Androgen Ablation Augments Human HLA2.1-Restricted T cell Responses to PSA Self-Antigen in Transgenic Mice

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BACKGROUND. In recent years, there has been an increasing interest in targeting human prostate tumor-associated antigens (TAAs) for prostate cancer immunotherapy as an alternative to other therapeutic modalities. However, immunologic tolerance to TAA poses a significant obstacle to effective, TAA-targeted immunotherapy. We sought to investigate whether androgen deprivation would result in circumventing immune tolerance to prostate TAA by impacting CD8 cell responses.

METHODS. To this end, we generated a transgenic mouse that expresses the human prostate-specific antigen (PSA) specifically in the prostate, and crossed it to the HLA-A2.1 transgenic mouse to evaluate how androgen deprivation affects human HLA A2.1-restricted T cell responses following immunization of PSA-expressing mice by vaccinia-PSA (PROSTVAC).

RESULTS. Our PSA transgenic mouse showed restricted expression of PSA in the prostate and detectable circulating PSA levels. Additionally, PSA expression was androgen-dependent with reduced PSA expression in the prostate within 1 week of castration, and undetectable PSA by day 42 after castration as evaluated by ELISA. Castration of the PSA/A2.1 hybrid mouse prior to immunization with a PSA-expressing recombinant vaccinia virus resulted in a significant augmentation of PSA-specific cytotoxic lymphocytes.

CONCLUSIONS. This humanized hybrid mouse model provides a well-defined system to gain additional insight into the mechanisms of immune tolerance to PSA and to test novel strategies aiming at circumventing immune tolerance to PSA and other TAA for targeted prostate cancer immunotherapy. *Prostate 70: 1002–1011, 2010. © 2010 Wiley-Liss, Inc.

KEY WORDS: prostate cancer; immune tolerance; immunotherapy; vaccinia; androgen deprivation therapy

INTRODUCTION

To date, clinical trials of PCA immunotherapy included immunization with defined antigenic preparations such as synthetic peptides [1–3], antigen-[4,5] or mRNA-loaded dendritic cells [6], manipulated tumor cells [7], or with plasmid DNA [8] or viral vectors engineered to express immunogenic genes [9]. In particular, recombinant vaccinia virus expressing prostate-specific antigen (PSA) was tested in clinical trials in combination with recombinant PSA-expressing fowlpox virus [10,11], demonstrating its ability to trigger a specific immune response [11] and an 8.5-month improvement in median overall survival in men with metastatic castration-resistant prostate cancer.

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cancer [10]. However, the survival benefit remains modest, suggesting an opportunity and need for significant improvement of efficacy in PCa immunotherapy.

In 1999, a study reported combined immunization with androgen deprivation to modulate antigen expression as a means of circumventing tolerance to a prostate tumor-associated antigen (TAA)-specific in the setting of a human PCa clinical trial that targeted PSA as a prototype PCa TAA [12]. The scientific basis for combining androgen deprivation and PCa TAA-specific immunization was subsequently validated in mouse models that focused on MHC class II-mediated helper T cell responses [13,14]. Androgen ablation results in a rapid involution of benign and neoplastic prostate tissue at both primary and metastatic sites, seemingly because of apoptosis of androgen-dependent epithelial cells [15–17]. This treatment has been shown to induce infiltration of lymphocytes, macrophages and dendritic cells into the prostate and trigger inflammation [18]. Such infiltration proved to be beneficial for PCa immunotherapy as it provides a synergistic help by increasing the number of tumor antigen-specific lymphocytes [13,19,20]. Additional supporting evidence for this concept was provided by the demonstration that testosterone can be immunosuppressive by stimulating tumors to secrete TGF-β, a cytokine that promotes the expansion of Treg [21]. Androgen deprivation has been shown to result in a decrease in expression of these genes in both patients undergoing androgen ablation therapy and in human PCa cell lines [22]. In mice, androgen deprivation-induced TAA gene downregulation has been shown to circumvent immune tolerance and enhance CD4 T cell responses to prostate TAA [13]. However, the effect of androgen deprivation on the generation of TAA-specific CD8 T cells has not been addressed before.

In this report, we describe the initial characterization of transgenic mice expressing the PSA transgene in an androgen-regulated and prostate-specific manner. To further refine current prostate cancer models, we developed a unique double-transgenic mouse model co-expressing PSA and HLA-A2.1 to facilitate the investigation of PSA-specific tolerance in the context of human MHC. Finally, we sought to determine whether castration of male mice prior to immunization to PSA improves class I MHC-restricted T cell responses to this clinically relevant target antigen.

**MATERIALS AND METHODS**

**Peptides, Proteins and Viruses**

Previously described HLA-A2.1-restricted, PSA-derived peptide PSA-3 (VISNDVCAQV) and its agonist PSA-3A (YISNDVCAQV) [23] and the H-2Kb restricted-, SV40 Tag-derived Tag-IV peptide (VYYDFLKC) [24,25] were purchased from the Macromolecular Resources Facility at Colorado State University (Fort Collins, CO). Human PSA was purchased from Calbiochem (San Diego, CA). Vac-PSA was obtained from Therion Biologics Corporation (Cambridge, MA). Vac-mTag was generated in our laboratory [26–28].

**Generation of Transgenic rPB-PSA Mice, Construction of PSA Transgenic Map**

Human prostate-specific antigen (PSA) prepro-cDNA (accession no. M26663, NID g618463) was ligated to rat probasin (rPB) promoter cassette (GI:10000942, –426 to +28) [29] at the introduced BamH1 site. The rPB-PSA was ligated to a bovine growth hormone (bGH, GI:2168498, 1,872–2,028) terminator sequence at the BglIII site. The final rPB-PSA transgenic construct was microinjected into oocytes of foster mouse mothers and offspring were evaluated by PCR using PSA primers. The transgene transmission to the progeny of potential founder mice was verified by PCR and Southern blot. From among several candidate founders we have generated, the mouse expressing stably the highest levels of PSA as detected by ELISA was selected as the founder to propagate hybrid transgenic colonies.

The rPB-PSA transgenic map was constructed to confirm boundary sequences between the rat probasin promoter region, PSA open reading frame (ORF) and the bGH terminator. Primer sets located in the promoter-PSA and PSA-terminator flanking regions were designed and utilized to generate overlapping PCR fragments for complete DNA sequencing of the boundary regions.

**Generation of PSA/HLA-A2.1 Double Transgenic Mouse**

Hybrid PSA and HLA-A2.1 double transgenic mice (PSA/A2.1) were generated by cross-breeding PSA mice with HLA A2.1/H2Kb transgenic mice (Harlan Sprague Dawley, Indianapolis, IN). All double transgenic mice were heterozygous for each of their transgenes and as confirmed by PCR.

**Genotyping of PSA Transgenic Mice**

Tail snippets from 3-week-old transgenic mice were collected and their DNA isolated. PCR was used to genotype transgenic progeny using the following PSA primers (Invitrogen): 

PSA-FOR: 5’ ACCATGTGGGCTCCCGTTG 3’ and PSA-REV: 5’ TCAGGGGTGCCCCGATG 3’.
DNA quality was tested with a β-globin PCR performed concurrently in each of the PSA reactions using the following primers: FOR: 5′ GCCAATCTGCTCACACAGGATA 3′ and REV: 5′ CATGCAGCTTGTGCACTGGA 3′.

PCR reactions contained 200–300 ng of genomic DNA template, 10 mM Tris/HCl pH 8.3, 50 mM KCl, 200 μM dNTPs, 1 μM each of forward and reverse primers, 1.25 U Taq DNA polymerase (Applied Biosystems) and 1.5 mM MgCl₂ in a 25 μl final reaction volume. PSA-β-globin PCR was performed using PSA and globin primers in 25 μl reaction at 94°C, 3 min followed by 30 cycles of 94°C for 1 min, 55°C for 1 min, 72°C for 2 min, and final elongation at 72°C for 10 min. Samples were stored at 4°C until gel electrophoresis was performed using a 1% agarose gel in 1× TAE buffer.

Extraction of RNA and RT-PCR

Total RNA was isolated from the prostate and other tissues using the Nucleospin RNA II Kit (Clontech, Mountain View, CA). RT-PCR was performed using a reverse transcriptase kit (Promega, Madison, WI). PSA primers were as described above, and β-actin primers that were used for control reactions as follows: β-actin FOR: 5′ TGTGATGTCACGCACGATTTCC 3′ and REV: 5′ TTTGATGTCACGCACGATTTCC 3′. A parallel RT-PCR reaction without reverse transcriptase served as a negative control. PCR products were visualized on a 1% agarose gel.

Castration of PSA Transgenic Mice

Mice were anesthetized with a ketamine–xylazine mixture (ratio 90 mg/kg ketamine to 4.5 mg/kg xylazine) administered i.p. A lower abdominal incision was made, the testicular blood supply isolated and ligated with electrocautery. The testes were removed and the vasa cauterized. Mice were re-explored at various time intervals (e.g. 7, 14, 21, 28, and 42 days) for harvesting of tissues.

Quantitation of PSA Levels by ELISA

Tissues were homogenized by mincing with surgical blade, resuspend in equal volume of Buffer A (50 mM Tris, KCl 1.15% pH 7.5) and Buffer C (10 mM KPO₄, 0.25 mM Sucrose pH 7.4) (max volume of 500 μl), and passed through a 21G needle with 1 ml of syringe. Cells were sonicated at 30% output (180W) for 10 sec pulse (three times). The lysate was centrifuged at 13,000 rpm for 10 min at room temperature. The protein concentration in the supernatant was calculated using the Bio-Rad protein assay kit (Bio-Rad Laboratories, Hercules, CA). 120 μl of lysate containing 50 μg of protein was used for PSA ELISA. Serum PSA concentration was measured by AxSYM PSA assay (AxSym system, Abbott Diagnostics, Abbott Park, IL), a Micro-particle Enzyme Immunoassay designed to quantitate PSA levels in human serum.

One hundred micrograms of total protein was placed in 250 μl of clear Hanks balanced salt solution and samples were submitted to the University of Michigan clinical pathology laboratory for PSA ELISA. Samples consisting of purified PSA from human semen or protein from wild type C57BL/6 mice were submitted to serve as positive and negative controls, respectively.

Mouse Whole Blood Processing and HLA A2.1/Kb FACS Analysis

One hundred microliters of blood was obtained from each mouse via orbital puncture and placed in a heparinized microfuge tube. Blood was placed in 2 ml lysis buffer (Mouse Erythrocyte Lysing Kit, R&D Systems Inc.) for 10 min at room temperature. Samples were centrifuged for 5 min at 250 × g, the supernatant was discarded, and the pellets were washed with 2 ml of 1× PBS. One million peripheral blood cells were incubated with 0.5 μg anti-mouse FcγR antibody (Mouse Fc Block, BD PharMingen) for 10 min at 4°C. 0.5 μg of FITC-conjugated anti-HLA A2 antibody (One Lambda, Inc., Canoga Park, CA) or isotype control was then added and the reaction mixture was incubated in a final volume of 20 μl at 4°C for 1 hr. Cells labeled with isotype control were used to assess background fluorescence, and 10,000 viable cells were analyzed in a FACScan microfluorometer (Becton Dickinson, Sunnyvale, CA).

Immunization of Castrated PSA/A2.1 Transgenic Mice

PSA/A2.1 mice were anesthetized with a ketamine–xylazine mixture (ratio 90 mg/kg ketamine to 4.5 mg/kg xylazine) administered i.p. A lower abdominal incision was made, the testicular blood supply isolated, and ligated with electrocautery. The testes were removed and the vasa cauterized. Four weeks post-castration, mice were immunized intravenously with 10⁶ PFU of a recombinant vaccinia virus expressing either the entire sequence of PSA (vac-PSA) [30] or the modified SV40 T antigen (vac-mTag) we previously generated in our laboratory [26–28].

Generation of Bone-Marrow-Derived Dendritic Cells

Dendritic cells (DC) were generated from bone marrow using IL-4 and GM-CSF (PeproTech, Rocky Hill, NJ), and purified on a metrizamide gradient. DC
maturation was evaluated by measuring MHC Class II, CD11c, CD40, CD80, and CD86 surface expression. Peptides (final 20 μg/ml) or proteins (50 μg/ml) were loaded on DC at 37°C for 4 hr or overnight, respectively. Loaded DC were washed and used as antigen-presenting cells in the ELISPOT assay.

**IFN-γ ELISPOT Assay**

MultiScreen 96-well plates were first coated with purified anti-mouse IFN-γ antibody (capture antibody) (4 μg/ml in 1× PBS; PharMingen) overnight at 4°C. Plates were blocked with PBS/1% BSA (PBS-BSA) at room temperature for 30 min and then washed thrice with 1× PBS before seeding the cells. CD8+ T cells were isolated from mice 3 weeks after immunization with 106 pfu/mouse vac-PSA or 107 vac-mTag. One million CD8+ T cells were seeded into each well (E/S = 10) and incubated at 37°C/5% CO2 for 24 hr with irradiated (5,000 rad) peptide- or protein-loaded DC. Plates were washed thrice with 1× PBS and then four times with 1× PBS/0.025% Tween-20 (PBS-TW20) before adding biotin rat anti-mouse IFN-γ antibody (2 μg/ml in PBS-BSA; PharnMingen) overnight at 4°C. The plates were washed four times with PBS-TW20 and incubated with anti-biotin antibody (1:1,000 dilution; Vector; Burtingame, CA) at room temperature for 90 min, followed by washing four times with PBS. Plates were then developed with NBT/BCIP before subjecting to an ELISPOT reader (Cellular Technology Laboratories, Ltd.; Cleveland, OH) to count spots.

**RESULTS**

**Prostate-Specific, Androgen-Dependent Expression of PSA in a Transgenic Mouse**

The rat probasin gene promoter has been demonstrated to be both developmentally and hormonally regulated in the mouse and demonstrates a high ability to direct transgene expression specifically to the prostate tissue [31]. The promoter has been previously utilized to generate transgenic mice that express viral [29] and human oncogenes [32,33] in the prostate to study prostate cancer development. We adopted a similar strategy to generate a transgenic mouse that exhibits prostate-specific expression of human PSA. To this end, an expression cassette containing human PSA cDNA, the rat probasin promoter, and the bovine growth hormone terminator sequence (Fig. 1A) served to generate transgenic founders. The founder that produced progeny that had stable and high levels of PSA expression was selected to propagate the colony.

To address the specificity of the PSA expression, we isolated RNA from prostate, testis, seminal vesicle, spleen, pancreas, liver, lung, and nodal tissue from multiple progeny. Using RT-PCR, we confirmed that the presence of RNA transcript for PSA was restricted to the prostate tissue (Fig. 1B). In line with this finding, analysis of protein extracts from these tissues by ELISA detected PSA in the prostate but not in any of the other tissues (Fig. 2A).

To test androgen-responsiveness of the PSA transgene, mice were castrated and killed at various time points, and the presence of PSA mRNA was tested by RT-PCR. As shown in Figure 2B, expression of the transgene decreased in a time-dependent fashion after castration. Only a minimal expression was present at day 28 post-castration, and no transcripts were detected at day 42.

**Generation of a PSA/HLA-A2.1 Hybrid Mouse**

To generate a hybrid mouse that expresses both human HLA-A2.1 and PSA, PSA and HLA A2.1/Kb transgenic mice were crossed and the progeny tested by PCR of genomic DNA (data not shown). Progeny that possessed copies of both transgenes (data not shown) were further tested for HLA-A2.1 expression by flow cytometry. These double-positive mice demonstrated expression of the HLA A2.1/Kb chimeric receptor on peripheral blood lymphocytes as demonstrated by FACS analysis. Some of these mice were killed and protein extracts from mouse prostate tissue contained high levels of PSA as demonstrated by ELISA (data not shown).

**Androgen Deprivation Augments CD8 T Cell Responses to Prostate TAA**

Consistent with the known immunosuppressive effects of androgens, castration of prostate tumor-bearing mice has been shown to enhance antigen-specific CD4 T cell responses to a model prostate TAA [13]. We therefore sought to determine whether CD8 responses are affected by androgen deprivation using the PSA/A2.1 as a model and PSA or PSA-derived peptides as immunogens. Male A2.1 and female PSA/A2.1 mice were used as controls.

Mice were immunized with recombinant vaccinia virus constructs that express either PSA (vac-PSA) or SV40 Tag (vac-mTag). Splenocytes from immunized mice were re-stimulated in vitro with peptide- or full protein-loaded dendritic cells and IFN-γ-releasing CD8 T cells were quantitated by ELISPOT and tetramer staining. Our data show that splenocytes from sham-castrated male, castrated male, and female PSA/A2.1 mice immunized with vac-mTag responded in a similar fashion to restimulation with Tag-IV, while no response was observed with splenocytes from vac-PSA immunized mice. A similar response to Tag IV was
obtained in control vac-mTag-immunized A2.1 mice, indicating a complete absence of tolerance to exogenous antigen in these mice (Fig. 3A). Immunization with vac-PSA resulted in a strong and specific response in A2.1 mice upon re-stimulation with full PSA-loaded DC. However, a very weak response was observed in PSA/A2.1. Castrated mice responded better ($P < 0.001$), although this response was still significantly lower than male A2.1 and female PSA/A2.1 mice ($P < 0.001$) (Fig. 3B). Splenocytes from vac-PSA-immunized mice showed a much lower number of IFN-γ-releasing, PSA-specific CD8 T cells upon re-stimulation with DC loaded with the peptides PSA3A (Fig. 3C) and PSA3 (Fig. 3D). Such a weak response to PSA peptides as compared with full PSA protein is possibly because the contribution of H2-Db to the response to potential H2-Db-restricted, PSA-derived epitopes in this mouse model. Sham-castrated male PSA/A2.1 showed no response, whereas castrated mice showed a weak response. Because of the overall signal to noise window for ELISPOT evaluation of A2.1-restricted PSA peptide responses (females vac-mTag controls showed low but significant activity of PSA peptides, Fig. 3C and D), we turned to tetramer assay to ascertain detection of A2.1-restricted, PSA-specific CTL. To this aim, we labeled in vitro-restimulated splenocytes with a PSA-3A-A2.1 tetramer and used flow cytometry to determine the fraction of CD8(+) T cells that bound the tetramer. Consistent with the ELISPOT data, tetramer staining revealed elevated amounts of PSA-specific CD8 T cells in spleens of immunized non-transgenic mice. These amounts were significantly lower (comparable to background level as seen in vac-mTag-immunized mice) in PSA transgenic mice, reflecting a deep immune tolerance to PSA. Splenocytes from castrated immunized male
PSA/A2.1 mice exhibited significantly higher numbers of tetramer-positive CD8 cells than did their non-castrated counterparts. The effect of castration was similar to the one observed in A2.1-PSA females, but still lower than in A2.1 male mice (Fig. 4).

**DISCUSSION**

A large body of evidence suggests the feasibility of PSA-based vaccines for prostate cancer [2,3,12,34–41]. However, because PSA is a self-antigen, these vaccines face the challenge of immune tolerance to PSA and are yet to demonstrate clinical efficacy. Hence, immunotherapy for PCa will most likely consist of combined approaches that simultaneously target tumors through TAA and interfere with tolerizing mechanisms that hinder immunity to tumors. These mechanisms include innate and adaptive immune responses, activation/inhibition of co-stimulatory/inhibitory molecules (e.g., CTLA-4), elimination of regulatory/suppressive cells (e.g., regulatory T cells—Treg) and soluble factors, and manipulation of hormonal pathways. So far, interference with the inhibitory co-stimulatory signals mediated by CTLA-4 using monoclonal antibodies has provided the most promising results, although a myriad of adverse autoimmune responses generated by this strategy still represent a sizeable obstacle to its implementation in clinic (reviewed in Ref. [42]).

In prostate cancer, androgen deprivation is a therapeutic modality that aims at depriving the prostate from testosterone to suppress its growth. Interestingly, this deprivation also results in strengthening immunity to tumors as shown in mice [13] and humans [12]. The mechanisms whereby androgen deprivation affects immunity to tumors are poorly understood, partly because of the lack of appropriate mouse models.

Although the impact of androgen deprivation on prostate tumor immunity has been addressed in previous work, these failed to look at the extent to which CTL responses to a human prostate TAA are affected in vivo in humanized mice.

In this work, we describe a transgenic mouse that expresses human PSA specifically in the prostate. Tissue-specific expression was demonstrated at both RNA and protein levels. We additionally show that expression of PSA in this mouse is regulated by androgens, confirming an optimal performance of the rat probasin promoter in this setting. This is evidenced by a time-dependent decrease of PSA levels in castrated male mice. At day 42 post-castration, no PSA was detected.

We crossed the PSA transgenic mouse with HLA-A2.1/Kb transgenic mouse. The resulting male offspring offers a valuable tool that allows investigation of immune tolerance to a human prostate antigen in a way that closely emulates human biology.
As expected, immunization of this hybrid mouse to a non-self-antigen (SV40 Tag) resulted in a strong CTL response, whereas only a very weak response was generated towards a self-antigen (PSA). In contrast, castration of male mice 4 weeks prior to immunization resulted in a significant augmentation of CTL response to PSA, while it did not affect the response to SV40 Tag. Interestingly, female hybrid mice showed a stronger response to PSA than did their castrated male counterparts.

Although interference of castration with immune tolerance to prostate specific antigens has been reported in two previous studies [13,14], these studies either focused on CD4 responses [13] or did not show any effect of castration when applied before immunization [14]. In the latter study, a prime-boost immunization protocol was applied and immune responses following primary immunization were not evaluated.

Overall, our present work describes a PSA/A2.1 transgenic mouse that might represent an attractive animal model to investigate immune tolerance to human prostate TAA and for preclinical development and refinement of PSA-targeted vaccines. It also

**Fig. 3.** Castration of male PSA/A2.1 mice augments PSA-specific cellular immune responses. Castrated male mice were immunized with either vac-mTag or vac-PSA and antigen-specific CD8 responses were evaluated by ELISPOT and tetramer staining. Non-castrated male mice and female mice were used as controls. For splenocyte restimulation in vitro, bone marrow-derived DC were loaded with either H-2Kb restricted, SV40 Tag-derived epitope IV (Tag-IV/DC) (A), with PSA (PSA/DC) (B) or with HLA-A2.1-restricted, PSA-derived epitopes (PSA3A/DC) (C) and (PSA3/DC) (D). IFN-γ-releasing CD8 T cells were revealed using an ELISPOT assay. Data shown as mean ± SD, representative of one experiment with from three animals per group.
provides compelling evidence supporting the use of androgen deprivation as a modality to circumvent immune tolerance to prostate TAA.

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