Androgen receptor activity promotes resistance to BRAF-targeted melanoma therapy

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Abstract

Treatment with molecularly-targeted therapy has revolutionized cancer care, including BRAF/MEK-targeted melanoma therapy. However responses are heterogenous and frequently not long-lasting. Novel strategies to target resistance are needed. We studied a cohort of patients with resectable metastatic melanoma treated with neoadjuvant BRAF/MEK-targeted therapy (n=52) and noted a strong sexual dimorphism in response to treatment, with female patients demonstrating significantly higher rates of a major pathologic response (MPR) (p=0.0001). RNA sequencing of tumors demonstrated enrichment of androgen-related genes in those failing to achieve MPR. Pre-clinical studies validated these findings, with significantly increased tumor growth in male vs female mice treated with BRAF/MEK inhibitors (BRAF/MEKi) (p=0.0005). Androgen receptor (AR) expression was upregulated in tumors of BRAF/MEKi-treated mice, and modulation of AR signaling via AR-blockade or castration was associated with significantly slower tumor growth (p=0.0001 and p=0.00004, respectively). Together, these results have important implications in the context of treatment with BRAF/MEKi-targeted therapy.

Main Text

To date, multiple studies have demonstrated that male sex is associated with worse outcomes in patients with melanoma, including patients with stage III and IV disease treated with targeted therapy1-3. While these differences have been noted in melanoma as well as other malignancies, the driving factors are incompletely understood4-8. Variations in the immune background9,10, tumor microenvironment11-14 and tumor cell susceptibility to targeted therapy9,15 have been posited as driving factors of this sexual dimorphism in treatment outcomes. Additionally, innate hormonal differences in the estrogen and androgen pathways have been suggested as potential mechanisms driving observed disparities in melanoma preclinical models12,13,16-18 of immune checkpoint blockade19,20 and targeted therapy21. However further insights from patient cohort and pre-clinical studies are needed to help derive rational combinatorial strategies to improve patient survival.

We and others have recently shown neoadjuvant targeted therapy to be of significant clinical benefit in patients with high-risk, surgically resectable locoregional metastatic melanoma22,23. These studies provide a unique opportunity for in-depth molecular characterization and longitudinal study of biospecimens in a more homogeneous patient population. Furthermore, this work has demonstrated that early endpoints, including major pathologic response after neoadjuvant chemotherapy correlate with long-term outcomes22,24. In one of the first trials performed, we noted a strong sexual dimorphism in response to treatment with neoadjuvant BRAF/MEK targeted therapy in patients with locoregional melanoma metastases22,25, though sample size was limited (Supplemental Figure 1A).

To build on these findings, we pooled data from a cohort of melanoma patients with locoregional metastases who were treated with neoadjuvant BRAF-MEK targeted therapy within and outside of clinical trials (Figure 1A, Supplementary Table 1). Patients were treated with neoadjuvant BRAF-MEK targeted therapy for a duration of 8-12 weeks prior to definitive surgical resection. Pathologic responses were
assessed, and patients were dichotomized into those having a major pathologic response (MPR) (\( \leq 10\% \) viable tumor) versus those without a major pathologic response (>10% viable tumor) as defined by the International Neoadjuvant Melanoma Consortium\(^{26}\). In this pooled cohort, we again noted a strong sex difference in therapeutic response, with female patients experiencing a significantly higher rate of MPR on neoadjuvant dabrafenib and trametinib (DT) (22/30, 66%) compared to male patients (3/22, 14%) (odds ratio of 12.7 95%CI [3.0-53.2] \( p=0.001 \)) (Figure 1B, Supplementary Table 1). No other clinical factors were significantly associated with achievement of MPR on logistic regression analysis, including age (\( p=0.70 \)), stage III or D (\( p=0.45 \)), stage IV (\( p=0.53 \)), BRAF V600 E versus V600 non-E mutation (\( p=0.57 \)), ECOG performance status (\( p=0.94 \)), body mass index (BMI) (>30) (\( p=0.80 \)), serum lactate dehydrogenase (LDH) (\( p=0.27 \)), or recurrent disease (\( p=0.29 \)). Within the female subset, there was no association between menopausal status (pre- vs post-) and MPR; 9/13 (69%) pre-menopausal women and 11/17 (65%) post-menopausal women achieved a MPR (\( p=0.79 \)). Females also had significantly improved progression-free survival as compared to male patients (Figure 1C, \( p=0.043 \)).

To explore the potential molecular mechanisms underlying this observed sex-difference in response to neoadjuvant therapy, we analyzed the gene expression changes using available RNA sequencing analysis of pre-treatment samples of patient tumors from the initial clinical trial cohort (n=4 female, 5 male). From this analysis, we observed an enrichment of androgen-related pathways in non-responders, and estrogen-related pathways in responders (Figure 1D, Supplementary Figure 1B). We next assessed androgen receptor (AR) expression via immunohistochemical staining for androgen receptor percent positive nuclei in available matched pre- and on-treatment tumor biopsy specimens (n=4 female, n=4 male). This identified a trend toward higher AR expression in tumors after treatment with BRAF-MEK targeted therapy in both male and female patients (Figure 1E, \( p=0.06 \), Supplemental Table 2), with all male patient tumors tested exhibiting increased expression post-treatment.

Intrigued by these results, we then functionally evaluated the impact of sexual dimorphism on response to neoadjuvant targeted therapy using the \( Braf^{V600E}/Pten^-/- \) (BP) syngeneic murine melanoma model (female genotype). To do this, C57Bl/6 (immune competent) or CD-1 nude (immune incompetent) male and female mice were implanted subcutaneously with BP cells and treated daily with dabrafenib + trametinib (DT) once tumors reached an average size of \( \sim 400-600 \) mm\(^3\) (Supplemental Fig. 3A). In these studies, DT treatment was associated with delayed tumor outgrowth in all mice compared to vehicle-treated mice as expected. However male mice demonstrated significantly impaired tumor control on DT compared to female mice in an immunocompetent model as well as in nude mice (\( p < 0.0005 \) and \( p < 0.00001 \) respectively, Figure 2A, B and Supplemental Figure 3B, C), corroborating our findings in the clinical cohort, and suggesting that the mechanism is not immune-mediated.

Building upon the observation that AR staining was elevated is post-treatment samples samples of patients on neoadjuvant DT, we next analyzed AR expression via bulk transcriptome analysis and immunofluorescence in tumors of vehicle- and DT-treated mice with the hypothesis that DT treatment would be elevated in DT-treated mice. In these studies, a modest increase in AR mRNA expression was noted in females and males upon treatment with DT (\( p<0.005 \) and \( p=0.16 \) respectively) (Figure 2C).
However, AR protein expression was significantly increased in male DT-treated mice versus vehicle control (p=0.0003, Figure 2D, E), suggesting a potential role for testosterone in stabilizing AR and mediating resistance. Interestingly, there was no difference in AR protein expression in female DT-treated mice compared to vehicle control (p=0.9, Figure 2D, E). Reverse phase protein array (RPPA) of tumor samples increased phosphorylation (activation) of components of the MAPK and PI3K/mTOR pathways in male versus female mice following treatment with DT, which may also underlie differential response (Figure 2F).

Following this, we next sought to test whether AR signaling was underlying the observed treatment resistance. To do this, we evaluated the effect of modulating AR signaling on DT response through either AR blockade or castration in male mice, or via administration of testosterone in female mice (Supplementary Figure 3A,B). Results demonstrated significantly increased tumor outgrowth in female mice receiving DT + testosterone versus DT alone (Figure 3A; p<0.00001, Supplementary Figure 3C). Conversely, male mice treated with DT + AR blockade (enzalutamide) or castration demonstrated delayed tumor outgrowth compared to male mice treated with DT alone (Figure 3B; p=0.0001 and p=0.00004, respectively, Supplementary Figure 3D,E). Notably, treatment with DT + testosterone in males was associated with increased tumor growth over DT alone (Figure 3B; p=0.0005), suggesting that testosterone-regulated activity of AR contributes to the differential response to DT. Treatment with exogenous testosterone reversed the protective effects of both castration and AR blockade with enzalutamide in male mice (Figure 3C and 3D; p< 0.00001 and p= 0.005, respectively). Treatment with AR blockade + DT in female mice was associated with delayed tumor outgrowth compared to DT treatment alone (Figure 3E; p=0.007) suggesting that this combination may also be effective in females. Similar to males, treatment with exogenous testosterone + DT and AR blockade tended to abrogate the effects of AR blockade on DT treatment (p=0.08). (Figure 3E; p=0.02).

We next assessed levels of testosterone and AR expression in tumors of male and female mice treated with DT with or without modulation of AR signaling (via AR blockade, castration, or exogenous testosterone administration). In these studies, treatment with strategies to reduce AR signaling in male mice (castration, enzalutamide) were associated with reduced intra-tumoral testosterone levels (Figure 3F; 138.7 and 4.6 fold decrease, respectively). Conversely, strategies to increase AR signaling (exogenous testosterone) in female mice was associated with increased intra-tumoral testosterone levels (Figure 3G; 263.5 fold increase). Androgen receptor staining mirrored these findings, with a trend for decreased AR expression in DT + enzalutamide and a significant reduction in the DT + castration groups versus DT alone in male mice, which was abrogated with administration of exogenous testosterone (Figure 3H and 3I, p=0.3 for DT + enzalutamide, p=0.0003 for DT + castration). Although there is a slight increase in the percentage of AR-positive cells following DT treatment, expression remains very low. While a trend toward an increase in AR positive nuclei in mice receiving DT + testosterone was noted as well (Figure 3H and 3I, p=0.04, p=0.06 respectively, Supplemental Figure 3F).

To validate that androgen receptor signaling was driving resistance to BRAF-MEK targeted therapy, we
next generated AR knockout melanoma cells using CRISPR technology and implanted these into male and female syngeneic mice. These studies demonstrated that the modulation of response to DT treatment via administration of testosterone or enzalutamide was lost (Figure 3J and 3K), suggesting that the phenotypes observed in our prior studies are directly related to androgen receptor signaling. Negligible AR expression was confirmed via immunofluorescence (Supplementary Figure 3G). Taken together, these data demonstrate that AR stabilization by testosterone contributes to resistance to BRAF-MEK targeted melanoma therapy, and highlights the potential for AR blockade by enzalutamide as a therapeutic strategy to overcome this resistance. Though our current studies only explored these relationships in melanoma, studies interrogating AR signaling in the context of MAPK-targeted therapy are warranted across cancer types.

**Methods**

**Clinical cohorts:**

Patients enrolled in the initial clinical trial cohort were as previously described.1 Briefly, patients ≥18 years old with histologically-proven clinical stage III or oligometastatic stage IV \( \text{BRAF}^{V600E/K} \) melanoma deemed resectable by multidisciplinary consensus and measurable disease by RECIST 1.1 criteria were enrolled and those randomized to the experimental arm received 8 weeks of neoadjuvant dabrafenib (150mg PO BD) plus trametinib (2mg PO daily) prior to surgical resection, followed by up to 44 weeks of adjuvant dabrafenib and trametinib (neoDT)(n=12). Additional patients (n=20) from a single-arm study of neoDT along with an additional retrospective cohort of patients treated off protocol for logistic reasons with neoDT (n=16), Dabrafenib (n=3), and Encorafenib plus Binimetinib (n=1) were included. Radiographic responses to neoadjuvant therapy were determined at week 8 prior to surgery and pathologic responses were determined by microscopic examination of the complete surgical specimen by a melanoma pathologist, including SOX10 immunostains when applicable to confirm presence or absence of viable melanoma cells.

**RNA sequencing:**

Tumor biopsies were obtained as feasible by punch or core biopsy prior to and during the neoadjuvant treatment period. Fresh-frozen tumor biopsy material was used for RNA sequencing library preparation. Total RNA was extracted from snap-frozen tumor specimens using the AllPrep DNA/RNA/miRNA Universal Kit (Qiagen) following assessment of tumor content by a Pathologist, and macrodissection of tumor bed if required. RNA quality was assessed on an Agilent 2100 Bioanalyzer using the Agilent RNA 6000 Nano Chip with smear analysis to determine DV200 and original RNA concentration. Based on RNA quality, 40-80ng of total RNA from each sample then underwent library preparation using the Illumina TruSeq RNA Access Library Prep kit according to the manufacturer's protocol. Barcoded libraries were pooled to produce final 10-12 plex pools prior to sequencing on an Illumina NextSeq 500 sequencer using one high-output run per pool of 76bp paired-end reads, generating 8 fastq files (4 lanes, paired reads) per sample.
**RNA sequencing data processing:**

RNA-seq FASTQ files were first processed through FastQC (v0.11.5)\(^2\), a quality control tool to evaluate the quality of sequencing reads at both the base and read levels. Reads having \(^315\) contiguous low-quality bases (phred score <20) were removed from the FASTQ files prior to STAR 2-pass alignment (v2.5.3)\(^3\) with default parameters to generate one BAM file for each sequencing event. After that, RNA-SeQC (v1.1.8)\(^4\) was used to generate quality control metrics including read counts, coverage, and correlation. A matrix of Spearman correlation coefficients amongst all sequenced samples was subsequently generated by RNA-SeQC and after careful review the sequencing data generated from one library pool that showed poor correlation with other library pools from the same RNA sample was removed before sample-level merging of BAM files.

HTSeq-count (v0.9.1)\(^5\) tool was applied to aligned RNA-seq BAM files to count how many aligned reads overlapped with the exons of each gene. The raw read counts generated from HTSeq-count were normalized into fragments per kilobase of transcript per million mapped reads (FPKM) using the RNA-seq quantification approach suggested by the bioinformatics team of NCI Genomic Data Commons (GDC) [https://docs.gdc.cancer.gov/Data/Bioinformatics_Pipelines/Expression_mRNA_Pipeline/]. Briefly, FPKM normalizes read count by dividing it by the gene length and the total number of reads mapped to protein-coding genes using a calculation described below:

\[
\text{FPKM} = \frac{\text{RC}_g \times 10^9}{\text{RC}_{pc} \times L}
\]

RC\(_g\): number of reads mapped to the gene; RC\(_pc\): number of reads mapped to all protein-coding genes; L, length of the gene in base pairs (calculated as the sum of all exons in a gene). The FPKM values were then log2-transformed for further downstream analyses.

**RNA sequencing differential expression and gene set enrichment analysis:**

The HTSeq normalized read count data for all expressed coding transcripts were processed by DESeq2 (v3.6)\(^6\) software to identify differentially expressed genes (DEGs) between two response (responder vs. non-responder) groups. A cut-off for gene expression fold change of \(\geq 2\) or \(\leq -0.5\) and a FDR (q-value) of \(<.05\) was applied to select the most differentially expressed genes. Gene set enrichment analyses were performed using the GSEA software developed at the Broad Institute, using vst-normalised\(^7\) input transcriptome expression data and querying the MSigDB Hallmark gene sets with default parameters.

**Animals and xenograft studies:**

Female or male C57BL/6 (strain code: 0000664, purchased from Jackson Lab), aged 6 to 12 weeks, and weighing approximately 20 to 25 g were used for in vivo studies. Female or male CD-1 nude mice (strain code: 086, purchased from Charles River Laboratories), aged 6 to 12 weeks, and weighing approximately
25-40 g were used for the immunodeficient model in vivo studies. Animal health was monitored daily by observation and sentinel blood sample analysis. Animal experiments were conducted in accordance with the Guideline of IACUC at MDACC.

BP cells were scaled up in DMEM culture media supplemented with 10% FBS, harvested, and prepared so that each mouse received 0.8x10^6 cells in 0.2 mL PBS. Cells were implanted subcutaneously in the right flank of each mouse. For some male mice, physical castration was required and performed two weeks before treatment or before cell implantation. Testosterone pellets (Innovative Research of America) were implanted subcutaneously in the left flank one week before treatment. Trametinib (Chemietek) at 1 mg/kg and Dabrafenib (Chemietek) at 30 mg/kg were suspended at concentrations as needed in an aqueous vehicle containing 0.5% Hydroxypropylmethylcellulose and 0.2% Tween 80 in distilled water and adjusted to pH 8.80 with diluted NaOH solution. Enzalutamide (Chemietek) at 10 mg/kg was suspended in 1% carboxymethyl cellulose (Sigma), 0.1% Tween-80, 5% DMSO.

BP tumors were monitored by caliper before randomly sorting and dividing into experimental groups (n=10 mice per group). Treatment was started from day 14 to 17 days post-implantation depending on mouse strain and gender. Vehicle controls, mixture of Trametinib at 1 mg/kg and Dabrafenib at 30 mg/kg or Enzalutamide at 10 mg/kg were given orally using a sterile 1-mL syringe and 18-gavage needle for 21 days. Dosing was 5 hours apart between administration of Trametinib + Dabrafenib and Enzalutamide for these specific treatment groups.

Tumor volume was calculated using the following formula: 
\[V = \frac{L \times (D^2)}{2}\]
(in which L = length of tumors; W = width of tumor). Tumor and plasma were harvested 4 hours after the last dose. Tumors were snap frozen and the plasma was divided for monitoring drug concentrations and hormone levels. Endpoint samples were collected and processed for gene expression using Quant-seq and protein expression via Reverse Phase Protein Array (RPPA). RPPA analysis was performed at the MD Anderson Cancer Center's functional proteomic RPPA core facility as previously described.

Quant-seq library construction and sequencing.

1000 ng or 500 ng of DNase-treated RNA samples were converted to cDNA using a QuantSeq 3’ mRNA according to the manufacturer's protocol (Lexogen, Vienna, Austria). The libraries were amplified with 12 or 13 PCR Cycles and purified using the provided Lexogen. The purified libraries were quantified using a Kapa library quantification kit (KAPA biosystems) and loaded on NextSeq 500 Sequencer (Illumina, San Diego, CA) at a final concentration of 2.6 pM to perform cluster generation, followed by 1x76 bp sequencing on NextSeq 500 Sequencer (Illumina).

Androgen Receptor CRISPR KO in BP cell lines:

Three guide sequences were designed to target a 100 bp region of Exon 1 using HorizonDiscovery's CRISPR Design Tool. Guide 1 (gacctgggtagtctacatgg AGG) was cloned into pLentiCRISPR.v2 according to
addgene lentiCRISPRv2 and lentiGuide oligo cloning protocol for the purpose of pool selection. Guide 2 (gcttgatacgggcgtgtggat GGG) and Guide 3 (ctggagaacccattggacta CGG) were ordered as crRNA's (IDT). The Neon electroporation system was used to transfect ribonucleoprotein (RNP) complex + plasmid into the BP cells. The crRNAs were rehydrated to 200 nM and pooled in equal volume (3 µL each) for annealing with Alt-R CRISPR-Cas9 tracrRNA-ATTO 550 (IDT; 1075928); 95°C x 5 min, slowly cool to 10°C at 0.1°C/sec. 5 µL of crRNA:tracrRNA were combined with 5 µL of Alt-R S.p. HiFi Cas9 nuclease (IDT; 1081060) at room temperature for 10 min to generate the RNP complex. Electroporation Enhancer (2 µL) (IDT; 1075916) and 2 µg of plasmid were added to the RNP. During the annealing reaction, 1x10^6 BP cells were pelleted at 600xg for 3 min. The total volume of RNP + Plasmid + Electroporation Enhancer was transferred to an aspirated cell pellet. Next, 95 µL of R buffer was added to the cells and they were gently resuspended to single cells for an immediate 100 µL electroporation reaction. The Neon settings for the BP cells were two 30 ms pulses at 1150 volts. The cells were transferred to a single well of a 6 well plate and allowed to recover in growth media. After 24 hrs, the media was replaced and 1.5 µg/µL of Puromycin for a 48-hr selection. The cells were allowed to recover from selection for 48 hrs before single-cell clone selection. After clones were selected and expanded, AR protein was analyzed by Western Blot to confirm AR knockout. Briefly, cells were lysed for 30 min at 4 °C in 150 µL RIPA buffer with phosphatase and protease inhibitors. Protein (15 µg) was run on 10% SDS Page (Bio-Rad) and transferred overnight onto PDF membranes using TRIS Glycine Methanol buffer. After blocking for 1 hour at room temperature in 5% milk diluted in TSBS-T, AR antibody (Abcam) was incubated at 1:2000 dilution for 24 hrs. GAPDH was used as a loading control. The lentiviral expression vectors pLV-105 were purchased from Genecopeia to express either GFP as a transduction control or mouse Ar (NM_013476.3) for an overexpression control. Lentivirus was generated using standard protocols and psPAX2 and pMD2.G as the packaging vectors. BP cells were transduced at 90% efficiency with viral supernatant and selected for 48 hours with 2 µg/mL puromycin.

**Measuring testosterone levels from plasma:**

Testosterone quantification was determined using Agilent’s UHPLC Infinity II and 6495 triple quadrupole mass spectrometer, and MassHunter workstation software (8.0.8.23.5). Briefly, plasma samples were extracted with tert-butyl methyl ether (Sigma 34875), dried, and derivatized using hydroxylamine hydrochloride (Sigma 431362). The recovered ketoxime steroids were reconstituted in methanol/water (1:1 v/v) and injected into the Infinity II UHPLC. Ketoxime steroids were separated using a Chromolith reverse phase column (RP-18 endcapped 100-2mm, Sigma 152006) and introduced into a JetStream source (Agilent) for triple quadrupole analysis. Data were analyzed and quantified using MassHunter software (Agilent) and GraphPad PRISM 8 software was used to graph and perform statistics (two-sided Students t-test)\(^9,10\).

**Immunofluorescence**
FFPE blocks were sectioned (5 μm), mounted on charged microscope slides (Leica 38002092), and dried at 37°C overnight. Slides were then baked at 60°C for 1 hour in an oven (Biocare DRY2008US), deparaffinized in 3 changes of xylene, and then rehydrated in 3 changes of 100% ethanol followed by a series of 95%, 70%, and 50% ethanol and distilled water (5 min. each). Antigen retrieval (10 mM sodium citrate, pH 6.0 with 0.05% Tween 20) was performed by heating slides to 95°C for 15 minutes in a microwave (Biogenex EZ Retriever System v.3) followed by a 30 minute cool down at room temperature. Slides were then washed with TBST (Thermo TA-999-TT). The area around each section was traced with a PAP pen (Sigma Aldrich Z672548), blocked with Background Sniper (Biocare BS966) for 10 minutes, and then washed with TBST.

Primary antibody (rabbit anti-Androgen Receptor [EPR1535(2)], Abcam ab133273, 1:300, diluted in Dako S3022) was incubated overnight at 4°C, followed by a TBST wash. Secondary antibody (goat anti-rabbit Alexa-647 conjugate, Thermo A32733, 1:400, diluted in Fluorescence antibody Diluent, “FAD”, Biocare FAD901L) was added and allowed to incubate for 1 hour at room temperature, followed by a TBST wash. A sequential incubation with fluorophore-conjugated primary antibody was performed (rabbit anti-Sodium Potassium ATPase [EP1845Y] Alexa-488 conjugate, Abcam ab197713, 1:500, were diluted in FAD) for 1 hour at room temperature, followed by a TBST wash. Finally, slides were incubated with DAPI (4′,6-Diamidino-2-Phenylindole, Dihydrochloride (5 mg/mL stock in DMF, Thermo D1306) at 0.25 μg/mL (diluted in TBST) for 10 minutes at room temperature. Slides were then washed with a series of TBST, followed by TBS and then distilled H₂O. Excess liquid was removed and slides were mounted with ProLong Diamond Gold (Life Tech P36930) and allowed to harden overnight. For patient samples, a TSA-amplification of AR signal was performed. Briefly, following antigen retrieval/tissue tracing with PAP pen, performed as above, slides were incubated with treated with Bloxall to remove endogenous peroxidase (Vector Labs SP-6000) for 10 minutes at RT. Slides were then rinsed in water (30 seconds) and TBST (30 seconds). Slides were blocked with 2.5% Normal Horse Serum (Vector Labs MP-7401 Component) for 25 minutes at RT and then block was tapped off. Slides were then blocked with Opal-specific PE/Diluent/Block (AKOYA ARD1001EA) for 10 minutes at RT and then block was tapped off. Primary antibody (rabbit anti-Androgen Receptor [EPR1535(2)], Abcam ab133273, 1:300, diluted in PE Diluent AKOYA ARD1001EA) was incubated 0/N at 4°C, followed by a TBST wash for 3 minutes. Anti-Rabbit secondary HRP polymer was then added to slides (Impress MP-7401) for 25 minutes at RT after which slides were washed in TBST. Opal signal was generated by adding Opal 570 fluorophore (AKOYA FP1488001KT diluted 1:200 in 1x Plus Amplification Diluent AKOYA FP1609) for 10 minutes at RT and then washed in TBST. All following steps e.g. staining with the membrane marker (Abcam ab197713), DAPI, and coverslip mounting were identical to the steps described above.

**IF Image acquisition and Analysis**

Slides were imaged on a Vectra 3 using the A UPlanSApo 10x/0.40 air objective first. Images were acquired using all available channels with the Vectra software (3.0.5) and the raw data was saved as
Regions of interest (ROIs) were created using PhenoChart (1.0.10) and these areas were then imaged again at 20X magnification on the Vectra. 20x images were then spectrally unmixed using inForm software and saved as Component TIFFs. The patient samples were imaged on a Vectra Polaris at 10x and saved as “.qptiff” files. Files were opened in QuPath software, channels were split, and saved individually (or merged) as TIFFs. An APP was created in Visiopharm to segment cells and assess intensity of AR immunofluorescence. Percent positive cells was calculated for each sample. Raw analyzed data was exported as a.csv file and graphed/statistics run using GraphPad PRISM 8 software. A t-test test was performed to test significance.

Methods References

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**Declarations**

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**AUTHOR CONTRIBUTIONS**

CPV, MGW, MCA, JW – formulated research goals and aims
CPV, MGW, MCA, JW – developed and designed methodology

MGW, MCA, HZ – designed and executed pipelines and analytic code

CPB, MGW, MCA, HZ, GM – conducted statistical analysis, built models and plotted results CPV, JRM, JM – designed and conducted immunological assays and pre-clinical studies generated and annotated meta data, created cell lines

JK, SJ, NF, GG – in vitro studies

CPV, JRM – scientific drivers, planned and oversaw all murine experiments and studies

JRD, MM – AR staining

MM – RPPA and quant-seq data

MP – generation of AR knockout cell lines

LF, JRD, JM, JW, prepared the manuscript

TH – Scientific oversight

All authors reviewed and edited the manuscript

JW - supervised and oversaw all aspects of study including design, conduct and analyses

DISCLOSURE

J. Wargo is an inventor on a US patent application (PCT/US17/53.717) submitted by the University of Texas MD Anderson Cancer Center that covers methods to enhance immune checkpoint blockade responses by modulating the microbiome. J. Wargo reports compensation for speaker’s bureau and honoraria from Imedex, Dava Oncology, Omniprex, Illumina, Gilead, PeerView, Physician Education Resource, MedImmune and Bristol-Myers Squibb. J. Wargo serves as a consultant / advisory board member for Roche/Genentech, Novartis, AstraZeneca, GlaxoSmithKline, Bristol-Myers Squibb, Merck, Biothera Pharmaceuticals and Microbiome DX. J. Wargo also receives research support from GlaxoSmithKline, Roche/Genentech, Bristol-Myers Squibb, and Novartis.

CNS is a contributor on a US patent application (PCT/US17/53.717) submitted by the University of Texas MD Anderson Cancer Center that covers methods to enhance immune checkpoint blockade responses by modulating the microbiome.

VG is a co-inventor on a US patent application (PCT/US17/53 717) submitted by The University of Texas MD Anderson Cancer Center that covers methods to enhance checkpoint blockade therapy by the microbiome. V.G. has served as a consultant for Microbiome DX, and reports honoraria from ExpertConnect and Kansas Society of Clinical Oncology.
JLM serves as a consultant for Merck and in advisory board for Bristol-Myers Squibb.

MCA reports advisory board participation and honoraria from Merck Sharp and Dohme, outside the submitted work.

RNA receives research funding from Merck, Bristol-Myers Squibb, Genentech, Novartis, Iovance and is a consult for Iovance and Nektar.

MAD is an advisory board member/consultant for BMS, Novartis, Array, Roche/Genentech, GSK Sanofi-Aventis. PI of research grant to institution from Astrazeneca, Roche/Genentech, GSK, Myriad, Oncothyreon, Sanofi-Aventis.

All other authors report no COI directly relevant to this work.

MATERIALS AND CORRESPONDENCE

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contacts, Timothy Heffernan (TPHeffernan@mdanderson.org), Joseph Marszalek (JRMarszalek@mdanderson.org), or Jennifer A. Wargo (jwargo@mdanderson.org).

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**Figures**
Figure 1

Strong sexual dimorphism noted in response to neoadjuvant BRAF-MEK inhibition therapy in melanoma patients. Sex of the patient influences response to targeted therapy in melanoma. A) Schema of Clinical cohort of patients treated with targeted therapy, B) Major pathologic response with ≤ 10% viable tumor in females and males, C) Recurrence-free survival by sex, D) Differentially expressed genes in pre-treatment specimens between responders and non-responders in the hallmark estrogen and androgen response pathways, E) Androgen receptor staining pre- and post-treatment in males (blue) and females (pink).
Murine model of melanoma validates a sexual dimorphic response and suggests AR activity as a mechanism of resistance

A) Percent change in tumor volume for male and female C57Bl/6 mice implanted subcutaneously with BP (BRAFV600E, PTEN/-/) cells that were treated with Vehicle or DT (30 mpk Dabrafenib and 1 mpk Trametinib, PO, QD). n = 10 mice per group, and the experiment was performed three times (see Supplementary Figure 3B-C). B) Percent change in tumor volume for male and female CD-1 nude mice implanted and treated as in A. n = 10 mice per group. For A and B, p-values were calculated for the final time point using two-sided Student’s t-test.

C) Transcriptome based mRNA expression of Androgen Receptor (AR) from Quant-seq data in female and male mice treated with Vehicle and DT. p-values were calculated using two-sided Student’s t-test.

D) Percentage of cells with AR positive nuclei in BP tumors from male and female BP mice treated with vehicle or DT for 25 days. p-values were calculated using two-sided Student’s t-test.

E) Representative images of AR protein expression by
immunofluorescence across the four groups of mice in D. Blue = DAPI; Yellow = Androgen Receptor. Scale = 100 μm. F) Differential protein expression for several proteins measured using RPPA. p-values comparing vehicle to treated samples were calculated using two-sided Student’s t-test and proteins with p<0.009 are shown.

Figure 3

Modulation of AR activity is associated with differential response to DT A) Percent change in BP tumor growth for female mice treated with vehicle, DT, or DT in combination with testosterone, and B) male mice treated with vehicle, DT, or DT in combination with enzalutamide, physical castration, or testosterone. C) Percent change in BP tumor growth in male mice treated with DT alone or DT in combination with physical castration or castration plus testosterone supplementation. D) Male and E) female mice treated
with DT alone or DT in combination with either enzalutamide or enzalutamide plus testosterone supplementation. (n = 6-10 mice/group). For A-E, p-values were calculated for the final time point using two-sided Student’s t-test. Note that the DT, DT + Castration, and DT + Enzalutamide groups in C and D are the same data that is shown in B. F and G) Level of testosterone (pg/mL) in plasma samples from BP tumor growth experiments in male and female mice, respectively. p-value was calculated using two-sided Student’s t-test. H) Quantification of the percent of AR+ nuclei by immunofluorescence in BP tumors from male and female mice treated for 25 days. p-values were calculated using two-sided Student’s t-test. I) Representative images from samples in H. Blue = DAPI; Yellow = Androgen Receptor. Scale = 100 μm. J-K) Percent change in AR-KO BP tumor growth for female (J) and male (K) mice treated with vehicle or DT in combination with either enzalutamide or testosterone, respectively (n = 10 mice/group).

**Supplementary Files**

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- Suppfig1.pdf
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