ligand for the PSM antigen is identified, toxin can be chemically conjugated to the ligands. Such conjugated ligands can be therapeutically useful. Examples of the toxins are daunomycin, chlorambucil, ricin, pseudomonas exotoxin, etc. Alternatively, chimeric construct can be created linking the cDNA of the ligand with the cDNA of the toxin. An example of such toxin is TGFα and pseudomonas exotoxin (35).

8. Others

The PSM antigen may have other uses. It is well known that the prostate is rich in zinc, if the antigen provides function relative to this or other biologic function the PSM antigen may provide utility in the treatment of other prostatic pathologies such as benign hyperplastic growth and/or prostatitis.

Because purified PSM antigen can be generated, the purified PSM antigen can be linked to beads and use it like a standard “affinity” purification. Serum, urine or other biological samples can be used to incubate with the PSM antigen bound onto beads. The beads may be washed thoroughly and then eluted with salt or pH gradient. The eluted material is SDS gel purified and used as a sample for microsequencing.

The sequences will be compared with other known proteins and if unique, the technique of degenerated PCR can be employed for obtaining the ligand. Once known, the affinity of the ligand will be determined by standard protocols (15).

REFERENCES OF THE FIRST SERIES OF EXPERIMENTS


SECOND SERIES OF EXPERIMENTS

Expression of the Prostate-Specific Membrane Antigen

Applicant's have recently cloned a 2.6-2.5 kb complementary DNA encoding PSMA, the prostate-specific membrane antigen recognized by the 7E11-C5.3 anti-prostate monoclonal antibody. Immunohistochemical analysis of the LNCaP, DU-145, and PC-3 prostate cancer cell lines for PSMA expression using the 7E11-C5.3 antibody reveals intense staining in the LNCaP cells, with no detectable expression in both the DU-145 and PC-3 cells. Coupled in vitro transcription/translation of the 2.65 kb full-length PSMA cDNA yields an 84 kDa protein corresponding to the predicted polypeptide molecular weight of PSMA. Post-translational modification of this protein with pancreatic canine microsomes yields the expected 100 kDa PSMA antigen. Following transfection of PC-3 cells with the full-length PSMA cDNA in a eukaryotic expression vector, the subject's activity confirms the expression of the PSMA glycoprotein by Western analysis using the 7E11-C5.3 monoclonal antibody. Ribonuclease protection analysis demonstrates that the expression of PSMA mRNA is almost entirely prostate-specific in human tissues. PSMA expression appears to be highest in hormone-deprived states and is hormonally modulated by steroids, with DHT downregulating PSMA expression in the human prostate cancer cell line LNCaP by 8-10 fold, testosterone downregulating PSMA by 3-4 fold, and corticosteroids showing no significant effect. Normal and malignant prostatic tissues consistently show high PSMA expression, whereas we have noted heterogeneous, and at times absent, expression of PSMA in benign prostatic hyperplasia. LNCaP tumors implanted and grown both orthotopically and subcutaneously in nude mice, abundantly express PSMA, providing an excellent in-vivo model system to study the regulation and modulation of PSMA expression.

EXPERIMENTAL DETAILS

Materials and Methods

Cells and Reagents:

The LNCaP, DU-145, and PC-3 cell lines were obtained from the American Type Culture Collection. Details regarding the establishment and characteristics of these cell lines have been previously published (5,7,8). Unless specified otherwise, LNCaP cells were grown in RPMI 1640 media supplemented with L-glutamine, nonessential amino acids,
US RE43,586 E

and 5% fetal calf serum (Gibco-BRL, Gaithersburg, Md.) in a 
CO₂ incubator at 37°C for 145 and PC-3 cells were grown in 
minimal essential medium supplemented with 10% fetal calf 
supernatant. All cell cultures were obtained from the MSKCC Media 
Preparation Facility. Restriction and modifying enzymes 
were purified from Gibco-BRL unless otherwise specified. 

We employed the avidin-biotin method of detection to 
analyze prostate cancer cell lines for PSM antigen expression (9A). Cell cytosplasm were made on glass slides using 5×10⁶ 
cells/100 µl per slide. Slides were washed twice with PBS and 
then incubated with the appropriate suppressor serum for 20 
minutes. The suppressor serum was drained off and the cells were 
incubated with diluted 7E11-CS.3 (5 g/ml) monoclonal 

antibody for 1 hour. Samples were then washed with PBS and 
sequentially incubated with secondary antibodies for 30 
minutes and avidin-biotin complexes for 30 minutes. Diamino-
benzidine served as our chromogen and color development 
followed by hematoxylin counter-staining and mounting. 
Duplicate cell cytosplasm were used as controls for each 
experiment. As a positive control, the anti-epidermal growth 
factor monoclonal antibody CAM 5.2 was used following the same 
procedure described above. Human EJ bladder carcinoma cells 
served as a negative control.

In Vitro Transcription/Translation of PSM Antigen

Plasmid 55A containing the full length 2.65 kb PSM cDNA in the 
plasmid pSPORT1 (Gibco-BRL) was transcribed in vitro using the Promega TNT system (Promega Corp. Madison, Wis.). T7 RNA polymerase was added to the cDNA in a 
reaction mixture containing rabbit reticulocyte lysate, an 

amino acid mixture lacking methionine, buffer, and 35S-Methionine (Amersham) and incubated at 30°C for 90 minutes. 
Post-translational modification of the resulting protein was 
accomplished by the addition of pancreatic canine 

micromoles into the reaction mixture (Promega Corp. Madison, Wis.). Protein products were analyzed by electrophoresis on 10% SDS-PAGE gels which were subsequently treated with 
Amplify autoradiography enhancer (Amershams, Arlington 
Heights, Ill.) according to the manufacturers instructions 
and dried at 80°C in a vacuum dryer. Gels were autoradiographed overnight at ~70°C using Hyperfilm MP (Amersham).

Transfection of PSM into PC-3 Cells

The full length PSM cDNA was subcloned into the pREP7 
eukaryotic expression vector (Invitrogen, San Diego, Calif.). 
Plasmid DNA was purified from transformed DH5-alpha bacte-
ria (Gibco-BRL) using Qiagen maxi-prep plasmid isolation 
columns (Qiagen Inc., Chatsworth, Calif.). Purified plasmid 
DNA (6.10 g) was diluted with 900 µl of Optimem media 
(Gibco-BRL) and mixed with 30 µl of Lipofectin reagent 
(Gibco-BRL) which had been previously diluted with 9001 of 
Optimem media. This mixture was added to T-75 flasks of 
40-50% confluent PC-3 cells in Optimem media. After 24-36 
hours, cells were trypsinized and split into 100 mm dishes 
containing RPMI 1640 media supplemented with 10% fetal 
calf serum and 1 mg/ml of Hygromycin B (Calbiochem, La Jolla, Calif.). The dose of Hygromycin B used was previously 
determined by a time course/dose response ctkt assay. Cells were maintained in this media for 2-3 weeks with changes of media and Hygromycin B every 4-5 days until 
discrete colonies appeared. Colonies were isolated using 
6 mm cloning cylinders and expanded in the same media. As a 
control, PC-3 cells were also transfected with the pREP7 
plasmid alone. RNA was isolated from the transfected cells and 
PMM mRNA expression was detected by both RNase 
Protection analysis (described later) and by Northern analy-

Western Blot Detection of PSM Expression

Crude protein lysates, were isolated from LNCaP, PC-3, 
and PSM-transfected PC-3 cells as previously described (10A). LNCaP cell membranes were also isolated according 
to published methods (10A). Protein concentrations were 
quantitated by the Bradford method using the BioRad protein 
reagent kit (BioRad, Richmond, Calif.). Following denatur-
ation, 20 g of protein was electrophoresed on a 10% SDS-
PAGE gel at 25 mA for 4 hours. Gels were electroblotted onto 
Immobilon P membranes (Millipore, Bedford, Mass.) 
overnight at 40C. Membranes were blocked in 0.15M NaCl/0.01M 
Tris-HCl (pH 7.5) plus 5% BSA followed by a 1 hour incubation 
with 7E11-CS.3 monoclonal antibody (10 g/ml). Blots were 
washed 4 times with 0.15M NaCl/0.01M Tris-HCl/0.05% 
Triton-X 100 (TS-X) and incubated for 1 hour with rabbit 
anti-mouse IgG (Accurate Scientific, Westbury, N.Y.) at 

concentration of 10 µg/ml.

Blots were then washed 4 times with TS-X and labeled with 
125I-Protein A (Amersham, Arlington Heights, Ill.) at a 
concentration of 1 million cpm/ml. Blots were then washed 4 
times with TS-X and dried on Whatman 3MM paper, fol-

lowed by overnight autoradiography at ~70°C using Hy-

perm film (Amersham). Orthotopic and Subcutaneous LNCaP Tumor Growth in 
Nude Mice

LNCaP cells were harvested from sub-confluent cultures 
by a one minute exposure to a solution of 0.25% trypsin 
and 0.02% EDTA. Cells were resuspended in RPMI 1640 media 
with 5% fetal bovine serum, washed and diluted in either 
Matrigel (Collaborative Biomedical Products, Bedford, 
Mass.) or calcium and magnesium-free Hank's balanced salt 
solution (HBSS). Only single cell suspensions with greater 
than 90% viability by trypan blue exclusion were used for in 
vivo injection. Male athymic Swiss (nu/nu) nude mice 4-6 
weeks of age were obtained from the Memorial Sloan-Ket-
ter Cancer Center Animal Facility. For subcutaneous 
tumor cell injection one million LNCaP cells resuspended in 
0.2 ml of Matrigel were injected into the hindlimb of each 
mouse using a disposable syringe fitted with a 28 gauge 
needle. For orthotopic injection, mice were first anesthetized 
with an intraperitoneal injection of Pentobarbital and placed in 
the supine position. The abdomen was cleansed with Beta-
dine and the prostate was exposed through a midline incision. 
2.5 million LNCaP tumor cells in 0.1 ml were injected 
directly into either posterior lobe using a 1 ml disposable 
syringe and a 28 gauge needle. LNCaP cells with and without 
Matrigel were injected. Abdominal closure was achieved in 
one layer using Autoclip wound clips (Clay Adams, Parsippany, N.J.). Tumors were harvested in 6-8 weeks, confirmed 
histologically by faculty of the Memorial Sloan-Kettering 
Cancer Center Pathology Department, and frozen in liquid 
nitrogen for subsequent RNA isolation.

RNA Isolation

Total cellular RNA was isolated from cells and tissues by 
standard techniques (11,12) as well as by using RNAzol B 
(Cinna/Biotech, Houston, Tex.). RNA concentrations and 

quality were assessed by UV spectrophotometry on a Beckman DU 
640 spectrophotometer and by gel analysis. Human tissue 
total RNA samples were purchased from Clontech Laborato-
ries, Inc., Palo Alto, Calif.

Ribonuclease Protection Assays

A portion of the PSM cDNA was subcloned into the plasmid 
vector pSPORT1 (Gibco-BRL) and the orientation of 
the cDNA insert relative to the flanking T7 and SP6 RNA poly-
merase promoters was verified by restriction analysis. Lin-
erization of this plasmid upstream of the PSM insert fol-

lowed by transcription with SP6 RNA polymerase yields a
400 nucleotide antisense RNA probe, of which 350 nucleotides should be protected from RNase digestion by PSm RNA. This probe was used in Fig. 20. Plasmid IN-20, containing a 1 kb partial PSM cDNA in the plasmid pCR II (Invitrogen) was also used for riboprobe synthesis. IN-20 linearized with Xmn I (Gibco-BRL) yields a 298 nucleotide anti-sense RNA probe when transcribed using SP6 RNA polymerase, of which 260 nucleotides should be protected from RNase digestion by PSm mRNA. This probe was used in Figs. 21 and 22. Probes were synthesized using SP6 RNA polymerase (Gibco-BRL), rNTNT (Gibco-BRL), RNAse (Promega), and 32P-CTP (NEN, Wilmington, Del.) according to published protocols (13). Probes were purified over NENSORB purification columns (NEN) and approximately 1 million cpm of purified, radiolabeled probe was mixed with 10 g of each RNA and hybridized overnight at 45°C using buffers and reagents from the RPA II kit (Ambion, Austin, Tex.). Samples were processed as per manufacturer’s instructions and analyzed on 5% polyacrylamide/7M urea denaturing gels using Sequ ACRYL reagents (ISS, Natick, Mass.). Gels were pre-heated to 55°C and run for approximately 1-2 hours at 25 watts. Gels were then fixed for 30 minutes in 10% methanol/10% acetic acid, dried onto Whatman MM paper at 80°C in a BioRad vacuum dryer and autoradiographed overnight with Hyperfilm MP (Amer sham). Quantitation of PSm expression was determined by using a scanning laser densitometer (LKB, Piscataway, N.J.).

Steroid Modulation Experiment

LNCaP cells (2 million) were plated onto T-75 flasks in RPMI 1640 media supplemented with 5% fetal calf serum and grown 24 hours until approximately 30-40% confluent. Flasks were then washed several times with phosphate-buffered saline and RPMI medium supplemented with 5% charcoal-extracted serum was added. Cells were then grown for another 24 hours, at which time dihydrotestosterone, testosterone, estradiol, progesterone, and dexamethasone (Steraloids Inc., Wilton, N.H.) were added to a final concentration of 2 nM. Cells were grown for 24 hours and RNA was then harvested as previously described and PSm expression analyzed by ribonuclease protection analysis.

Experimental Results

Immunohistochemical Detection of PSm:

Using the 7E11-C5.3 anti-PSm monospec antibody, PSm expression is clearly detectable in the LNCaP prostate cancer cell line, but not in the PC-3 and DU-145 cell lines (Fig. 17) in agreement with previously published results (4A). All normal and malignant prostatic tissues analyzed stained positively for PSm expression (unpublished data).

In-Vivo Transcription/Translation of PSm Antigen:

As shown in Fig. 18, coupled in-vitro transcription/translation of the 2.65 kb full-length PSm cDNA yields an 84 kDa protein species in agreement with the expected protein product from the 750 amino acid PSm open reading frame. Following post-translational modification using pancreatic canine microsomes we obtained a 100 kDa glycosylated protein species consistent with the mature, native PSm antigen. Detection of PSm Antigen in LNCaP Cell Membranes and Transfected PC-3 Cells:

PC-3 cells transfected with the full length PSm cDNA in the pREP7 expression vector were assayed for expression of SM mRNA by Northern analysis (data not shown). A clone with high PSm mRNA expression was selected for PSm antigen analysis by Western blotting using the 7E11-C5.3 antibody. In Fig. 19, the 100 kDa PSm antigen is well expressed in LNCaP cell lysate and membrane fractions, as well as in PSm-transfected PC-3 cells but not in native PC-3 cells. This detectable expression in the transfected PC-3 cells proves that the previously cloned 2.65 kb PSm cDNA encodes the antigen recognized by the 7E11-C5.3 anti-prostate monoclonal antibody and that the antigen is being appropriately glycosylated in the PC-3 cells, since the antibody recognizes a carbohydrate-containing epitope on PSm.

PSm mRNA Expressions

Expression of PSm mRNA in normal human tissues was analyzed using ribonuclease protection assays. Tissue expression of PSm appears predominantly within the prostate, with very low levels of expression detectable in human brain and salivary gland (Fig. 20). No detectable PSm mRNA expression was evident in non-prostatic human tissues when analyzed by Northern analysis (data not shown). We have also noted on occasion detectable PSm expression in normal human small intestine tissue, however this mRNA expression is variable depending upon the specific riboprobe used (data not shown). All samples of normal human prostate and human prostatic adenocarcinoma assayed have revealed clearly detectable PSm expression, whereas we have noted generally decreased or absent expression of PSm in tissues exhibiting benign hyperplasia (Fig. 21). In human LNCaP tumors grown both orthotopically and subcutaneously in nude mice we detected abundant PSm expression with or without the use of matrigel, which is required for the growth of subcutaneously implanted LNCaP cells (Fig. 21). PSm mRNA expression is distinctly modulated by the presence of steroids in physiologic doses (Fig. 22). DHT downregulated expression by 8-10 fold after 24 hours and testosterone diminished PSm expression by 3-4 fold. Estradiol and progesterone also downregulated PSm expression in LNCaP cells, perhaps as a result of binding to the mutated androgen receptor known to exist in the LNCaP cell. Overall, PSm expression is highest in the untreated LNCaP cells grown in steroid-depleted media, a situation that we propose simulates the hormone-deprived (castrate) state in vivo. This experiment was repeated at steroid dosages ranging from 2-200 nM and at time points from 6 hours to 7-days with similar results; maximal downregulation of PSm mRNA was seen with DHT at 24 hours at doses of 2-20 nM.

Experimental Discussion

In order to better understand the biology of the human prostate in both normal and neoplastic states, we need to enhance our knowledge by studying the various proteins and other features that are unique to this important gland. Previous research has provided two valuable prostatic biomarkers, PAP and PSA, both of which have had a significant impact on the diagnosis, treatment, and management of prostate malignancies. Our present work describing the preliminary characterization of the prostate-specific membrane antigen (PSm) reveals it to be a gene with many interesting features. PSM is almost entirely prostate-specific as are PAP and PSA, and as such may enable further delineation of the unique functions and behavior of the prostate. The predicted sequence of the PSM protein (3) and its presence in the LNCaP cell membrane as determined by Western blotting and immunohistochemistry, indicate that it is an integral membrane protein. Thus, PSM provides an attractive cell surface epitope for antibody-directed diagnostic imaging and cytotoxic targeting modalities (14). The ability to synthesize the PSM antigen in vitro and to produce tumor xenografts maintaining high levels of PSM expression provides us with a convenient and attractive model system to further study and characterize the regulation and modulation of PSM expression. Also, the high level of PSM expression in the LNCaP cells provides an excellent in-vitro model system. Since PSM expression is
hormonally-responsive to steroids and may be highly expressed in hormone-refractory disease (15), it is imperative to elucidate the potential role of PSM in the evolution of androgen-independent prostate cancer. The detection of PSM mRNA expression in minute quantities in brain, salivary gland, and small intestine warrants further investigation, although these tissues were negative for expression of PSM antigen by immunohistochemistry using the 7E11-C5.3 antibody (16). In all of these tissues, particularly small-intestine, we detected mRNA expression using a probe corresponding to a region of the PSM cDNA near the 3' end, whereas we were unable to detect expression when using a 5' end PSM probe. These results may indicate that the PSM mRNA transcript undergoes alternative splicing in different tissues. Previous protein studies have suggested that the 7E11-C5.3 antibody may actually detect two other slightly larger protein species in addition to the 5Da PSM antigen (17). These other protein species can be seen in the LNCap lysate and membrane samples in Fig. 19. Possible origins of these proteins include alternatively spliced PSM mRNA, other genes distinct from but closely related to PSM, or different post-translational modifications of the PSM protein. We are currently investigating these possibilities.

Applicant’s approach is based on prostate tissue specific promoter/enzyme or cytokine chimeras. We will examine promoter specific activation of prodrugs such as non toxic gancyclovir which is converted to a toxic metabolite by herpes simplex thymidine kinase or the prodrug 4-(bis(2chloroethyl)amino)benzoyl-1-glutamic acid to the benzoic acid mustard alkylating agent by the pseudomomas carboxypeptidase G2. As these drugs are activated by the enzyme (chimer) specifically in the tumor the active drug is released only locally in the tumor environment, destroying the surrounding tumor cells. We will also examine the promoter specific activation of cytokines such as IL-2, IL-2 or GM-CSF for activation and specific antitumor vaccination. Lastly the tissue specific promoter activation of cellular death genes may also prove to be useful in this area.

Gene Therapy Chimeras

The establishment of “chimeric DNA” for gene therapy requires the joining of different segments of DNA together to make a new DNA that has characteristics of both precursor DNA species involved in the linkage. In this proposal the two pieces being linked involve different functional aspects of DNA, the promoter region which allows for the reading of the DNA for the formation of mRNA will provide specificity and the DNA sequence coding for the mRNA will provide for therapeutic functional DNA.

DNA-Specific Enzyme or Cytokine mRNA:

When effective, antitumor drug activity is the requirement to achieve both a long enough time (t) and high enough concentration (c) (oct) of exposure of the tumor to the toxic drug to assure sufficient cell damage for cell death to occur. The drug also must be “active” and the toxicity for the tumor greater than for the hosts normal cells (22). The availability of the drug to the tumor depends on tumor blood flow and the drugs diffusion ability. Blood flow to the tumor does not provide for selectivity as blood flow to many normal tissues is often as great or greater than that to the tumor. The majority of chemotherapeutic cytotoxic drugs are often toxic to normal tissues as to tumor tissue. Dividing cells are often more sensitive than non-dividing normal cells, but in many slow growing solid tumors such as prostatic cancer this does not provide for antitumor specificity (22).

Previously a means to increase tumor specificity of antitumor drugs was to utilize tumor associated enzymes to activate nontoxic prodrugs to cytotoxic agents. (19). A problem with this approach was that most of the enzymes found in tumors were not totally specific in their activity and similar substrate active enzymes or the same enzyme at only slightly lower amounts was found in other tissue and thus normal tissues were still at risk for damage.

To provide absolute specificity and unique activity, viral, bacterial and fungal enzymes which have unique specificity for selected prodrugs were found which were not present in human or other animal cells. Attempts to utilize enzymes such as herpes simplex thymidine kinase, bacterial cytosome deaminase and carboxypeptidase G-2 were linked to antibody targeting systems with modest success (19). Unfortunately, antibody targeted enzymes limit the number of enzymes available per cell. Also, most antibodies do not have a high tumor target to normal tissue ratio thus normal tissues are still exposed reducing the specificity of these unique enzymes.

Antibodies are large molecules that have poor diffusion properties and the addition of the enzymes molecular weight further reduces the antibodies diffusion.

Gene therapy could produce the best desired result if it could achieve the specific expression of a protein in the tumor and not normal tissue in order that a high local concentration of the enzyme be available for the production in the tumor environment of active drug (21).

Cytokines:

Applicant’s research group has demonstrated that Applicant’s can specifically and non-toxically “cure” an animal of an established tumor, in models of bladder or prostate cancer. The prostate cancer was the more difficult to cure especially if it was grown orthotopically in the prostate.

Our work demonstrated that tumors such as the bladder and prostate were not immunogenic, that is the administration of irradiated tumor cells to the animal prior to subsequent administration of non-irradiated tumor cells did not result in a reduction of either the number of tumor cells to produce a tumor nor did it reduce the growth rate of the tumor. But if the tumor was transfected with a retrovirus and secreted large concentrations of cytokines such as IL-2 then this could act as an antitumor vaccine and could also reduce the growth potential of an already established and growing tumor. IL-2 was the best, GM-CSF also had activity whereas a number of other cytokines were much less active. In clinical studies just using IL-2 for immunostimulation, very large concentrations had to be given which proved to be toxic. The key to the success of the cytokine gene modified tumor cell is that the cytokine is produced at the tumor site locally and is not toxic and that it stimulates immune recognition of the tumor and allows specific and non toxic recognition and destruction of the tumor.

The exact mechanisms of how IL-2 production by the tumor cell activates immune recognition is not fully understood, but one explanation is that it bypasses the need for cytokine production by helper T cells and directly stimulates tumor antigen activated cytotoxic CD8 cells. Activation of antigen presenting cells may also occur.

Tissue Promotor-Specific Chimera DNA Activation Non-Prostatic Tumor Systems:

It has been observed in non-prostatic tumors that the use of promotor specific activation can selectively lead to tissue specific gene expression of the transfected gene. In melanoma the use of the tyrosinase promotor which codes for the enzyme responsible for melanin expression produced over a 50 fold greater expression of the promotor driven reporter gene expression in melanoma cells and not non melanoma cells. Similar specific activation was seen in the melanoma
cells transfected when they were growing in mice. In that experiment no non-melanoma or melanocyte cell expressed the tyrosinase drive reporter gene product. The research group at Welcome Laboratories have cloned and sequenced the promoter region of the gene encoding for carciinoembryonic antigen (CEA). CEA is expressed on colon and colon carcinoma cells but specifically on metastatic cytotoxic deaminase which converts 5-fluorouracil into 5-fluorouracil and observed a large increase in the ability to selectively kill CEA promoter driven colon tumor cells but not dividing not dividing normal liver cells. In vivo they observed that bystander tumor cells which were not transfected with the cytotoxic deaminase gene were also killed, and that there was no toxicity to the host animal as the large tumors were regressing following treatment. Herpes simplex virus, (HSV), thymidine kinase similarly activates the prodruk gancyclovir to be toxic towards dividing cancer cells and HSV thymidine kinase has been shown to be specifically activatable by tissue specific promoters.

Prostatic Tumor Systems:
The therapeutic key to effective cancer therapy is to achieve specificity and spare the patient toxicity. Gene therapy may provide a key to specificity in that non-essential tissues such as the prostate and prostatic tumors produce tissue specific proteins, such as acid phosphatase (PAP), prostate specific antigen (PSA), and a gene which we cloned, prostate-specific membrane antigen (PSM). Tissues such as the prostate contain selected tissue specific transcription factors which are responsible for binding to the promoter region of the DNA of these tissue specific mRNA. The promoter for PSA has been cloned and we are investigating its use as a prostatic specific promoter for prostatic tumor cells. Usually patients who are being treated for metastatic prostatic cancer have been put on androgen deprivation therapy which dramatically reduces the expression of mRNA for PSA. PSM on the other hand increases in expression with hormone deprivation which means it would be even more intensely expressed on patients being treated with hormone therapy. Preliminary work in collaboration with Dr. John Isaacs’ Laboratory demonstrates that PSM is expressed when the human chromosome region containing the human PSM gene is transferred to the rat tumor AT-6. AL-6 is a metastatic androgen independent tumor. The same chromosome transferred into non prostate derived tissues or tumors is not expressed and thus these cells could be used as an animal model for these experiments. PSA, PSM positive HUANLNCAP cells will be used for testing in nude mice.

REFERENCES OF THE SECOND SERIES OF EXPERIMENTS

14. Personal Communication from Cytogen Corporation, Princeton, NJ.


THIRD SERIES OF EXPERIMENTS

Sensitive Detection of Prostatic Hematogenous Micrometastases Using PSA and PSMA-Derived Primers in the Polymerase Chain Reaction

We have developed a PCR-based assay enabling sensitive detection of hematogenous micrometastases in patients with prostate cancer. We performed "nested PCR", amplifying mRNA sequences unique to prostate-specific antigen and to the prostate-specific membrane antigen, and have compared their respective results. Micrometastases were detected in 52 patients (6.7%) by PCR with PSA-derived primers, while PSMA-derived primers detected tumor cells in 19/16 patients (63.3%). All 8 negative controls were negative with both PSA and PSMA PCR. Assays were repeated to confirm results, and PCR products were verified by DNA sequencing and Southern analysis. Patients harboring circulating prostatic tumor cells as detected by PCR, and not by PSA-PCR included 8 patients previously treated with radical prostatectomy and with non-measurable serum PSA levels at the time of this assay. The significance of these findings with respect to future disease recurrence and progression will be investigated.

Improvement in the overall survival of patients with prostate cancer will depend upon earlier diagnosis. Localized disease, without evidence of extra-prostatic spread, is successfully treated with either radical prostatectomy or external beam radiation, with excellent long-term results (2.3). The major problem is that approximately two-thirds of men diagnosed with prostate cancer already have evidence of advanced extra-prostatic spread at the time of diagnosis, for which there is at present no cure (4). The use of clinical serum markers such as prostate-specific antigen (PSA) and prostatic acid phosphatase (PAP) have enabled clinicians to detect prostatic carcinomas earlier and provide useful parameters to follow responses to therapy (5). Yet, despite the advent of sensitive serum PSA assays, radionuclide bone scans, CT scans and other imaging modalities, we are still unable to detect the presence of micrometastatic cells prior to their establishment of solid metastases. Previous work has been done utilizing the polymerase chain reaction to amplify mRNA sequences unique to breast, leukemia, and other malignant cells in the circulation and enable early detection of micrometastases (6,7). Recently, a PCR-based approach utilizing primers derived from the PSA DNA sequence was published (8). In this study 3/2 patients with advanced, stage D prostate cancer had detectable hematogenous micrometastases.

We have recently identified and cloned a 2.65 kb cDNA encoding the 100 kDa prostate-specific membrane antigen (PSM) recognized by the anti-prostate monoclonal antibody 7E11-C3.3 (9). PSM appears to be an integral membrane glycoprotein which is highly expressed in prostatic tumors and metastases and is almost entirely prostate-specific (10). Many anaplastic tumors and bone metastases have variable and at times no detectable expression of PSA, whereas these lesions appear to consistently express high levels of PSM. Prostatic tumor cells that escape from the prostate gland and enter the circulation are likely to have the potential to form metastases and are possibly the more aggressive and possibly anaplastic cells, a population of cells that may not express high levels of PSA, but may retain high expression of PSM. We therefore chose to utilize DNA primers derived from the sequences of both PSA and PSM in a PCR assay to detect micrometastatic cells in the peripheral circulation.

Despite the high level of amplification and sensitivity of conventional RNA PCR, we have utilized a "nested" PCR approach in which we first amplify a target sequence, and subsequently use this PCR product as the template for another round of PCR amplification with a new set of primers totally contained within the sequence of the previous product. This approach has enabled us to increase our level of detection from one prostatic tumor cell per 10,000 cells to better than one cell per ten million cells.

EXPERIMENTAL DETAILS

Materials and Methods

Cells and Reagents:

LNCaP and MCF-7 cells were obtained from the American Type Culture Collection (Rockville, Md.). Details regarding the establishment and characteristics of these cell lines have been previously published (11,12). Cells were grown in RPMI 1640 media supplemented with L-glutamine, nonessential amino acids, obtained from the MSKCC Media Preparation Facility, and 5% fetal calf serum (Gibco-BRL, Gaithersburg, Md.) in a CO2 incubator at 37C. All cell media was obtained from the MSKCC Media Preparation Facility. Routine chemical reagents were of the highest grade possible and were obtained from Sigma Chemical Company, St. Louis, Mo.

Patient Blood Specimens

All blood specimens used in this study were from patients seen in the outpatient offices of urologists on staff at MSKCC. Two anti-coagulated (purple top) tubes per patient were obtained at the time of their regularly scheduled blood draws. Specimen procurement was conducted as per the approval of the MSKCC Institutional Review Board. Samples were promptly brought to the laboratory for immediate processing. Serum PSA and PAP determinations were performed by standard techniques by the MSKCC Clinical Chemistry Laboratory. PSA determinations were performed using the Tandem PSA assay (Hybritech, San Diego, Calif.). The eight blood specimens used as negative controls were from 2 males with normal serum PSA values and biopsy-proven BPH, one healthy female, 3 healthy males, one patient with bladder cancer, and one patient with acute promyelocytic leukemia.

Blood Sample Processing/RNA Extraction

4 ml of whole anti-coagulated venous blood was mixed with 3 ml of ice cold phosphate buffered saline and then carefully layered atop 8 ml of Ficoll (Pharmacia, Uppsala, Sweden) in a 15-ml polyvynene tube. Tubes were centrifuged at 200 x g for 30 min. at 4C. Using a sterile Pasteur pipette, the buffy coat layer (approx. 1 ml) was carefully removed and rediluted up to 50 ml with ice cold phosphate buffered saline in a 50 ml polypolyene tube. This tube was then centrifuged at 2000 x g for 30 min at 4C. The supernatant was carefully decanted and the pellet was allowed to drip dry. One ml of RNAzol B was then added to the pellet and total RNA was isolated as per manufacturers directions (Cinna/Biotech, Houston, Tex.). RNA concentrations and purity were determined by UV spectroscopy on a Beckman DU 640 spectrophotometer and by gelanalyzer.
Determination of PCR Sensitivity

RNA was isolated from LNCaP cells and from mixtures of LNCaP and MCF-7 cells at fixed ratios (i.e. 1:100, 1:1000, etc.) using RNAzol B. Nested PCR was then performed as described below with both PSA and PSM primers in order to determine the limit of detection for the assay. LNCaP-MCF-7 (1:100,000) cDNA was diluted with distilled water to obtain concentrations of 1:1,000,000 and 1:10,000,000. MCF-7 cells were chosen because they had been previously tested and shown not to express PSM by PCR.

Polymerase Chain Reaction

The PSA outer primers used span portions of exons 4 and 5 to yield a 486 bp PCR product and enable differentiation between cDNA and possible contaminating genomic DNA amplification. The upstream primer sequence beginning at nucleotide 494 in PSA cDNA sequence is 5'-TACCCTCGCATCAGGAACA-3' (SEQ. ID. No. 39) and the downstream primer at nucleotide 960 is 5'-CTTGAAGCACACTTACA-3' (SEQ. ID. No. 40). The PSA inner upstream primer (beginning at nucleotide 550) 5'-ACACCGCAGGTTTACATCAG-3' (SEQ. ID. No. 41) and the downstream primer (at nucleotide 894) 5'-GTCGACGTTCGACACAG-3' (SEQ. ID. No. 42) yield a 355 bp PCR product. All primers were synthesized by the MSCKC Microchemistry Core Facility. 5 µl of total RNA was reverse-transcribed into cDNA in a total volume of 201 using Superscript reverse transcriptase (Gibco-BRL) according to the manufacturers recommendations. 11 µl of this cDNA served as the starting template for the outer primer PCR reaction. The 201 PCR mix included: 0.5U Taq polymerase (Promega Corp., Madison, Wis.), Promega reaction buffer, 1.5MM MgCl2, 200M dNTPs, and 1.0M of each primer. This mix was then transferred to a Perkin Elmer 9600 DNA thermal cycler and incubated for 25 cycles. The PCR profile was as follows: 94°C for 15 sec., 60°C for 15 sec., and 72°C for 45 sec. After 25 cycles, samples were placed on ice, and 11 of this reaction mix served as the template for another round of PCR using the inner primers. The first set of tubes were returned to the thermal cycler for 25 additional cycles. PSM-PCR required the selection of primer pairs that also spanned an intron in order to be certain that cDNA and not genomic DNA were being amplified. Since the genomic DNA sequence of PSM has not yet been determined, this involved trying different primer pairs until a pair was found that produced the expected size PCR product when cDNA was amplified, but with no band produced from a genomic DNA template, indicating the presence of a large intron. The PSM outer primers yielded a 946 bp product and the inner primers a 434 bp product. The PSM outer primer upstream used was 5'-ATGGGTTTGGTTACATTC-3' (SEQ. ID. No. 43) (beginning at nucleotide 1401) and the downstream primer (at nucleotide 2348) was 5'-GCTTGGAGCGCATGATCATGC-3' (SEQ. ID. No. 44). The PSM inner upstream primer (at nucleotide 1581) was 5'-ACTCCTCTAAGACGGTGCGG-3' (SEQ. ID. No. 45) and the downstream primer (at nucleotide 2015) was 5'-AACATTATCTCAGGAAAG-3' (SEQ. ID. No. 46). cDNA used was the same as for the PSA assay. The 501 PCR mix included: 1U Tag Polymerase (Promega), 250M dNTPs, 10M mercaptoethanol, 2 mM MgCl2, and 51 of a 10X buffer mix containing: 160mM NH4SO4, 670mM Tris pH 8.8, and 2 mg/ml of nucleylated BSA. PCR was carried out in a Perkin Elmer 4800 DNA thermal cycler with the following parameters: 94°C for 4 minutes for 1 cycle, 94°C for 30 sec., 58°C for 1 minute, and 72°C for 1 minute for 25 cycles, followed-by 72°C 10 minutes. Samples were then cycled and 21 of this reaction mix used as the template for another 25 cycles with new reaction mix containing the inner PSM primers. cDNA quality was verified by performing control reactions using primers derived from -actin yielding a 446 bp PCR product. The upstream primer used was 5'-AGGCCAAGCCGAGA-3' (SEQ. ID. No. 47) (exon 3) and the downstream primer was 5'-ATGTCCATGGAGAAG-3' (SEQ. ID. No. 48) (exon 4). The entire PSA mix and 101 of each PSM reaction mix were run on 1.5-2% agarose gels, stained with ethidium bromide and photographed in an Eagle Eye Video Imaging System (Stratagene, Torrey Pines, Calif.). Assays were repeated at least 3 times to verify results.

Cloning and Sequencing of PCR Products

PCR products were cloned into the pCR II plasmid vector using the TA cloning system (Invitrogen). These plasmids were transformed into competent E. coli cells using standard methods (13) and plasmid DNA was isolated using Magic Miniprep (Promega) and screened by restriction analysis. TA clones were then sequenced by theideoxy method (14) using Sequenase (U.S. Biochemical). 3-4g of each plasmid was denatured with NaOH and ethanol precipitated. Labeling reactions were carried out according to the manufacturers recommendations using [35S]-ATP (NEB), and the reactions were terminated as discussed in the same protocol. Sequencing products were then analyzed on 6% polyacrylamide/7M urea gels run at 120 watts for 2 hours. Gels were fixed for 20 minutes in 10% methanol/10% acetic acid, transferred to Whatman 3MM paper and dried down in a vacuum dryer for 2 hours at 80°C. Gels were then autoradiographed at room temperature for 18 hours.

Southern Analysis

Ethidium-stained agarose gels of PCR products were soaked for 15 minutes in 0.2N HCL followed by 30 minutes each in 0.5N NaOH/1.5M NaCl and 0.1M Tris pH 7.5/1.5M NaCl. Gels were then equilibrated for 10 minutes in 10× SSC (1.5M NaCl/0.15M Sodium Citrate. DNA was transferred onto Nitran nylon membranes (Schleicher and Schuell) by pressure blotting in 10× SSC with a Posi-blottter (Stratagene). DNA was cross-linked to the membrane using a UV Stratalinker (Stratagene). Blots were pre-hybridized at 65°C for 2 hours and subsequently hybridized with denatured 32P-labeled, random-primed cDNA probes (either PSA or PSM) (9,15). Blots were washed twice in 1x SSPE/0.5% SDS at 42°C and twice in 0.1x SSPE/0.5% SDS at 50°C for 20 minutes each. Membranes were air-dried and autoradiographed for 30 minutes to 1 hour at ~70C with Kodak X-Omat film.

Experimental Results

Our technique of PCR amplification with nested primers improved our level of detection of prostatic cells from approximately one prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using either PSA or PSM-derived primers (FIGS. 26 and 27). This represents a substantial improvement in our ability to detect minimal disease. Characteristics of the 16 patients analyzed with respect to their clinical stage, treatment, serum PSA and PAP values, and results of our assay are shown in table 1. In total, PSA-PCR detected tumor cells in 2/30 patients (6.7%), whereas PSM-PCR detected cells in 19/30 patients (63.3%). There were no patients positive for tumor cells by PSA and not by PSM, while PSM provided 8 positive patients not detected by PSA. Patients 10 and 11 in table 1, both with very advanced hormone-refractory disease were detected by both PSA and PSM. Both of these patients have died since the time these samples were obtained. Patients 4, 7, and 12, all of whom were treated with radical prostatectomies for clinically localized disease, and all of whom have non-measurable serum PSA values 1-2 years postoperatively were positive for circulating prostatic tumor cells by PSA-PCR, but negative by PSM-PCR. A representative ethidium stained gel photograph
for PSM-PCR is shown in FIG. 28. Samples run in lane A represent PCR products generated from the outer primers and samples in lanes labeled B are products of inner primer pairs. The corresponding PSM Southern blot autoradiograph is shown in FIG. 29. The sensitivity of the Southern blot analysis exceeded that of ethidium staining, as can be seen in several samples where the outer product is not visible on FIG. 28, but is detectable by Southern blotting as shown in FIG. 29. In addition, sample 5 on FIGS. 28 and 29 (patient 6 in FIG. 30) appears to contain both outer and inner bands that are smaller than the corresponding bands in the other patients. DNA sequencing has confirmed that the nucleotide sequence of these bands matches that of PSM, with the exception of a small deletion. This may represent either an artifact of PCR, alternative splicing of PSM mRNA in this patient, or a PSM mutation. We have noted similar findings with other samples on several occasions (unpublished data). All samples sequenced and analyzed by Southern analysis have been confirmed as true positives for PSA and PSM.

Experimental Details

The ability to accurately stage patients with prostate cancer at the time of diagnosis is clearly of paramount importance in selecting appropriate therapy and in predicting long-term response to treatment, and potential cure. Pre-surgical staging presently consists of physical examination, serum PSA and PAP determinations, and numerous imaging modalities including transrectal ultrasoundography, CT scanning, radionuclide bone scans, and even MRI scanning. No present modality, however, addresses the issue of hematogenous micrometastatic disease and the potential negative impact on prognosis that this may produce. Previous work has shown that only a fractional percentage of circulating tumor cells will inevitably go on to form a solid metastasis (16), however, the detection of and potential quantification of circulating tumor cell burden may prove valuable in more accurately staging disease. The long-term impact of hematogenous micrometastatic disease must be studied by comparing the clinical courses of patients found to have these cells in their circulation with patients of similar stage and treatment who test negatively.

The significantly higher level of detection of tumor cells with PSM as compared to PSA is not surprising to us, since we have noted more consistent expression of PSM in prostate carcinomas of all stages and grades as compared to variable expression of PSA in more poorly differentiated and anaplastic prostate cancers. We were surprised to detect tumor cells in the three patients that had undergone radical prostatectomies with subsequent undetectable amounts of serum PSA. These patients would be considered to be surgical “cures” by standard criteria, yet they apparently continue to harbor prostatic tumor cells. It will be interesting to follow the clinical course of these patients as compared to others without PCR evidence of residual disease. We are presently analyzing larger numbers of patient samples in order to verify these findings and perhaps identify patients at risk for metastatic disease.

REFERENCES


FOURTH SERIES OF EXPERIMENTS

EXPRESSION OF THE PROSTATE SPECIFIC MEMBRANE ANTIGEN (PSM) DIMINISHES THE MITOGENIC STIMULATION OF AGGRESSIVE HUMAN PROSTATIC CARCINOMA CELLS BY TRANSFERRIN

An association between transferrin and human prostate cancer has been suggested by several investigators. It has also been observed that the expressed prostatic secretions of patients with prostate cancer are enriched with respect to the content of transferrin and that prostate cells are rich in transferrin receptors (J. Urol. 143, 381, 1990). Transferrin derived from bone marrow has been shown to selectively stimulate the growth of aggressive prostate cancer cells (PNAS 89, 6197, 1992). We have previously reported the cloning of the cDNA encoding the 100 kDa PSM antigen (Cancer Res. 53, 208, 1993). DNA sequence analysis has revealed that a portion of the coding region, from nucleotide 1250 to 1700 possesses a 54% homology to the human transferrin receptor. PC-3 cells do not express PSM mRNA or protein and exhibit...
increased cell growth in response to transferrin, whereas LNCaP prostate cancer cells which highly express PSM have a very weak response to transferrin. To determine whether PSM expression by prostatic cancer cells impacts upon their mitogenic response to transferrin we stably transfected the full-length PSM cDNA into the PC-3 prostate cancer cells. Clones highly expressing PSM mRNA were identified by Northern analysis and expression of PSM protein was verified by Western analysis using the anti-PSM monoclonal antibody 7E11-C5.3.

We plated 2x10^6 PC-3 or PSM-transfected PC-3 cells per well in RPMI medium supplemented with 10% fetal bovine serum and at 24 hrs. added 1 μg per ml of holotransferrin to the cells. Cells were counted 1 day to be highly mitogenic to the PC-3 cells. Cells were counted at 1 day to determine plating efficiency and at 5 days to determine the effect of the transferrin. Experiments were repeated to verify the results.

We found that the PC-3 cells experienced an average increase of 275% over controls, whereas the LNCaP cells were only stimulated 43%. Growth kinetics revealed that the PSM-transfected PC-3 cells grew 30% slower than native PC-3 cells. This data suggests that PSM expression in aggressive, metastatic human prostate cancer cells significantly abrogates their mitogenic response to transferrin.

The use of therapeutic vaccines consisting of cytokine-secreting tumor cell preparations for the treatment of established prostate cancer was investigated in the Dunning R3327-MatLyLu rat prostatic adenocarcinoma model. Only IL-2 secreting, irradiated tumor cell preparations were capable of curing animals from subcutaneously established tumors, and engendered immunological memory that protected the animals from another tumor challenge. Immunotherapy was less effective when tumors were induced orthotopically, but nevertheless led to improved outcome, significantly delaying, and occasionally preventing recurrence of tumors after resection of the cancellous prostate. Induction of a potent immune response in tumor bearing animals against the nonimmunogenic MatLyLu tumor supports the view that active immunotherapy of prostate cancer may have therapeutic benefits.

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| 125 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
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Glu Pro Pro Pro Pro Gly Tyr Glu Asn Val Ser Asp Ile Val Pro Pro 145 150 155 160
Phe Ser Ala Phe Ser Pro Glu Gly Met Pro Glu Gly Asp Leu Val Tyr 165 170 175
Val Asn Tyr Ala Arg Thr Glu Asp Phe Phe Lys Leu Glu Arg Asp Met 180 185 190
Lys Ile Asn Cys Ser Gly Lys Ile Val Ile Ala Arg Tyr Gly Lys Val 195 200 205
Phe Arg Gly Asn Lys Val Lys Asn Ala Glu Leu Ala Gly Ala Lys Gly 210 215 220
Val Ile Leu Tyr Ser Asp Pro Ala Asp Tyr Phe Ala Pro Gly Val Lys 225 230 235 240
Ser Tyr Pro Asp Gly Thr Asp Leu Pro Gly Gly Val Gin Arg Gly 245 250 255
Asn Ile Leu Asn Leu Gly Ala Gly Asp Pro Leu Thr Pro Gly Tyr 260 265 270
Pro Ala Asn Glu Tyr Ala Tyr Arg Arg Gly Ile Ala Glu Ala Val Gly 275 280 285
Leu Pro Ser Ile Pro Val His Pro Ile Gly Tyr Tyr Asp Ala Gin Lys 290 295 300
Leu Leu Glu Lys Met Gly Gly Ser Ala Pro Asp Ser Ser Trp Arg 305 310 315 320
Gly Ser Leu Lys Val Pro Tyr Asn Val Gly Pro Gly Phe Thr Gly Asn 325 330 335
Phe Ser Thr Gin Lys Val Lys Met His Ile His Ser Thr Asn Glu Val 340 345 350
Thr Arg Ile Tyr Asn Val Ile Gly Thr Leu Arg Gly Ala Val Glu Pro 355 360 365
Asp Arg Tyr Val Ile Leu Gly Gly His Arg Asp Ser Trp Val Phe Gly 370 375
Gly Ile Asp Pro Gin Ser Gly Ala Ala Val Val His Gly Ile Val Arg 385 390 395 400
Ser Phe Gly Thr Leu Lys Lys Gly Thr Arg Pro Arg Arg Thr Ile 405 410 415
Leu Phe Ala Ser Trp Asp Ala Glu Phe Gly Leu Leu Gly Ser Thr 420 425 430
Glu Trp Ala Glu Glu Asn Ser Arg Leu Leu Gin Arg Gly Val Ala 435 440 445
Tyr Ile Asn Ala Asp Ser Ile Glu Gly Asn Tyr Thr Leu Arg Val 450 455 460
Asp Cys Thr Pro Leu Met Tyr Ser Leu Val His Asn Leu Thr Lys Glu 465 470 475 480
Leu Lys Ser Pro Asp Glu Gly Phe Glu Gly Lys Ser Leu Tyr Glu Ser 485 490 495
Trp Thr Lys Lys Ser Pro Ser Pro Glu Phe Ser Gly Met Pro Arg Ile 500 505 510
Ser Lys Leu Gly Ser Gly Asn Arg Phe Glu Val Phe Phe Gin Arg Leu 515 520 525
Lys Ile Ala Ser Gly Arg Ala Arg Tyr Thr Lys Asn Trp Glu Thr Asn 530 535 540
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Leu Val Glu Lys Phe Tyr Asp Pro Met Phe Lys Tyr His Leu Thr Val 565 570 575
Ala Glu Val Arg Gly Gly Met Val Phe Glu Leu Ala Asn Ser Ile Val 580 585 590
Leu Pro Phe Asp Cys Arg Asp Tyr Ala Val Val Leu Arg Lys Tyr Ala 595 600 605
Asp Lys Ile Tyr Ser Ile Ser Met Lys His Pro Glu Glu Met Lys Thr 610 615 620
Tyr Ser Val Ser Phe Asp Ser Leu Phe Ser Ala Val Lys Asn Phe Thr 625 630 635 640
Glu Ile Ala Ser Lys Phe Ser Glu Arg Leu Gln Asp Phe Asp Lys Ser 645 650 655
Asn Pro Ile Val Leu Arg Met Met Asn Asp Glu Leu Met Phe Leu Glu 660 665 670
Arg Ala Phe Ile Asp Pro Leu Gly Leu Pro Asp Arg Pro Phe Tyr Arg 675 680 685
His Val Ile Tyr Ala Pro Ser Ser His Asn Lys Tyr Ala Gly Glu Ser 690 695 700
Phe Pro Gly Ile Tyr Asp Ala Leu Phe Asp Ile Glu Ser Lys Val Asp 705 710 715 720
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tgtgtgtcgc accatctctca gcaagtcgcg aggccgcttg ttcagccccaa ttgatggaga 180
cactgctct ganagmcgtt ggaaggtgc gatcanntt tctc gttaaggg tmngacmnaa 240
ccaagcgag gannngccga gantaagtgg gtaacagact tggacaaatt ccatgaaaga 300
caggaagatt ctgaactctt tcgggtctat ccaaggtatc gagaaacctc atcggtatgt 360
tgtgtagtgga gcccagcagag acctctgggg cccaggggctg gtaaagcctg gcaacttgac 420
tgttagatatg ctgaactctt cccctgttct atcagacacta tggaaacagc aggggtcaca 480
accggagga agcagactct tctgctgtctg gatgcgcagga gatcagcggag cttgaggggc 540	actgagaattc tgctctggg accegtccat gctgcatcacg aaagctctcttctgcctcatc 600
ngcttgtagt ctccagctct ggagacagcc catgtcsaga tttctgcgcag cccctttgctg 660
tatagctgaggggatttattgagaggg ctggagactc cacagacagt ctccagagcgc 720
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gaaagaaaca agatatacct acaatcccttg cgyttagaaaa ggatagatgaa acaagaccc 180
tctaattgta ttagagagcc agaagagagc tggggctcct ggtnttctgg aatcoctgtg 240
tggataccct cttctntgtgt aaaaacctgg ccacagattc cagatagtac tcaaaaagat 300
gattagagc ccaacagagc cattatcttt gcaacgtaga cttggaagga cattagagt 360
gttggtcctga ctagtggtct gggaggtgct cttctctctc tgcctctaa mngntctctc 420
actaacatta atnctgtgata aagtgctcct ggtatagac aacctcaagg tttctgcag 480
ccccatatc tatatcacta tgggaagag atgtcaggag nctgtaatgac tcgganmn 540
nnnttgatgg aataatatact tataagaaac gtaatggat tagcaaaatt gagcaacatt 600
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catccttggga ggatataagag gttttcttga acagcaagtc atagttcttg tagggcccaa 180
gagagctgaa tggggttctgg gatagtggaa ttcnncggttg aggcaagct cttctatttg 240
aacctggaca gattgtctct taaaactggtt gttctccacc cagcagaagca 300
ctcttttcgc cagttggaagag ctagtgaggg ctagtgctct cttgcaccact gatgtgctag 360
aggyatacct tcgtenoct gcattaag gcttcacgt aattaactt ggtaaagcg

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1 5

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<210> SEQ ID NO 34
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<210> SEQ ID NO 35
<211> LENGTH: 6
<212> TYPE: PRT
<213> ORGANISM: human
<400> SEQUENCE: 35
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1 5
What is claimed is:

1. A purified monoclonal antibody which binds to a fragment of an outer membrane domain of prostate specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO: 35).

2. A purified monoclonal antibody which binds to a fragment of an outer membrane domain of prostate specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO: 36).

3. A purified monoclonal antibody which binds to a fragment of an outer membrane domain of prostate specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO: 37).

4. A purified monoclonal antibody which binds to a fragment of an outer membrane domain of prostate specific membrane antigen, which fragment has within its structure each of the following amino acid sequences:
   (a) Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO: 35);
   (b) Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO: 36);
   (c) Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO: 37); and

5. A purified monoclonal antibody which binds to a fragment of an outer membrane domain of prostate specific membrane antigen, which fragment has within its structure each of the following amino acid sequences:
   (a) Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO: 35);
   (b) Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO: 36); and
   (c) Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO: 37).

6. A purified monoclonal antibody which binds to a fragment of prostate specific membrane antigen, which fragment corresponds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

7. A purified monoclonal antibody which binds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO: 35).

8. A purified monoclonal antibody which binds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO: 36).

9. A purified monoclonal antibody which binds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO: 37).

10. A purified monoclonal antibody which binds to an outer membrane domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

11. A purified monoclonal antibody which binds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

12. The purified antibody of any one of claims 1-11, wherein the antibody is a monoclonal antibody.

13. A composition of matter comprising [the] a monoclonal antibody [of any one of 1-11] and an agent conjugated [to the monoclonal antibody] thereto, wherein the monoclonal antibody binds to an outer membrane domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

14. The composition of matter of claim 13, wherein the agent is a radiisotope or toxin.

16. A method of imaging prostate cancer in a subject which comprises administering to the subject the composition of matter of claim 13, wherein the agent is an imaging agent under conditions permitting formation of a complex between the composition of matter and prostate specific membrane antigen, and obtaining an image of any complex so formed.

17. A monoclonal antibody having an antigen-binding region -specific for the extracellular domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

18. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a fragment of an outer membrane domain of prostate-specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO:35).

19. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a fragment of an outer membrane domain of prostate-specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO:36).

20. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a fragment of an outer membrane domain of prostate-specific membrane antigen, which fragment has within its structure the consecutive amino acid sequence Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO:37).

21. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a fragment of an outer membrane domain of prostate-specific membrane antigen, which fragment has within its structure each of the following amino acid sequences: (a) Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO:35); (b) Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO:36); and (c) Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO:37).

22. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a fragment of prostate-specific membrane antigen, which fragment corresponds to a hydrophilic region of an outer membrane domain of prostate-specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

23. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a hydrophilic region of an outer membrane domain of prostate-specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID NO:35).

24. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a hydrophilic region of an outer membrane domain of prostate-specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID NO:36).

25. A composition of comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a hydrophilic region of an outer membrane domain of prostate-specific membrane antigen, which hydrophilic region has within its structure the consecutive amino acid sequence Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID NO:37).

26. A composition of matter comprising a monoclonal antibody and an agent conjugated thereto, wherein the monoclonal antibody binds to a hydrophilic region of an outer membrane domain of prostate specific membrane antigen, the amino acid sequence of which antigen is set forth in SEQ ID NO:2.

27. The composition of matter of any of claim 18-20, or 21-26, wherein the agent is a radioisotope or toxin.


29. A method of imaging prostate cancer in a subject which comprises administering to the subject the composition of matter of any of claim 18-20 or 21-26, wherein the agent is an imaging agent, under conditions permitting formation of a complex between the composition of matter and prostate-specific membrane antigen, and obtaining an image of any complex so formed.


31. The composition of claim 30, wherein the agent is a radioisotope.

32. The composition of claim 30, wherein the agent is a toxin.

33. The method of claim 29, wherein the imaging agent is a radioisotope.

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