## **RAPID COMMUNICATION**

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# PSMA expression in the Hi-Myc model; extended utility of a representative model of prostate adenocarcinoma for biological insight and as a drug discovery tool

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Prostate-specific membrane antigen (PSMA), also known as glutamate carboxypeptidase II (GCPII), is highly overexpressed in primary and metastatic prostate cancer (PCa). This has led to the development of radiopharmaceuticals for targeted imaging and therapy under current clinical evaluation. Despite this progress, the exact biological role of the protein in prostate cancer development and progression has not been fully elucidated. This is in part because the human PSMA and mouse PSMA (mPSMA) have different patterns of anatomical expression which confound study in the most widely utilized model organisms. Most notably, mPSMA is not expressed in the healthy murine prostate. Here, we reveal that mPSMA is highly upregulated in the prostate adenocarcinoma of the spontaneous Hi-Myc mouse model, a highly accurate and well characterized mouse model of prostate cancer development. Antibody detection and molecular imaging tools are used to confirm that mPSMA is expressed from early prostatic intraepithelial neoplasia (PIN) through adenocarcinoma.

#### KEYWORDS

drug development, mouse models, prostate cancer, PSMA

### **1** | INTRODUCTION

Targeted imaging and therapeutic delivery to prostate-specific membrane antigen (PSMA) is currently of great biomedical interest. Antibodies, peptides, and small molecules are all under investigation both preclinically and in late stage clinical trial for diagnostic benefit and therapeutic potential. Indeed, at this time there are 127 active or recruiting clinical trials using PSMA targeted agents or cells for

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diagnostic or therapeutic purpose in the clinicaltrials.gov database. PSMA protein is expressed in the kidney, gut, brain, peripheral nerves, salivary glands, and prostate of humans. In malignant tissues, expression can be detected in the neovasculature of multiple cancer types, and is highly upregulated in prostate cancer tissues. Expression in prostate cancer increases with disease stage of and the majority of both soft-tissue and osseous metastases highly express the antigen.<sup>1,2</sup>

Discovered independently in several different tissues including as glutamate carboxypeptidase II (GCPII) in the gut, PSMA in the prostate and N-acetyl-L-aspartyl-L-glutamate peptidase I of NAAG peptidase I (NAALADase I) in the brain, the enzyme encoded by FOLH1 has different functions in different tissues. Within the brain it catabolizes NAAG, the most prevalent peptidic neurotransmitter, to N-acetylaspartate and glutamate; in the intestine it cleaves terminal glutamates from dietary folic-polyglutamates. The biological role of PSMA in the prostate and in cancer is not fully known.

Mice are commonly utilized to more deeply understand the etiological basis and impact of protein expression, as well as to evaluate imaging agents and experimental agents in models of disease, and should be a useful tool to investigate PSMA function. Human and mouse PSMA share 86% sequence identity, with identical enzymatic activity.<sup>3</sup> However, PSMA expression patterns in mice are different than those found in man. Some reports have indicated low<sup>3</sup> or a lack<sup>4</sup> of expression of mouse FOLH1 (Folh1) in the mouse prostate, while others report moderate to high expression in normal prostate and in the TRAMP genetically engineered mouse model (GEMM) of prostate cancer.5-7 Investigation of mPSMA in cancer progression and development have used transgenic mPSMA expressing tumor models<sup>8</sup> and PSMA knockout mice crossed with TRAMP mice<sup>7,9,10</sup> to suggest that PSMA promotes prostate cancer formation and progression. Some of this discrepancy may be due to multiple similar family members that can interfere with specific detection of PSMA. Here we use multiple PSMA specific detection methods, including highsensitivity non-invasive in vivo positron emission tomographic (PET) imaging, to show little or no PSMA expression in normal mouse prostate, but highly upregulated expression in the Hi-Myc model of prostate cancer. This unexpected finding has significant import to improve our understanding of PSMA in disease biology and provides a validated research model for drug evaluation.

## 2 | RESULTS

In order to confirm the ability of antibodies used for immunohistochemistry to specifically detect PSMA expression in mouse tissues, we performed protein detection on tissues known to express PSMA. In kidney, PSMA expression was detected in proximal renal tubules and in Bowman's capsule with the expected membranous staining pattern; representative low (top) and high (bottom) magnification micrographs shown (Figure 1A). Specificity was confirmed by repeating the staining conditions in PSMA (*Folh1*) knockout mice (Figure 1B). PSMA expression was undetectable in all examined tissues from knockout mice, suggesting no apparent cross reactivity with other PSMA family members. Little to no expression was detected in normal prostate (Figure 1D) from wild-type mice.

In human prostate cancer samples (Figure 1C), high levels of PSMA expression was detected using the same antibody and revealed the expected pattern of membranous staining with concentration at the luminal border of epithelial cells. Surprisingly, in Hi-Myc mice (Figures 1E and 1F), PSMA expression was detected in PIN lesions (Figure 1E) and in invasive adenocarcinoma (Figure 1F) with a membranous expression pattern similar to human cancer. Tissue lysates prepared from wild type and Hi-Myc mice were similarly assayed for PSMA expression using the same antibody used for immunohistochemistry (Figure 1G). A doublet band of the expected size (approximately 110 kDa) was detected in kidney, brain, and salivary gland. In prostate tissue, only a very faint band was present in lysates from wild-type mice, but PSMA expression was detected in the anterior and lateral lobes of 6-month-old Hi-Myc mice. Myc expression and cancer burden in the Hi-Myc mice is not equal in all prostate lobes, but has been reported to have mosaic expression. Hi-Myc mice show very high Myc expression and high volume disease present in the lateral prostate lobe, and scattered Myc expression and low volume disease present in the anterior prostate.<sup>11</sup>

High levels of expression were detected in preinvasive (PIN) lesions and invasive adenocarcinoma, so we assayed for PSMA expression at very early time points to determine when PSMA expression began in Hi-Myc mice. In this model, the AR-driven Probasin promoter drives Myc expression as early as 2 weeks of age, and PIN lesions begin to develop shortly after Myc expression begins. At 4 weeks of age, no PSMA was detectable despite the presence of Myc-induced PIN lesions in the prostate (Figure 2A). At 8 weeks of age, PIN is widespread in the Hi-Myc prostate, and PSMA is expressed at high levels in PIN lesions (Figure 2B). PSMA expression continues at 6 months (Figure 2C), and most mice will have PSMA positive invasive lesions at this time point.<sup>32</sup>

Myc expression and PIN development in Hi-Myc mice is heterogeneous in most prostate lobes, and areas of normal prostate cells interspersed between PIN lesions served as negative internal controls. In order to determine if PSMA expression is a feature of all prostate cancer models, or Hi-Myc in particular, we assayed for PSMA expression in TRAMP (Figures 2D and 2E) and prostate-specific PTEN deleted mice (Probasin-Cre x PTEN fl/fl, Figure 2F). No PSMA expression was detected in TRAMP PIN lesions (Figure 2D), invasive TRAMP lesions (Figure 2E), or PB-Cre PTEN lesions (Figure 2F).

To this point, all detection of PSMA expression was accomplished using a single antibody reagent in genetically engineered control and diseased mice. To further confirm our findings of PSMA expression in a mouse model, and to accomplish this in an in vivo setting, we utilized the exemplary GCPII-ligand 2-(3-(1-carboxy-5-[(6-[<sup>18</sup>F]fluoro-pyridine-3carbonyl)-amino]-pentyl)-ureido)-pentanedioic acid ([<sup>18</sup>F]-DCPyL).<sup>12,13</sup> Animals under isoflurane anesthesia were administered 9.25 MBq [<sup>18</sup>F]-DCPyL and imaged at 60 min post-injection. Representative volume rendered datasets are shown in Figure 1. PET volumes are shown with dual intensity scales (for without and with bladder segmentation) as



**FIGURE 1** Specific detection of PSMA in the Hi-Myc model of prostate cancer. Immunohistochemistry using anti-PSMA antibodies in wild-type mouse kidney (A) shows high expression in proximal tubules. Top (low 20×) and bottom (high 40×) magnification of the same representative sections. Arrows indicate PSMA staining of the Bowman's capsule of the glomerulus. Negative staining in the kidney of a *Folh1* knockout mouse (B) confirms antibody specificity. Human prostate cancer (C) shows high PSMA expression concentrated at cell membranes. Normal mouse prostate (D) shows no staining. Hi-Myc PIN (E) and adenocarcinoma (F) show high PSMA expression with a pattern similar to human cancer. Western blot (G) shows similar results with expected expression in kidney, brain, and salivary gland (SG), but not in wild-type prostate anterior lobe (AP) or lateral lobe (LP). Hi-Myc prostates show PSMA expression proportional to the expected tumor burden in AP (low burden) and LP (high burden) [Color figure can be viewed at wileyonlinelibrary.com]



**FIGURE 2** PSMA expression is limited to the Hi-Myc model. Hi-Myc mice (A-C) show no PSMA expression in PIN at 4 weeks of age (A), but high expression is present at 8 weeks (B) and 6 months (C). TRAMP PIN (D), TRAMP invasive tumors (E), and PTEN deleted tumors (F) show no PSMA expression. Top (low 20×) and bottom (high 40×) magnification of the same representative sections [Color figure can be viewed at wileyonlinelibrary.com]

bladder activity concentration exceeds 100%IA/mL. In the two perspectives, one can visualize the caudally oriented dorsal and lateral prostate lobes, with some signal ascending rostrally along the anterior lobular projections of the prostate (also known as the coagulating glands). No prostatic focus of [<sup>18</sup>F]-DCPyL-PET signal is detected in the TRAMP mice (notably with significantly greater volume of disease in these age matched animals.<sup>14</sup> Likewise, wild-type mice have no detectable prostatic signal (dissection results shown below).

To confirm these in vivo results, the genitourinary package was promptly removed in its entirety, microdissected at 4°C and arrayed (Figure 4A). The tissues were rescanned and uptake, corrected for decay and injection activity, was analyzed (Figures 4B and 4C). Recapitulating the in vivo imaging and pathobiological findings, signal in the lobes of the prostate bearing Myc-driven disease displayed considerable accumulation of [<sup>18</sup>F]-DCPyL.

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#### 3 | DISCUSSION

In these experiments, we have established the presence of PSMA in a spontaneous, autochthonous, and well regarded model of prostate adenocarcinoma development. Beginning with immunohistochemical evaluation, we have verified that the reagents used to interrogate mouse and human tissues were specific for mouse and human PSMA. Knockout animals, which have intact *GcpIII (Folh2)*, did not stain positive and served to underline the specificity of our approach. In contrast with some previous reports, we detected minimal to no expression of PSMA in normal and TRAMP prostates. We relied on detection of PSMA protein using a monoclonal antibody and a small molecule, while some reports have detected *Folh1* RNA. The reason for this discrepancy is unclear, but differing sensitivity of these methods or detection of highly similar homologs may explain these differences.

There are many agents under development for improved detection and targeted therapy of prostate adenocarcinoma.<sup>15-19</sup> The widespread use of high-sensitivity and robust methods to detect PSMA in particular in the locally advanced and metastatic settings by PET and SPECT are poised to revolutionize management for those afflicted with the disease. Further, many of these agents can be derivatized to deliver radioisotopes with therapeutic intent.<sup>20-22</sup> This creates the urgent need to better understand the underlying biology of the target protein, its expression in disease states and its response to novel and conventional therapies. We pursued in vivo imaging in order to evaluate whether the expression levels detected by molecular biology and histological tools were sufficient to annotate disease status non-invasively in the Hi-Myc model. Utilizing the urea-based [<sup>18</sup>F]-DCFPyL, a high-resolution

GCPII-ligand with optimized pharmacokinetics in late-stage clinical evaluation (NCT03181867, NCT02981368), expression in the prostate can be detected in these animals. This is notable given the considerable background signal provided by the bladder when using this renally excreted agent (Figure 3). No detectable signal was observed in wild-type or TRAMP model animals. Excised tissue imaging was performed in order to more clearly visualize expression in the lobes of the prostate. Differential disease development is known in this model; within the dorsolateral and ventral lobes are concentrations of proliferating lesions at earlier time points and to a greater degree<sup>11,15,23</sup> and to quantitate uptake at the sub-organ level. These values also correspond with immunohistochemical findings in Figure 1.

There is no definitive explanation for the absence of GCPII expression in normal mouse prostate, in contrast to that in man. One partial explanation for this is that models of prostatic disease are limited by several factors, key among these is that rodents, used for the bulk of biological evaluation in mammalian systems, do not naturally develop prostate cancer. Another is that within mice, PSMA expression patterns vary from sites in man. Some organs, including the gut, kidney and brain share common GCPII profiles across species. For example, while it has become evident that GCPII is often expressed in the neovasculature of several types of cancer in man, the presence of the protein is absent in xenograft models of cancer in mice.<sup>24</sup>

Our finding of PSMA expression in the pre- and cancerous prostate of a well regarded mouse model is significant as it may be useful in elucidating the reason for the differences in expression in human and mouse tissues. Moreover, the use of PSMA-targeting







**FIGURE 4** PSMA-radioligand uptake in excised and dissected prostate. A, Photograph of representative dissected genitourinary package including seminal vesicle (SV) and anterior prostate (AP), dorsal (DP), lateral (LP), and ventral (VP) from three mice strains Hi-Myc<sup>FVB</sup>, TRAMP, and WT-C57BI/6. Muscle tissue for reference background is included in the central sample (scale bar 5 mm). B, Ex vivo microPET acquisitions of the dissected tissues as above. Uptake is present in the Hi-Myc samples, notably in the VP, and DLP. C, Quantification of volumes of interest of the tissues recapitulates the in vivo and ex vivo imaging results [Color figure can be viewed at wileyonlinelibrary.com]

agents is now being firmly established in clinical practice for diagnostic and therapeutic intent. We have previously found mouse models of considerable value in order to study the pharmacokinetic and radiobiological profile of established imaging agents and emerging radiotherapies.<sup>30,31</sup> This work shows that the ability to direct molecularly specific ligands to diseased tissues may recapitulate clinical absorbed dose profiles at the tumor and off-target tissues is of considerable value to assess and optimize methods.

To conclude, our work extends the biological utility of this genetically engineered model system for use towards understanding the role of this abundant prostate cancer cell-surface marker in disease progression, and in drug development applications for anti-PSMA imaging and therapy evaluation.

#### 4 | METHODS

#### 4.1 | Mice

All experimental procedures were approved by the Johns Hopkins Institutional Animal Care and Use Committee (IACUC). Wild-type C57BL/6J, FVB/NJ, and TRAMP mice (C57BL/6-Tg(TRAMP)8247Ng/ J) were obtained from Jackson Laboratories (Bar Harbor, ME, Stocks 664, 1800, and 3135). FVB-Tg(ARR2/Pbsn-MYC)7Key (Hi-Myc, Strain 01XF5) mice were obtained from NCI Mouse Repository (Frederick, MD). PSMA-null mice were a gift of Dr Joseph Neale (Georgetown University, Washington, DC), who had received them originally from Warren Heston (Cleveland Clinic Lerner Research Institute, Cleveland, OH).<sup>25</sup> Briefly, this GEMM has excised exons 3-5 of *Gcpll/Folh1* and backcrossed in C57Bl/6>10 generations to generate homozygous PSMA-null mice. PSMA is undetectable in the knockout by PCR or Western blot, and knockout mice express no basal phenotypic differences.<sup>26,27</sup> Probasin-Cre x PTEN<sup>fl/fl</sup> mice were generated as previously described.<sup>28</sup> Genotyping was performed using primer sets and protocols recommended by the vendor. Genomic DNA for PCR was isolated from tails.

#### 4.2 | Immunohistochemistry

Slides prepared from formalin fixed tissues were deparaffinized and rehydrated before steaming in EDTA pH 8 (Invitrogen) for 40 min. Endogenous peroxidases were quenched with BLOXALL (Vector Labs), and the slides were blocked for 1 h with Serum Free Protein Block (Dako). Slides were incubated with antibodies directed against PSMA (Cell Signaling Technology rabbit monoclonal antibody (D718E). Staining was visualized with ImmPRESS Polymer detection kit and ImmPACT DAB (Vector Labs).

#### 4.3 Western blot

Lysates were prepared from flash frozen tissue samples using RIPA buffer with protease and phosphatase inhibitors added (Sigma). After separation by SDS-PAGE, samples were transferred to PVDF membranes and blocked with 5% NFDM solution. Blots were incubated with antibodies directed against PSMA (Cell Signaling Technology rabbit monoclonal antibody (D7I8E) diluted in 5% NFDM for 1 h, followed by peroxidase labeled secondary antibody and visualized with SuperSignal chemiluminescent substrate (Thermo Fisher).

#### 4.4 | In vivo and ex vivo PET

Age matched wild-type (FVB and C57BI/6), TRAMP, Hi-MYC<sup>FVB</sup>, and Hi-MYC<sup>C57BI/6</sup> male mice were administered 9.25-13 MBg (250-300  $\mu$ Ci) [<sup>18</sup>F]DCFPyL intravenously (*n* = 2). After 1 h uptake time, animals were anesthetized with isoflurane and placed prone on in the center of the field of view of a dedicated rodent microPET R4 (Concorde Microsystems). List-mode data were captured in order to acquire  $25 \times 10^6$  coincident events (approximately 15 min) using a  $\gamma$ -ray energy window of 350-750 keV, and a coincidence timing window of 6 ns. Following imaging, representative animals were sacrificed and the genitourinary (GU) package and bladder removed. The excised GU was placed in a Petri dish on an ice slurry and dissected using a stereomicroscope (Zeiss Stemi 2000). The separated tissues were placed in a clean 4 cm diameter dish with 0.5 mL of ice water and rescanned in the microPET R4 for whichever transpired first:  $5 \times 10^{6}$ coincident events or 30 min. Data were sorted into two-dimensional histograms by Fourier re-binning, and images were reconstructed by a maximum a posteriori and 3D filtered back projection. Image data were normalized to correct for non-uniformity of response of the PET, deadtime count losses, positron branching ratio, and physical decay to the time of injection and quantification determined relative to injected

activity (to yield percent injected activity per gram [%IA/g] or per mL [%IA/mL]). No attenuation, scatter, or partial-volume averaging correction was applied. Images were analyzed by using ASIPro (Concorde Microsystems). Regions of interest were delineated on sequential two dimensional slices to compute uptake values per mL of tissue specimen. For in vivo 3D rendered PET data a weighted average of the intensities along projections through the tomographic image is computed, and overlaid with a second volume in which the bladder and kidneys have been thresholded out. The image intensity scales represent the full image intensity space of 0-100% (white-red) and 0-2.5% (blue-green) using Amira 6.5 (FEI). [<sup>18</sup>F]DCFPyL was obtained from the Johns Hopkins University School of Medicine PET Center and synthesized with high specific activity (>150 GBq/µmol) as previously reported.<sup>29</sup>

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#### CONFLICT OF INTEREST

The authors have no disclosures or affiliations that are relevant for the subject matter of this report.

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