Cooperation between Polycomb and androgen receptor during oncogenic transformation

Citation

Published Version
doi:10.1101/gr.131508.111

Permanent link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:14169379

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dashboard.current.terms-of-use#LAA

Share Your Story
The Harvard community has made this article openly available. Please share how this access benefits you. Submit a story.

Accessibility
Cooperation between Polycomb and androgen receptor during oncogenic transformation

Jonathan C. Zhao, Jianjun Yu, Christine Runkle, et al.

Genome Res. 2012 22: 322-331 originally published online December 16, 2011
Access the most recent version at doi:10.1101/gr.131508.111

Supplemental Material

http://genome.cshlp.org/content/suppl/2011/12/16/gr.131508.111.DC1.html

References

This article cites 28 articles, 9 of which can be accessed free at:
http://genome.cshlp.org/content/22/2/322.full.html#ref-list-1

Article cited in:
http://genome.cshlp.org/content/22/2/322.full.html#related-urls

Creative Commons License

This article is distributed exclusively by Cold Spring Harbor Laboratory Press for the first six months after the full-issue publication date (see http://genome.cshlp.org/site/misc/terms.xhtml). After six months, it is available under a Creative Commons License (Attribution-NonCommercial 3.0 Unported License), as described at http://creativecommons.org/licenses/by-nc/3.0/.

Email alerting service

Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here

To subscribe to Genome Research go to:
http://genome.cshlp.org/subscriptions

© 2012, Published by Cold Spring Harbor Laboratory Press
Cooperation between Polycomb and androgen receptor during oncogenic transformation

Jonathan C. Zhao,1 Jianjun Yu,1 Christine Runkle,1 Longtao Wu,1 Ming Hu,2 Dayong Wu,3 Jun S. Liu,2 Qianben Wang,3 Zhaohui S. Qin,4 and Jindan Yu1,5,6

1Division of Hematology/Oncology, Department of Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois 60611, USA; 2Department of Statistics, Harvard University, Cambridge, Massachusetts 02138, USA; 3Department of Molecular and Cellular Biochemistry and the Comprehensive Cancer Center, Ohio State University, Columbus, Ohio 43210, USA; 4Department of Statistics and Bioinformatics, Emory University, Atlanta, Georgia 30322, USA; 5Robert H. Lurie Comprehensive Cancer Center, Northwestern University Feinberg School of Medicine, Chicago, Illinois 60611, USA

Androgen receptor (AR) is a hormone-activated transcription factor that plays important roles in prostate development and function, as well as malignant transformation. The downstream pathways of AR, however, are incompletely understood. AR has been primarily known as a transcriptional activator inducing prostate-specific gene expression. Through integrative analysis of genome-wide AR occupancy and androgen-regulated gene expression, here we report AR as a globally acting transcriptional repressor. This repression is mediated by androgen-responsive elements (ARE) and dictated by Polycomb group protein EZH2 and repressive chromatin remodeling. In embryonic stem cells, AR-repressed genes are occupied by EZH2 and harbor bivalent H3K4me3 and H3K27me3 modifications that are characteristic of differentiation regulators, the silencing of which maintains the undifferentiated state. Concordantly, these genes are silenced in castration-resistant prostate cancer rendering a stem cell–like lack of differentiation and tumor progression. Collectively, our data reveal an unexpected role of AR as a transcriptional repressor inhibiting non-prostatic differentiation and, upon excessive signaling, resulting in cancerous dedifferentiation.

[Supplemental material is available for this article.]

The transcriptional regulation by androgen receptor (AR), stimulated by androgen, is critical for prostate differentiation and development, as well as malignant transformation. Previous studies have extensively shown that AR induces prostate-specific gene expression driving prostatic differentiation during development (Cunha et al. 2004) and leading to oncogenic transformation during cancer (Heinlein and Chang 2004; Lamont and Tindall 2011). The blockage of AR signaling through androgen deprivation has thus been the mainstream treatment of advanced prostate cancer. While almost all metastatic prostate cancers are initially responsive to androgen ablation therapies, in most cases, however, the disease reemerges in a castration-resistant form. Notably, evidence suggests that this castration-resistant prostate cancer (CRPC) remains dependent on the expression and transcriptional activity of AR through hypersensitive AR activation in the milieu of very low androgen (Chen et al. 2004). The AR pathway therefore remains a leading therapeutic target in CRPC. Recent genomic studies have begun to reveal AR-regulated genes or pathways that might contribute to CRPC.

With the advent of expression microarrays, a large number of genes were found to be regulated by androgen (Wang et al. 2007b). Using chromatin immunoprecipitation (ChIP)–based assays, global AR binding events have been gradually mapped first by region-limited DNA microarrays (Massie et al. 2007) and later by genomictiling arrays (Wang et al. 2009) and ChIP-seq assays (Jia et al. 2008; Lin et al. 2009; Yu et al. 2010b; Massie et al. 2011). These studies have provided unprecedented understanding of AR transcriptional regulation at the genome scale. For example, >90% of AR binding events were found at enhancers >10 kb away from the transcription start sites (TSS) of coding genes. AR is able to regulate an anabolic transcriptional network to fuel prostate cancer (Massie et al. 2011) and to induce cell cycle genes specifically in CRPC cells (Wang et al. 2009). While a number of these studies observe androgen-repressed genes, these genes and their functions, however, have been somewhat neglected in the favor of androgen-induced genes. Whether or not they represent direct AR target genes and how their repression contributes to prostate cancer are poorly understood.

Previous studies have shown that androgen-repressed genes may also play important roles in prostate cancer cell growth and metastasis (Prescott et al. 2007). Their reexpression during androgen ablation therapy is thought to contribute to disease regression, and they may become repressed once again in CRPC. Despite this importance, only a few studies have reported AR inhibition of a handful of genes (Grosse et al. 2011), a majority of which, however, suggested indirect mechanisms involving inhibition of co-factor proteins with transcriptional functions such as SP1 (Verras et al. 2007; Liu et al. 2008; Baniwal et al. 2009; Song et al. 2010). Few of them, indeed, suggested direct AR binding to DNA, however, often through an altered DNA binding specificity (Lanzino et al. 2010; Qj et al. 2011). The evidence to support AR as a transcriptional repressor is lacking, and the exact mechanism largely unknown. Systematic analysis of genomic data will be essential to firmly establish AR as a globally acting transcriptional repressor.

In this study, we present evidence that AR directly inhibits a large number of genes by binding to their regulatory elements containing the consensus ARE motifs. Mechanistically, this repression is mediated by the Polycomb group protein EZH2 and subsequently repressive chromatin remodeling. These genes are development regulators functionally involved in cell differenta-
tion and tumor suppression. Importantly, AR-repressed genes are down-regulated in CRPC cells in an androgen-independent manner. AR-mediated transcriptional repression thus may be an overlooked, yet critical, mechanism to prostate cancer progression and castration resistance.

**Results**

**AR directly suppresses a large number of target genes**

To identify genes whose expression levels are regulated by androgen, we performed microarray profiling of the androgen-dependent LNCaP prostate cancer cells treated with 1 nM synthetic androgen R1881 for 0, 3, 6, 12, 24, and 48 h. The expression of each probe across all time points was normalized to 0 h to yield fold changes. The probes with greater than threefold change (either up or down) in at least two time points and greater than fourfold change in at least one time point were defined as “androgen-regulated.” K-mean clustering followed by heatmap view demonstrated three major clusters: probes gradually induced by androgen \( (n=944) \), probes gradually repressed by androgen \( (n=939) \), and a small cluster of probes strongly up-regulated at 12 h but down-regulated at 48 h \( (n=193) \) (Fig. 1A). We defined the first two clusters as “androgen-induced” and “androgen-repressed,” corresponding to a unique set of 442 androgen-induced and 428 androgen-repressed RefSeq genes, respectively (Supplemental Table S1). Androgen thus appears to be able to repress as many genes as it induces. To ensure this bidirectional regulation, we further examined these genes in previously published time-course studies of androgen treatment and confirmed their respective differential regulation by androgen (Supplemental Fig. S1A, Wang et al. 2007a; Massie et al. 2011). We then selected several top repressed genes and confirmed remarkable (>10-fold) down-regulation by androgen in prostate cancer cells (Fig. 1B). In addition, this repression occurred within the physiological range of androgen in a dose-dependent manner (Fig. 1C). To determine whether this androgen repression is mediated by AR, we performed RNA interference of AR in LNCaP cells. We demonstrate substantial reduction of AR and well-known AR-induced genes such as KLK3 (also known as PSA) and TMPRSS2 upon the removal of AR (Fig. 1D). In contrast, AR knockdown led to remarkable derepression of androgen-repressed genes (Fig. 1E). Interestingly, we found that AR overexpression significantly suppressed the repressed genes even in the absence of androgen (Supplemental Fig. S1B, C). In order to conclude that the repressed genes are directly regulated by AR, however, it is important to determine whether AR binds to their respective regulatory regions.

From the expression microarray data, we noticed that a majority of the androgen-induced and -repressed genes start to show corresponding trends of response at 6 h, which become quite apparent at 12 h but do not reach peak response until 24 h after androgen treatment (Fig. 1A). We thus performed ChiP-seq analysis of AR in hormone-deprived LNCaP cells treated with ethanol or androgen for 16 h to ensure capture of maximum AR binding events (Yu et al. 2010b). We assigned all AR binding events to the nearest genes. We observed that 340 (77%) androgen-induced and 296 (69%) of androgen-repressed genes contain at least one AR binding event within their regulatory regions. Because these numbers appeared high relative to other similar studies intersecting binding events and expression changes, we selected a set of androgen-unregulated genes that were detected in the microarray but showed no differential regulation by androgen. Out of 445 genes that...
showed <1.2-fold differences across all time points, only 185 (41%) contain at least one AR binding event. Overall, AR binding frequency on the repressed genes was only marginally ($P = 0.013$, Pearson’s $\chi^2$ test) lower than the induced genes but was significantly ($P = 4.52 \times 10^{-16}$) higher than the unregulated genes (Fig. 2A; Supplemental Fig. S2A). To ensure that this is not an observation limited to our data set, we analyzed AR binding on genes induced or repressed by androgen in a previously published data set (Wang et al. 2007a). We found that >65% of the genes induced or repressed by androgen contain one AR binding event, which is again significantly ($P < 0.001$, Pearson’s $\chi^2$ test) higher than the unregulated genes (Supplemental Fig. S2B). We next examined whether our finding holds true in other independent prostate cancer model systems. Taking a similar approach, we identified genes that are up- or down-regulated by androgen in VCaP cells and analyzed AR binding on these genes using AR ChIP-seq data derived from VCaP cells. Our results confirmed that in VCaP cells there is also significantly more AR binding on androgen-induced ($P < 2.2 \times 10^{-16}$) and -repressed genes ($P = 9.4 \times 10^{-7}$) compared with unregulated control genes (Supplemental Fig. S2C). This strongly indicates that, control genes, a majority of the androgen-repressed genes are also directly targeted by AR.

Next, we investigated the distribution of AR binding sites and found that <5% of the AR binding events, and to a lesser degree (only 2%) for the repressed genes, locate within the promoter regions (between 5 kb upstream and TSS) of target genes (Fig. 2B). This low number of androgen-repressed genes that contains AR binding sites within the promoters might account for the lack of direct AR-repressed genes reported in the literature. With AR binding events around the repressed genes observed, we next attempted to understand how AR is recruited to the repressed genes. To determine whether any specific DNA motif recruits AR to the target genes, we performed de novo motif search. Our results revealed nearly identical ARE motifs present in the binding sites of both AR-induced and -repressed genes (Fig. 2C). However, AR binding peaks of the induced genes contain significantly ($P < 0.001$) more ARE motifs than the repressed ones, which, in turn, harbor significantly ($P < 0.02$) more than the control regions (Supplemental Fig. S2D). Consistent with these findings, the induced genes contain more strongly enriched AR binding events than the repressed ones (Supplemental Fig. S2E). Therefore, ARE motif appears essential in recruiting AR to both the induced and repressed genes, with relatively more AR enrichment on the induced genes. We next attempted to confirm individual candidate AR-repressed genes.

**Figure 2.** AR directly binds regulatory elements of androgen-repressed genes. (A) AR binding density of regions surrounding the highest peak (±5 kb) of genes induced, unregulated, or inhibited by androgen. AR ChIP-seq was performed in LNCaP cells treated with ethanol (EtO) or synthetic androgen R1881 for 16 h. Each row represents one gene, and each column represents the binding intensity in each 100-bp window of the corresponding ChIP-seq experiment. Genes were sorted by the height of their highest AR binding peak in R1881 condition. AR ChIP-seq in EtO condition was ranked according to R1881 condition. (B) Distribution of AR binding sites relative to AR-induced or AR-repressed genes. AR binding sites within the regulatory regions of AR-induced or -repressed genes were further categorized into promoter (within 5 kb upstream of the TSS site), enhancer, exon, and intron regions. The most conserved DNA sequence motifs were found across the AR binding sites of AR-induced and -repressed genes. De novo motif search was performed using MDscan of the top 500 AR binding sites associated with either AR-induced or -repressed genes. (C) ChIP-seq AR binding peaks on the regulatory elements of candidate AR-repressed genes. Genomic PCR primers (indicated by red arrows) were designed to flank the binding peaks. (D) AR directly binds to the regulatory elements of candidate AR-repressed genes in LNCaP cells. AR ChIP was done in LNCaP cells treated with EthO or R1881 for 16 h. ChIP-PCR was performed using primers specific to each candidate gene (red arrows in D), KLK3, and a negative control, KIAA0066. Error bars: $n = 3$, mean ± SEM. (E) AR directly binds to the regulatory elements of candidate AR-repressed genes in LNCaP cells. AR ChIP was done in LNCaP cells treated with EthO or R1881 for 16 h. ChIP-PCR was performed using primers specific to each candidate gene (red arrows in D), KLK3, and a negative control, KIAA0066. Error bars: $n = 3$, mean ± SEM.
Using primers flanking the ChIP-seq AR binding sites (Fig. 2D), we validated AR occupancy on the regulatory elements of selected AR-repressed genes in prostate cancer cells (Fig. 2E). To demonstrate the pathological relevance, we analyzed two human prostate cancer tissues and confirmed AR binding on the repressed genes in primary tissues (Fig. 2F).

**AR transcriprional repression is dictated by the Polycomb group protein EZH2**

It is intriguing that AR recruitment to some genes turns on expression, while to others it turns off expression. It has been clearly demonstrated that AR is recruited to the induced genes by FOXA1 and active H3K4me1 and H3K4me2 histone modifications (Lupien et al. 2008). To determine whether the same mechanisms are involved in AR recruitment to the repressed genes, we analyzed ChIP-seq data of AR, FOXA1, H3K4 methylation, and H3 acetylation. Because AR primarily binds enhancer regions, we examined ChIP-seq read density within 50 kb upstream to 5 kb downstream of the TSS of target genes. As previously noted by others (He et al. 2010), H3K4me2 and H3 acetylation were found at both enhancers and promoters, while AR, FOXA1, and H3K4me1 predominantly localize at enhancers. Heatmap view of binding densities revealed clear colocalization of AR, FOXA1, H3K4me1, H3K4me2, and H3 acetylation, supporting their collaborative function (Fig. 3A). However, we found that the ChIP-seq read densities of these open chromatin marks are in general substantially stronger for AR-induced than AR-repressed genes.

To further demonstrate how chromatin remodeling and AR recruitment may be altered by androgen, we compared cofactor binding on representative AR-induced and AR-repressed genes before and after androgen stimulation (Fig. 3B). We found that both the induced and repressed genes are pre-marked by precursor modifications even in the absence of androgen. However, these precursor modifications are significantly more enriched on the induced genes with P-values of 0.04 and 0.01 for H3K4me1 and H3K4me2, respectively. In contrast, H3 acetylation, a marker of actively transcribed genes, was not significantly (P = 0.11) different.

**Figure 3.** Pioneering factors FOXA1 and H3K4 mono- and di-methylation colocalize with AR at target loci. (A) ChIP-seq binding density of regions surrounding the TSS (−50 kb to +5 kb) of AR-induced or -repressed genes. AR ChIP-seq was performed in hormone-starved LNCaP cells treated with ethanol (Ethl) and androgen (R1881). All other ChIP-seq was performed in LNCaP cells grown in regular medium. AR-induced and -repressed genes were each sorted by the distance of the highest AR (R1881) binding peak to the TSS of its corresponding genes; the genes with the highest AR binding peak most upstream of its TSS were ranked on top. Genes that do not contain a peak within −50 to 5 kb of its TSS were ranked at the bottom of each. The P-values indicate differences between the read densities of the two groups of genes. (B) QRT-PCR validation of reduced AR coactivating factor binding on the AR-regulated genes. ChIP of AR, FOXA1, H3K4me2, H3K4me2, and Acetyl H3 were performed in hormone-starved LNCaP cells treated with ethanol (Ethl) or androgen (R1881) for 16 h. PCR quantification was carried out using primers flanking the promoters of KLK3 and TMPRSS2 (negative control for AR and FOXA1 binding) and the enhancers of all genes. Marks with significant differences between AR-induced and -repressed genes include H3K4me1 (ethl), H3K4me2 (ethl), and H3 acetyl (R1881) with P-values equal to 0.04, 0.01, and 0.009, respectively. Error bars: n = 3, mean ± SEM.
suggesting that the differences in H3K4me1 and H3K4me2 are unlikely due to the basal expression activities of the AR-induced and AR-repressed genes prior to androgen stimulation. To further confirm this, we analyzed their expression levels in two time-course androgen-treatment studies (Wang et al. 2007a; Massie et al. 2011). Indeed, we observed no significant differences between basal expression levels of induced and repressed genes before AR activation (Supplemental Fig. S3). The expression levels of the induced genes only become significantly higher than the repressed genes after androgen stimulation. Being consistent with the expression status, ChIP-PCR showed that androgen stimulation successfully recruited AR to enhancers of both AR-induced and AR-repressed genes and led to significantly (P = 0.009) stronger H3 acetylation on the induced genes. Collectively, our data suggest that the preexisting H3K4me1/2 modifications are commonly used to pry open the nucleosomes to allow FOXA1 and AR binding. However, AR is much more likely to induce genes strongly marked by active histone modifications. With the mechanism of AR recruitment to the repressed genes essentially worked out, we next attempted to understand how AR suppresses gene expression by focusing on the promoter regions.

We examined the ChIP-seq read densities of Pol II, H3K4me3, H3K27me3, and H3K36me3 (Fig. 4A). As expected, AR-induced genes are significantly (P < 0.01) enriched for Pol II and H3K4me3 around the TSS and H3K36me3 on the transcribed regions. Intriguingly, H3K27me3 ChIP-seq read density is significantly (P = 0.004) more enriched around AR-repressed genes (Fig. 4A; Supplemental Fig. S4A). Similarly, a higher level of H3K27me3 around AR-repressed genes was also observed in VCaP cells (Supplemental Fig. S4B). By ChIP-PCR, we confirmed stronger enrichment of
H3K4me3 and Pol II on the induced gene promoters (Fig. 4B). In contrast, H3K27me3 is significantly \( (P = 0.05) \) more enriched on the repressed genes, especially after AR activation, supporting repressive chromatin remodeling for gene repression. Because H3K27me3 is enzymatically catalyzed by the Polycomb group protein EZH2, we asked whether EZH2 is involved in AR-mediated gene repression. We carried out ChIP using an anti-EZH2 antibody. Our results demonstrated that EZH2 occupancy on the AR-repressed genes is significantly \( (P < 0.03) \) increased following AR activation (Supplemental Fig. S4C).

To further examine whether EZH2 is required for AR-mediated transcriptional repression, we performed RNA interference to knock down EZH2 in LNCaP cells (Fig. 4C). Importantly, following EZH2 knockdown, three out of the four repressed genes tested were markedly derepressed (Fig. 4C). The gene that failed to respond may be regulated by additional repressive mechanisms such as promoter hypermethylation. To confirm the role of EZH2 on a global scale, we performed microarray profiling of these LNCaP cells treated with control and EZH2 siRNA. We identified a cohort of genes that are differentially expressed upon EZH2 knockdown (Supplemental Table S2). Gene Set Enrichment Analysis (GSEA) demonstrated highly significant \( (FDR = 0.003) \) up-regulation of AR-repressed genes upon EZH2 knockdown (Fig. 4D). Furthermore, HDAC inhibitor SAHA, which is known to reduce EZH2-mediated H3K27me3 (Yu et al. 2007a), derepressed AR-repressed genes alone or in synergy with AR knockdown (Supplemental Fig. S5A). We then asked whether the derepression of AR-repressed genes is due to the loss of EZH2. We established LNCaP cells with stable EZH2 knockdown using shRNA technology (Supplemental Fig. S5B). Using ChIP followed by PCR, we first verified significant enrichment of H3K27me3 \( (P < 0.03) \) and EZH2 \( (P < 0.02) \) on the AR-repressed gene promoters (Supplemental Fig. S5C). We then compared the levels of H3K27me3 and EZH2 in the control cells and the cells with EZH2 knockdown. Our results showed marked reduction of both EZH2 occupancy \( (P < 0.02) \) and H3K27me3 modification \( (P < 0.04) \) on the AR-repressed genes upon EZH2 knockdown (Supplemental Fig. S5D,E). Therefore, Polycomb group protein EZH2 appears to collaborate with AR on target gene repression. We next examined whether this regulation is functionally relevant.

**AR-repressed genes are developmental regulators suppressed by Polycomb in embryonic stem (ES) cells**

We sought to determine the potential function of the AR-repressed genes by using Molecular Concept Map (MCM) analysis of enrichment by predefined molecular concepts/gene sets in the Oncomine database (http://www.oncomine.com). Out of \( \sim 20,000 \) molecular concepts, only 445 (2%) showed significant \( (P < 0.001) \) overlap with AR-repressed genes (Supplemental Table S3). As expected, our AR-repressed genes are also found repressed by androgen stimulation in other in vitro and in vivo studies (Fig. 5A). In addition, many of these genes are down-regulated in aggressive subtypes of a variety of cancers including prostate, breast, and melanoma, suggesting that they may be general tumor-suppressor genes. Importantly, there are significant overlaps between AR-repressed genes and molecular concepts/gene sets relating to Polycomb group (PCG) proteins, such as genes occupied by EZH2, EED, SUZ12, and H3K27me3 in stem cells and genes suppressed in ES cells relative to their differentiated counterparts. Furthermore, MCM analysis showed enrichment of transcription factors such as LEF1, MEIS1, and MEF2 that are involved in development and differentiation. In contrast, AR-induced genes are enriched for prostate-specific gene sets and have no clear correlation with stem cell signatures. These findings pertaining to AR-repressed genes prompted us to further determine whether they may be developmental regulators whose repression maintains undifferentiation in ES cells but contributes to dedifferentiation in cancer.

Characteristic features of developmental regulators in ES cells include PcG protein occupancy and bivalent H3K4me3 and H3K27me3 histone modifications (Lee et al. 2006). We thus obtained a relevant ChIP-seq data set performed in ES cells (Ku et al. 2008) and examined Polycomb occupancy on the promoters of AR-repressed, AR-induced, and AR-unregulated genes. Not surprisingly, the unregulated genes showed no enriched occupancy by PcG proteins EZH2 and Ring1 (Fig. 5B). Rather, they are strongly occupied by H3K4me3 at the promoters and H3K36me3 at the transcribed regions, supporting their being widely expressed housekeeping genes. Interestingly, some of the AR-induced genes are occupied by PcG and contain bivalent histone modifications, being consistent with their roles in mediating prostatic differentiation. Surprisingly, AR-repressed genes demonstrated the strongest enrichment of PcG occupancy \( (P < 0.001) \), linear mixed effect model) and bivalent H3K4me3 and H3K27me3 \( (P < 0.001) \) modifications. Therefore, AR-repressed genes appear to be developmental regulators whose expression must be turned off in ES cells to maintain the undifferentiated, pluripotent state. In other words, the expression of these genes in ES cells may lead to cell differentiation, and their loss in differentiated cells is likely to cause dedifferentiation, a hallmark of cancer.

**AR-repressed genes remain down-regulated in castration-resistant prostate cancer (CRPC) in the milieu of low androgen**

Since AR remains indispensable for CRPC growth, presumably through regulation of target genes, we next sought to dissect out whether AR-repressed or AR-induced genes or both are accountable for growth of LNCaP-abl, an androgen-independent derivative of LNCaP cells. We hypothesized that if the repressed genes are responsible for CRPC growth, they should be repressed in LNCaP-abl without the need of androgen and vice versa for AR-induced genes. We thus compared the expression of these genes between LNCaP and LNCaP-abl cells, which are regularly maintained in androgen-depleted medium. Notably, the repressed genes were expressed in LNCaP-abl at a level similar to LNCaP in the presence of androgen, suggesting their androgen-independent down-regulation in CRPC cells (Fig. 6A). Since AR-repressed genes are likely tumor suppressor genes, their loss may contribute to CRPC cell growth. In contrast, AR-induced genes were barely up-regulated in LNCaP-abl due to the lack of androgen and thus are unlikely to be accountable for androgen-independent LNCaP-abl cell growth (Fig. 6B). However, AR-induced genes may be important for other androgen-independent cells as CRPC is a highly heterogeneous disease. Our data suggest that down-regulation of AR-repressed genes may provide new mechanisms for androgen-independent growth of CRPC cells.

Since AR repression in androgen-responsive cells is facilitated by EZH2, which is often up-regulated in castration-resistant prostate cancer, we next investigated whether the down-regulation of AR-repressed genes in CRPC is due to increased EZH2 regulation. We thus compared the level of Polycomb occupancy on AR-repressed genes between LNCaP-abl and LNCaP cells in the presence and absence of androgen. Concordant with our earlier data, we showed that androgen stimulation of LNCaP cells increases H3K27me3 on the
repressed genes (Fig. 6C). Remarkably, the level of H3K27me3 modification is significantly stronger in LNCaP-abl compared with LNCaP cells grown in the same androgen-depleted environment. In fact, H3K27me3 level on some genes is even markedly higher than stimulated LNCaP cells. These genes tend to be among the best down-regulated in LNCaP-abl. Our data thus suggest that AR-repressed genes are silenced in CRPC cells in an androgen-independent manner. To confirm this observation on a global scale, we performed microarray analysis of basal gene expression in LNCaP and LNCaP-abl cells in the absence of androgen. GSEA analysis indeed revealed that, while AR-induced genes showed no differential expression, AR-repressed genes are significantly down-regulated in LNCaP-abl cells compared with LNCaP (Fig. 6D).

Discussion
This is the first study to investigate the regulation and function of AR as a transcriptional repressor on the global scale. For decades, AR has been primarily considered as a transcriptional activator that induces prostate-specific gene expression. Although androgen stimulation has been shown to also repress genes, these genes are presumed to be indirectly regulated by AR. Little is known about the overall function of AR-repressed genes in prostate cancer. Taking a systems approach, this study establishes AR as a globally acting transcriptional repressor targeting as many genes as it induces. The pioneering factors and ARE motifs used by the AR-induced genes are also involved in recruiting AR to the repressed genes. Once bound to target loci, the repressive function of AR, however, is dictated, at least in part, by the Polycomb group protein EZH2 and consequent H3K27 trimethylation. EZH2 knockdown sufficiently derepresses AR-repressed genes. Furthermore, we provide evidence that, being concordant with the central role of EZH2 in ES cell maintenance, AR-repressed genes are essentially developmental regulators whose expression would otherwise lead to cell differentiation. Our results thus reveal an unexpected role of AR in suppressing cell differentiation (toward non-prostatic lineages). Therefore, AR, previously known as a lineage-specific transcription factor, may play dual roles; it promotes prostatic differentiation via inducing some genes, while simultaneously inhibiting non-prostatic differentiation by suppressing other developmental regulators.

This newly characterized role of AR may be important for normal prostate development and function, as well as malignant transformation. During development, while AR induces prostate-specific gene expression, its suppression of developmental regulators of other lineages is equally important to ensure concomitant silencing of non-prostatic pathways. In mature prostate cells, this newly characterized function of AR may work in concert with the AR-induced genes in stabilizing the prostatic cell type by suppressing potential trans-differentiation. During prostate tumorigenesis,
Figure 6. AR repression is maintained by Polycomb in castration-resistant prostate cancer. (A) QRT-PCR analysis of AR-repressed and -induced genes in androgen-sensitive LNCaP and castration-resistant LNCaP-abl cells. LNCaP cells were hormone-starved for 72 h before treatment with ethanol (E) or androgen (R) for 48 h. LNCaP-abl cells were maintained in a hormone-free medium. Target gene expression was first normalized to GAPDH and then to its level at LNCaP treated with ethanol. Error bars: n = 3, mean ± SEM. (B) QRT-PCR analysis of AR-induced genes in androgen-sensitive LNCaP and castration-resistant LNCaP-abl cells. (C) H3K27me3 modification on AR-repressed genes. LNCaP cells were hormone-starved for 72 h and treated with ethanol (E) or androgen (R) for 16 h before ChIP experiments. LNCaP-abl cells were maintained in a hormone-free medium. Error bars: n = 3, mean ± SEM. (D) GSEA showing enrichment of AR-repressed genes with genes down-regulated in LNCaP-abl compared with LNCaP. Gene expression in LNCaP and LNCaP-abl under an androgen-depleted environment was determined. Differentially expressed genes were ranked from up-regulated in LNCaP-abl to down-regulated in LNCaP-abl relative to LNCaP.

Methods

Cell lines and treatments

The prostate cancer cell lines LNCaP were obtained from ATCC and grown in RPMI supplemented with 10% FBS (Invitrogen). LNCaP-abl cells were kindly provided by Dr. Qianben Wang (OSU) and were grown in phenol-free RPMI (Invitrogen) supplemented with 10% charcoal-stripped FCS (Invitrogen). For androgen treatment, cells were hormone-starved for 3 d before being treated with 1 nM R1881 for 48 h (for expression assay) or 10 nM R1881 for 16 h (for ChIP). For RNA interference assays, cells were treated with non-targeting siRNA (D-00110-01, Dharmacon), or siRNAs specific to EZH2 (P-002079-01, Dharmacon) or AR (L-003400, Dharmacon). AR overexpression constructs were a gift from Dr. Stephen Plymate (University of Washington). All transfections were performed using Lipofectamine 2000 (Invitrogen).

ChIP and ChIP-seq assays

ChIP and ChIP-seq were performed as previously described (Yu et al. 2007a, 2010b). The data set is available at the GEO database as GSE14092. The ChIP antibodies used include AR (#06-680, Millipore), EZH2 (#17-662, Millipore), H3K4me3 (ab8580, Abcam), H3K27me3 (#07-449, Millipore), Acetyl-H3 (ab10812, Abcam), FOXA1 (ab23738, Abcam), SUZ12 (ab12073, Abcam), and RNA Pol II (ab817, Abcam).

ChIP-seq data analysis

ChIP-seq data were analyzed using HPeak (Qin et al. 2010) to generate window counts and visualized using TreeView. The ChIP-seq data set in embryonic stem cells were downloaded from GEO (GSE13084) (Ku et al. 2008). MCM analysis was performed using a query gene list to search for all concepts available in the OncoMine database as previously described (Yu et al. 2007b). Concepts with significant enrichment by the query concept were exported into a supplemental table. Representative concepts were selectively shown as a network in a figure.

The de novo motif search was performed using MDscan (Liu et al. 2002). AR binding sites associated with AR-induced and...
Gene expression microarrays

Total RNA was extracted using an RNeasy Kit (QiAGEN) and confirmed for RNA integrity. Microarray analysis was performed using HumanHT-12 v 4.0 Expression BeadChip (Illumina). The bead-level data were preprocessed using GenomeStudio (Illumina), and the expression data were analyzed using the beadarray package in Bioconductor, log_{2}-transformed, and quantile-normalized using the beadarray package. Enrichment of gene sets was analyzed using the Gene Set Enrichment Analysis (GSEA) tool (Broad Institute). All gene expression data have been uploaded to the GEO database.

Quantitative PCR

QRT-PCR or quantitative ChIP-PCR was performed using GoTaq qPCR Master Mix 2x (Promega) on an Applied Biosystems StepOne Plus Real Time PCR System as previously described (Yu et al. 2007b, 2010a).

Western blotting

After washing twice with cold PBS, cells were lysed and sonicated briefly in NP-40 buffer (Invitrogen) containing protease inhibitors (Roche Applied Science). Frozen tissue samples were homogenized in NP-40 buffer containing protease inhibitors. All lysates were centrifuged to remove cellular debris, and the supernatants were used for Western blotting. The antibodies used for Western blotting were anti-GAPDH (monoclonal, Abcam), anti-AR (polyclonal PG21, Millipore), and anti-EZH2 (BD).

Data access

New high-throughput data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE1978.

Acknowledgments

We thank Dr. Stephen Plymate for the AR overexpression constructs and Drs. Timothy Kuzel, Chung Lee, Raymond Bergan, and Robin Leikin for helpful discussions. We are grateful to Dr. Arul M. Chinnaiyan for his support. This work was supported by the NIH Robin Leikin for helpful discussions. We are grateful to Dr. Arul M. Chinnaiyan for his support. This work was supported by the NIH R01CA055689 (to J.Y.) and R01HG005119 (to Z.Q.), the U.S. Department of Defense and Dr. Timothy Kuzel, Chung Lee, Raymond Bergan, and Robin Leikin for helpful discussions. We are grateful to Dr. Arul M. Chinnaiyan for his support. This work was supported by the NIH R01CA055689 (to J.Y.) and R01HG005119 (to Z.Q.), the U.S. Department of Defense and

References


Weibbe DS, Xu J, Chen Y, Borsu L, Scher HI, Rosen N, Sawyers CL. 2009. Histone deacetylases are required for androgen receptor function in...

AR directly inhibits non-prostatic differentiation


Received September 2, 2011; accepted in revised form November 29, 2011.