Histone 2B-GFP Label-Retaining Prostate Luminal Cells Possess Progenitor Cell Properties and Are Intrinsically Resistant to Castration

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SUMMARY

The existence of slow-cycling luminal cells in the prostate has been suggested, but their identity and functional properties remain unknown. Using a bigenic mouse model to earmark, isolate, and characterize the quiescent stem-like cells, we identify a label-retaining cell (LRC) population in the luminal cell layer as luminal progenitors. Molecular and biological characterizations show that these luminal LRCs are significantly enriched in the mouse proximal prostate, exhibit relative dormancy, display bipotency in both in vitro and in vivo assays, and express a stem/progenitor gene signature with resemblance to aggressive prostate cancer. Importantly, these LRCs, compared with bulk luminal cells, maintain a lower level of androgen receptor (AR) expression and are less androgen dependent and also castration resistant in vivo. Finally, analysis of phenotypic markers reveals heterogeneity within the luminal progenitor cell pool. Our study establishes luminal LRCs as progenitors that may serve as a cellular origin for castration-resistant prostate cancer.

INTRODUCTION

The prostatic gland contains basal and luminal epithelial cells, together with rare neuroendocrine cells. Luminal cells express cytokeratin (CK) 18 and androgen receptor (AR) and are androgen dependent, whereas basal cells express CK5 and stem cell (SC) transcription factor p63 and are androgen independent (Shen and Abate-Shen, 2010). Developmentally, the murine prostate originates from an ancestral p63+AR+ basal SC population (Pignon et al., 2013). Studies using prostate regeneration assays demonstrate that multipotent SCs capable of differentiating into all prostatic cell types are localized in the basal layer (Kwon and Xin, 2014). Therefore, basal cells are thought to represent the main pool of prostate stem cells (PSCs) (Wang et al., 2013). However, lineage-tracing studies indicate that basal cells rarely generate luminal cells during adult tissue homeostasis but display plasticity under the inductive influence of embryonic urogenital mesenchyme (UGM) cells in tissue regeneration assays, acquiring facultative stem/progenitor cell properties and generating luminal cells (Choi et al., 2012; Wang et al., 2013). Consequently, both cell layers in adult murine prostate are self-sustained by lineage-restricted stem/progenitor cells (Choi et al., 2012), although more primitive SCs reside in the basal layer (Ousset et al., 2012). An in vitro organoid assay recently identified a small fraction (~1%) of luminal cells functionally defined as multipotent luminal progenitors in that they were able to generate organoids containing both basal and luminal cells (Karthaus et al., 2014). Beyond homeostasis, several rare luminal progenitor populations have been reported in regressed mouse prostates, including castration-resistant NKX3.1-expressing (CARN) (Wang et al., 2009), SCA-1+ (Kwon et al., 2016), and castration-resistant BMI1-expressing (CARB) (Yoo et al., 2016) cells. The precise relationship between these luminal progenitor cell populations remains unclear.

The prostate has been a model for studying tissue SCs, because it undergoes atrophy upon castration and regeneration upon re-administration of androgen, and this regression-regeneration cycle can be repeated multiple times. Somatic SCs are generally dormant and this cardinal slow-cycling feature is frequently utilized to identify putative SCs by labels that become diluted as a result of cell division (Tang, 2012). Studies have shown that label-retaining cells (LRCs) in many organs are enriched for SCs (dos Santos et al., 2013; Foudi et al., 2009; Szotek et al., 2009), SCA-1+ (Kwon et al., 2016), and castration-resistant BMI1-expressing (CARB) (Yoo et al., 2016) cells. The precise relationship between these luminal progenitor cell populations remains unclear.

Previously, 5-bromodeoxyuridine (BrdU) was employed to perform pulse-chase experiments to identify candidate SCs. In the prostate, a long-term chased BrdU+ cell population, encompassing both basal and luminal cells, which resides in the proximal region of mouse prostatic ducts and exhibits attributes of epithelial SCs was proposed as PSCs (Tsujimura et al., 2002). Whether these dormant cells truly represent SCs has not been answered mainly due to the technical infeasibility of purifying out live BrdU+ cells for functional studies. More recently, cell surface markers coupled with fluorescence-activated cell sorting (FACS) have been used to dissect the subsets...
of cells within a bulk population. These assays depend on known SC markers, and, notably, the majority of widely used markers (e.g., SCA-1, CD49f) preferentially identify prostate basal stem-like cells (Lawson et al., 2007; Lukacs et al., 2010a; Stoyanova et al., 2012; Xin et al., 2005), leaving the luminal cell compartment understudied. Lineage-tracing technology has greatly enhanced our understanding of SC development; however, lineage-tracing studies only suggest that a certain cell population harbors SCs, but could not pinpoint which precise cell(s) within the population is SC (Rycaj and Tang, 2015).

In this study, we employed a bigenic mouse model to identify, isolate, and characterize the stem-like properties and gene expression profiles of quiescent LRCs from mouse prostates expressing a tunable H2B-GFP driven by the promoter of a luminal lineage-preferential gene probasin (Suraneni et al., 2010). Biological and molecular studies show that long-term chased luminal LRCs are inherently resistant to castration and can generate organoids in vitro and prostatic glands in vivo. Notably, the LRC-derived organoids and prostatic glands contain both basal and luminal cells, suggesting the bipotency of LRCs. RNA sequencing (RNA-seq) analysis of this rare population reveals a progenitor-like transcriptome that resembles therapy-resistant prostate cancer (PCa). Analysis of a spectrum of phenotypic markers previously linked to epithelial cell stemness reveals heterogeneity in luminal progenitor populations. Collectively, our study augments our understanding of prostate luminal progenitors and provides valuable insights into their potential roles in prostate development and, possibly, cancer initiation and progression.

RESULTS

Establishing a Bigenic Mouse Model to Label Slow-Cycling Cells in Prostatic Epithelium

To study the dormant cell population in the luminal cell compartment, we adopted a Tet-Off system similar to that described previously (Tumbar et al., 2004) by engineering a Pb-tetVP16 transgenic line to express tetracycline repressor-VP16 controlled by probasin (Pbsn)-based ARR2Pb promoter (Zhang et al., 2000; Figure S1A). By crossing the Pb-tetVP16 mice with the tetracycline-responsive element-regulated mCMV/H2B-GFP reporter mice (Tumbar et al., 2004), we generated the bigenic mice, Pb-tetVP16-GFP, in which GFP expression is ultimately driven by Pbsn promoter (Figures S1A and S1B). In this way, without doxycycline (DOX) administration (pulse), the prostate tissues would be largely GFP+. Upon DOX administration (chase), the prostate will gradually lose the GFP signal due to cell division, while infrequently cycling and dormant cells would retain GFP for an extended period of time (Figure S1A). Indeed, the whole prostate or microdissected prostate branches from the unchased young adult (6 weeks) animals were green, and GFP intensity dropped accordingly at different intervals of chase (Figure 1A). These data demonstrate the successful establishment of a bigenic mouse model to fluorescently label slow-cycling cells in the prostatic epithelium.

H2B-GFP Primarily Labels Prostate Luminal Cells

We assessed the expression and distribution of GFP+ cells in the adult prostate glands. Consistent with the reported activity of endogenous Pbsn promoter (Zhang et al., 2000), GFP signal was higher in ventral prostate (VP) and dorsal and lateral prostate (DLP) than that in anterior prostate (AP) (Figure 1B). As a marker of differentiation and androgen action in the mouse prostate, Pbsn is primarily expressed in the luminal epithelial cells. Double immuno-fluorescence (IF) staining of CK5 or CK8 and GFP indicated that, as expected, GFP+ cells were mainly localized to the luminal compartment in unchased prostates (Figure 1C). Consistent with the varying gross GFP intensity in different prostate lobes (Figure 1B), the frequency of GFP+ cells was much higher in VP (79.4%) and DLP (79.4%) than that in AP (43.8%) (Figures 1C and S1C; Table S1). We also observed some GFP+CK5+ double-positive cells (Figure 1C, solid arrows) and non-epithelial GFP+ cells in the stromal compartment (Figure S1C), suggesting that the Pbsn promoter is also active in a small subset of basal and stromal cells, as reported previously (Valdez et al., 2012; Wang et al., 2006). Interestingly, during the course of this work and consistent with a prior study (Wang et al., 2014a), we frequently noticed many GFP+ luminal cells in the lumen (Figure 1C, dashed arrows), implying that luminal cells have a faster turnover than basal cells (see below).

Kinetics of H2B-GFP LRCs in Basal and Luminal Cell Layers

To characterize optimal chase time and kinetic changes in GFP+ cells, we quantified the percentage of GFP+ cells at different time points after chase according to their lineage identity (Figures 1C–1E and 2A). We started chasing the male mice at 6–8 weeks when the prostate is generally well developed. The initial luminal cell-labeling efficacy was 79.4%, 79.4%, and 43.8% for VP, DLP, and AP, respectively (Table S1). As expected, upon chase, prostatic cells gradually lost GFP signal (Figures 2B and S2A). By 9-week chase, the percentage of GFP+ luminal cells dropped from ~80% to <20% in VP, and by 14-week chase and after, GFP+ cells were rarely seen (Figures 2C and S2A). After a 12-week chase, the luminal GFP+ cells remained at 2%–6% in different lobes (Figure 2C). Consequently, we referred to the GFP+ cells that persisted for at least a 12-week chase as LRCs. This time point is in line with studies in other
organs using similar H2B-GFP mouse models (dos Santos et al., 2013; Szotek et al., 2008; Wang et al., 2012). Our model also labeled a small subset of CK5+ basal cells, with 2%–8% GFP+ cells in unchased lobes and 1%–3% GFP+ cells after a 12-week chase (Figure 2D; Table S1). Interestingly, compared with the relatively flat curve of basal GFP+ cell kinetics, luminal GFP+ cells decreased much faster as a function of chase (Figures 2C and 2D). IF analysis revealed Ki-67+ cells exclusively in luminal layer in all lobes (Figure 2E). Furthermore, we frequently and exclusively observed symmetrically dividing GFP+ cells in the luminal layer (Figure 2F). These observations validated faster proliferation and loss of GFP signal in luminal cells compared with basal cells. Quantitatively, unchased young adult bigenic mice displayed a small percentage of prostatic GFP+ cells that co-expressed the proliferation marker Ki-67 (4.03%, 0.42%, and 1.80% for AP, VP, and DLP, respectively); in contrast, after a 12-week chase no GFP+ cells were stained positive for Ki-67 (Figures 2G and S2B), confirming the quiescence of LRCs.

Luminal LRCs Are Enriched in the Proximal Prostatic Ducts and Mark Progenitors Capable of Regenerating Prostate Glands In Vivo
We next performed in vitro and in vivo SC-related assays, first in bulk GFP+ LRCs. The lineage-negative (Lin−) GFP+ cells freshly purified from the prostates chased for 12 weeks (Figure S3A) displayed higher capacity to form 2D colonies...
and 3D spheres with larger size (Figures 3A–3C) than matched GFP− cells. In vivo tissue regeneration assays demonstrated that both GFP+ and GFP− cells could readily regenerate prostate glandular structures with secretions in the lumen (Figure 3D). The origin of the recombinants was verified by GFP signal and staining (Figures 3D and S3B). It is worth noting that, after a 12-week chase, the LRCs were composed of both luminal and basal cells (ratio = 2:1; Table S1). As basal cells represent the main PSC pool, it is expected that GFP− population, which contained the majority of both luminal and basal cells, would also efficiently regenerate prostate tissues in vivo. Regardless, our

Figure 2. Dynamics and Characterization of LRCs
(A) Scheme for tracking the dynamics of prostatic GFP+ cells in hormonally intact mice, in which we started chasing the bigenic animals at 6 weeks of age and analyzed the prostate tissues at different time points post chase. (B) IF images of CK8 and GFP in the VP isolated from bigenic mice chased for different time intervals. (C and D) Quantification of GFP+ cells in luminal (C) and basal (D) cell populations as a function of chase time. At each time point, two to four mice were examined, and data are shown as the mean ± SE. (E) IF staining of Ki67 and CK8 in different prostate lobes in wild-type adult mice. Arrows indicate Ki67+/CK8− cells. (F) IF of GFP and CK8 in the AP of unchased adult bigenic mice showing symmetrical division of luminal cells. (G) IF staining of Ki67 and GFP in the VP of bigenic mice chased for 12 weeks. See also Figure S2 and Table S1.
data indicate that the long-term chased LRC population harbors stem/progenitor cells with tissue regeneration ability in vivo.

To “zoom in” on the biological differences between luminal and basal LRCs, we purified GFP+ cells from the two cell compartments using marker staining and FACS (Valdez et al., 2012) to identify basal (SCA-1+CD49fhi), luminal (SCA-1−/CD49f/c0), and stromal (SCA-1+CD49f/c0) cells (Figure S4A). Since the Pbsn promoter is active in only a small subset of basal cells (Valdez et al., 2012; Wang et al., 2006), we considered our Pb-tetVP16-GFP bigenic model non-optimal to characterize basal SCs. In support, basal LRCs exhibited limited advantages over basal non-LRCs in SC-related assays, exhibiting slightly increased 2D clonal capacity and nearly equivalent sphere-forming cell frequency (Figures S4B and S4C), although GFP+ basal cells did form larger 3D spheres than GFP− basal cells (Figure S4D). Overall, these results could be explained by low labeling efficiency in the basal cell layer and high SC activity of stochastic basal cells. Therefore, for the remainder of this study we focused on luminal LRCs.

Since our initial luminal cell-labeling efficacy was ~80% for VP and DLP, we first tested whether luminal progenitor cell activity was equally distributed between the labeled and unlabeled (~20%) luminal populations in unchased adult animals. Both 2D clonal (Figure 4E) and 3D organoid (Figure 4F) assays revealed similar activities in unchased luminal (L)-GFP+ and L-GFP− cells, validating the suitability of our model in comparing stem/progenitor cell activities in chased L-GFP+ and L-GFP− cells. Freshly purified luminal LRCs from 12-week-chased prostates exhibited significantly higher clonal and organoid-forming (Figures 4A and 4B) capacities than matched luminal GFP− cells. Importantly, luminal GFP+ cells formed large hollow organoids, whereas only a limited number of luminal GFP− cells formed small compact spheres (Figure 4C). Importantly, luminal LRC-derived organoids had a glandular structure reminiscent of normal prostate, with p63+ and CK5+ basal cells mainly residing in the peripheral layer (Figure 4G). This result is in line with recent findings that only multipotent luminal progenitors can form organoids in vitro (Karthaus et al., 2014). When we mixed L-GFP+ or L-GFP− cells (4.5 × 10^{3}) with mouse UGM and implanted subcutaneously into NOD-SCID-IL2Rγnull (NSG) mice, only L-GFP+ cells generated prostatic glandular structures with overt secretions in the lumen (Figure 4D), whereas even higher numbers of luminal GFP− cells (up to 18 × 10^{3}) failed to
Figure 4. Luminal LRCs Mark Progenitors Capable of Regenerating Prostate Glands In Vivo

(A–C) Freshly purified Lin− luminal GFP+ LRCs exhibit stem/progenitor activities. Shown are higher colony- (A) and organoid- (B) forming capabilities and larger sphere sizes (C) in L-GFP+ compared with L-GFP− cells. For (B) and (C), data are shown as the mean ± SD derived from technical replicates.

(D–F) Only luminal GFP+, but not GFP− cells, are capable of regenerating prostate tissues in vivo. Shown are H&E staining (D) and IF of CK5 and CK8 (E), and GFP (F) in prostate glands regenerated in vivo from sorted luminal GFP+ cells co-injected with mouse UGM.

(G and H) Enrichment of luminal LRCs in the proximal mouse prostate. Different prostate lobes dissected from bigenic mice at 10.5 weeks of DOX chase were divided longitudinally into two portions (distal and proximal) and subjected to IF of GFP and CK8. Shown are representative images (G) and quantification data (H) from two different mice. Results shown (A)–(F) are representative data of at least two to three independent experiments showing consistent results. For (G) and (H), two bigenic mice were analyzed and data represent the means from cell number counting of five to eight random high-magnification (×20) images of each indicated category.

See also Figure S4 and Table S1.
give rise to prostatic tissue (not shown). These LRC-regenerated glands were normal-appearing with a layer of CK8+ luminal cells lining the lumen and a discontinuous layer of CK5+ basal cells encapsulating the structure (Figure 4E). The identity of injected cells and the LRC origin of recombinants were verified by GFP staining (Figure 4F), and the level of GFP was still detectable after several rounds of cell proliferation during prostate regeneration (not shown). These results together indicate that luminal LRCs mark luminal progenitors with bilineage differentiation capacity in vitro and in vivo.

Previous studies using a BrdU label-retention strategy suggested that putative PSCs (including both basal and luminal cells) are slow cycling and enriched in the proximal region of the prostate (Tsujimura et al., 2002). We sought to examine whether our luminal LRCs are also enriched in the proximal prostate. Each prostate lobe from mice chased for 10.5 weeks was dissected longitudinally into two portions, representing the distal and proximal prostatic ducts. If analysis of GFP and CK8 indicated that the GFP+CK8+ luminal LRCs were significantly enriched in the proximal prostatic ducts adjacent to the urethra (figures 4G and 4H).

Interestingly, when we examined the mRNA and protein level of PBSN in the pelvic urogenital sinus (UGS) of newborn and 1-week-old male mice, we observed that the Pbsn mRNA and PBSN protein were minimally expressed in newborn (d1) UGS but readily detectable in 1-week-old UGS (figures S4H–S4J), suggesting that Pbsn is expressed early on in urogenital tissues and Pbsn promoter activity is not necessarily active only in mature fully differentiated luminal cells. Indeed, we detected GFP+ cells co-expressing CK8 and relatively high levels of PBSN protein in pelvic UGS of 1-week-old bigenic mice (Figure 4K). These results indicate that our LRC labeling system is also suitable in marking luminal progenitor cells at early developmental stages.

Luminal LRCs Display a Luminal Progenitor Gene Expression Signature

We next sought to determine the molecular features of L-GFP+ progenitors prepared from bigenic mice chased for 12 weeks. By performing genome-wide RNA-seq analysis using rRNA-depleted total RNAs (Figure 5A), we obtained an average of ~77 million reads per sample with an average mapping rate of 93% to the reference mouse genome (UCSC version mm10; Figure S5A). MA plot indicated appropriate normalization of our RNA-seq data (Figure S5B). Gene set enrichment analysis (GSEA) of RNA-seq transcriptome revealed exclusive enrichment in L-GFP+ (over L-GFP−) cells of gene signatures related to luminal epithelial progenitors and SCs (Figure 5B, plots 1–4). Significantly, a recently reported signature of an AR-deleted luminal cell population with a stem-like phenotype in the mouse prostate (Xie et al., 2017) was significantly enriched in our luminal LRCs (Figure 5B, plot 5). Furthermore, we recently developed an in vitro 2D culture system that allows enrichment of luminal progenitors from primary human prostate tissues (Zhang et al., 2017). The gene signature of this cultured-enriched human luminal progenitors (WIT-L) was also enriched in luminal LRCs (Figure 5B, plot 3). In contrast, and, as expected, signatures associated with non-progenitor cells (luminal progenitor_down), AR signaling and steroid/androgen hormone metabolism were only enriched in L-GFP+ cells (Figure S5C). It has been suggested that luminal progenitors are less dependent on AR signaling (Agarwal et al., 2015; Kwon et al., 2014). Consistent with an enrichment of AR/androgen signaling in L-GFP− cells (Figure S5C) and with AR+ luminal cell signature (Xie et al., 2017) in L-GFP+ cells (Figure 5B, plot 5), Ar mRNA expression was decreased in L-GFP+ cells (Figure 5C). If analysis of GFP and AR using titrated anti-AR antibody dilutions revealed significant heterogeneity of AR expression within luminal cell population and indicated that ~37% ± 1.6% of GFP+ luminal cells expressed low AR protein (Figures 5D and 5S). Interestingly, CK19, a marker for luminal transit amplifying cells (Hudson et al., 2001; Korsten et al., 2009; Wang et al., 2001), and CD24a, a cell surface marker frequently used to enrich mammary luminal progenitors (dos Santos et al., 2013; Rodilla et al., 2015), were dramatically increased in L-GFP+ cells (Figure 5C). Intriguingly, we observed higher levels of Pbsn in L-GFP+ versus L-GFP− cells (Figure 5C), perhaps indicative of the luminal nature of LRCs and also due to LRC identification by virtue of Pbsn promoter activities. Real-time qPCR analysis in two independently sorted biological samples confirmed slightly reduced Ar mRNA levels in L-GFP+ cells and also validated the differentially expressed patterns of about a dozen genes in L-GFP+ versus L-GFP− cells (Figure S5E; data not shown). Intriguingly, L-GFP+ cells, compared with L-GFP− cells, expressed higher levels of luminal markers (Nkx3.1, Krt8, and Krt18) but lower levels of basal genes (Tip63 and Krt5) (Figure S5E).

To further dissect the transcriptomic profiles in L-GFP+ cells (Table S2), we performed the pathway/network enrichment analysis by Ingenuity Pathway Analysis and DAVID gene ontology annotation. The two approaches generated similar results and revealed neurogenesis, SC and development, inflammation/immunity, and cell adhesion being the top gene categories overrepresented in L-GFP+ cells (Figures 5E and S5F). Interestingly, we have recently shown that normal human prostate basal/SCs (Zhang et al., 2016b) and PSA−ve prostate cancer SCs (Qin et al., 2012) also preferentially express many neurogenesis-related genes. Further GSEA corroborated that,
compared with L-GFP− cells, the slow-cycling L-GFP+ cells were highly enriched in gene networks involved in neural/neuronal development and functions (Figures 5B, plot 6, and S5G). Surprisingly, we observed no obvious enrichment of cell-cycle-related pathways in L-GFP+ cells, and a survey of key cell-cycle regulators revealed a generally
decreased expression of positive cell-cycle genes, while cell-cycle-negative regulators remained unchanged (Figure S8H). These results implicate potential involvement of other mechanisms (e.g., transforming growth factor β signaling [Salm et al., 2005]) in driving the quiescence of LRCs.

Neurogenesis and Inflammation Gene Signatures Link the Gene Expression Profile in Luminal LRCs to Aggressive and Castration-Resistant PCa

Our recent studies (Zhang et al., 2017) linked the gene expression profile in WIT-L human prostate luminal progenitors to aggressive subtypes of PCa such as CRPC (castration-resistant PCa) and neuroendocrine PCa. Further, attenuated AR activity and androgen deprivation therapy (ADT) promote a stem-like cell phenotype in PCa (Schroeder et al., 2014) and inflammation has been shown to expand a progenitor-like luminal cell pool (Liu et al., 2016) and decrease AR signaling in human prostatic luminal cells (Zhang et al., 2016a). These discussions prompted us to determine whether the gene expression profile in luminal LRCs could be linked to clinical features of PCa. GSEA revealed a significant enrichment of gene signatures of WIT-L human luminal progenitors in L-GFP+ cells (Figure 5B, plot 3), suggesting that the luminal LRC gene profile may associate with aggressive PCa phenotypes. In support, gene signatures in patient CRPC after failure of ADT were enriched in L-GFP+ cells (Figures 5F and S5I). These latter observations may suggest a global progenitor-like gene expression profile in human CRPC.

Of interest, a gene signature of mouse prostate tumors originated from basal cells (Wang et al., 2013) was modestly enriched in L-GFP+ cells (Figure SSJ), suggesting that prostate tumors with a basal cell origin may be maintained by luminal progenitor-like cells, consistent with an earlier study (Stoyanova et al., 2013). Importantly, a recently reported gene signature of a human CD38low prostatic luminal progenitor population was dramatically enriched in our L-GFP+ cells (Figure 5G). Of note, the CD38low human luminal progenitors are tightly associated with inflammation in PCa, express an inflammatory signature with reduced AR and androgen signaling, and associate with disease progression and poor outcome (Liu et al., 2016). Thus, our mouse luminal LRCs share molecular features with human prostatic luminal progenitors.

Luminal LRCs Resist Castration In Vivo

The cellular origin(s) of castration resistance is a central question in PCa research and a long-standing hypothesis is that stem/progenitor cells may preferentially survive ADT. Consequently, we evaluated the in vivo castration sensitivity of luminal LRCs. To first assess whether our model could label luminal cells intrinsically resistant to castration, we castrated the unchased Pb-tetVP16-GFP male mice at 6–8 weeks and then started DOX chase and examined the abundance of GFP+ cells in prostate lobes at different time points after castration. Numerous GFP+ cells (8.0%, 29.3%, and 20.1% for AP, VP, and DLP, respectively) were observed at even 12 weeks after castration (Figure S6A), suggesting that our model marks a proportion of luminal cells that are intrinsically castration resistant in physiological conditions. We next sought to evaluate whether long-term chased LRCs resist castration. We castrated 10- to 12-week-chased bigenic male mice and analyzed prostate 1–4 weeks later (Figures 6A–6D). As short as 1 week after castration, seminal vesicles and prostate glands shrank significantly (Figures 6A and 6C), suggesting successful androgen ablation. In all experimental settings, the frequency of luminal LRCs (GFP+CK8+) dramatically increased post castration, indicating that GFP+ cells possessed intrinsic survival advantages over GFP+ cells (Figures 6A–6D). For example, in uncastrated bigenic mice chased for 14 weeks, very few GFP+ cells were seen (Figure S2A); in contrast, abundant GFP+ cells remained in the prostates of mice castrated at 10–12 weeks and harvested at 13- to 14-week chase (Figures 6B–6D). We also examined whether LRCs resist apoptosis after castration by co-IF analysis of GFP and cleaved caspase-3 in the prostate castrated at 12- to 13-week chase and harvested at 3 days, 1 week, and 2 weeks post castration (Figure S6B). Apoptosis in mouse prostatic luminal cells generally peaks at 3–4 days after castration and persists, at declining levels, in the following 1–3 weeks (Kato et al., 2013). Notably, we did not detect any GFP+ LRC cells expressing cleaved caspase-3 in all three experimental settings (Figure S6B), suggesting that castration-induced apoptosis in LRCs is a rare event. To further determine whether the increased number of GFP+ LRCs was due to proliferation besides resistance to apoptosis, we performed co-IF of Ki67 and GFP in the above experimental settings. A small number of Ki67+ epithelial cells were observed in regressed prostate (e.g., 1-week [Figure S6C] or 2-weeks [Figure S6D] post castration), but no GFP+ Ki67+ LRCs were found (Figure S6D), consistent with the dormancy of LRCs. These results collectively demonstrate that luminal LRCs preferentially survive androgen deprivation.

Phenotypic Analysis of LRCs Reveals Heterogeneity in Luminal Progenitor Cells

Several surface markers have been reported for mouse mammary luminal progenitors, including SCA-1 (or LY6A), CD133, CD117, CD14, CD24 (Shehata et al., 2012), NOTCH3 (Lafkas et al., 2013), and NOTCH1 (Rodilla et al., 2015). Other than SCA-1 (Burger et al., 2005), similar phenotypic studies in the mouse prostate under physiological (i.e., homeostatic) conditions are generally lacking.
CK19 was proposed as a prostate luminal progenitor marker in several studies (Hudson et al., 2001; Korsten et al., 2009; Wang et al., 2001) and several SC factors such as BMI1 (Yoo et al., 2016), SOX2 (Kregel et al., 2013), STAT3 (Schroeder et al., 2014), and HOXB9 (de Pijnieux et al., 2001) have been associated with stem-like prostate (cancer) cells under castration condition. We performed double IF staining of GFP and about a dozen of reported mouse prostate/mammary luminal progenitor markers on prostate tissues collected from bigenic mice chased for 12 weeks and correlated with the marker mRNA levels in RNA-seq (Figure 7A). Most of these markers were undetectable in normal uncastrated prostates in both formalin-fixed and paraffin-embedded cryosections (Figure 7A). Consistent with the RNA-seq data (Figures 5C and 7A), L-GFP+ cells expressed relatively abundant CK19 protein compared with L-GFP− cells, although not all CK19high cells were GFP+ (Figures 7B and S7A). A subset of L-GFP+ cells co-expressed SCA-1 at the proximal prostate, and not all SCA-1+ cells were GFP+ (Figure 7B), consistent with a recent report that SCA-1 identifies a distinct subset of androgen-independent luminal progenitors in the proximal prostate (Kwon et al., 2016). The 12-week-chased L-GFP+ cells did not co-localize with BMI1-expressing cells (Figures 7B and S7A), a rare population that mainly resides in the basal cell layer (Lukacs et al., 2010b), although Bmi1+ cells have recently been reported to reside, infrequently, in the luminal layer and mark rare castration-resistant luminal progenitors (Yoo et al., 2016). We detected a relatively low and homogeneous expression of NOTCH1 (Figure 7B), suggesting that, unlike in mammary tissues (Rodilla et al., 2015), NOTCH1 is not a preferential marker for prostate luminal progenitors. The positive staining of BMI1 and SCA-1 in stromal cells, and cleaved caspase-3 in apoptotic cells shed into the lumen after castration, served as positive control for IF analysis (Figure S7B). Together, these data suggest that prostate luminal progenitors are phenotypically, and, likely, functionally (as reflected by heterogeneous AR expression), heterogeneous.

**DISCUSSION**

Recent studies, employing 2D (Zhang et al., 2017) and 3D organoid (Karthaus et al., 2014) in vitro culture systems for human prostate and BrdU pulse-chase (Tsujimura et al., 2002) and lineage-tracing (Choi et al., 2012; Ousset et al., 2012) techniques for murine prostate, have indicated the existence of luminal progenitors within the luminal cell lineage. In this study, we established a bigenic mouse model to specifically label slow-cycling cells in the prostate epithelium, and further defined luminal LRCs as functional progenitors distinct from bulk luminal cells. These luminal LRCs have the following features: (1) they are relatively dormant and, like BrdU-retaining LRCs (Tsujimura et al., 2002), enriched in the proximal prostate; (2) they display...
bipotential differentiation ability in both in vitro organoid and in vivo prostate regeneration assays; (3) they express a progenitor gene signature with preferential expression of neurogenesis- and inflammation-associated genes, and this gene signature is dramatically enriched in treatment-failed PCa; (4) they survive androgen ablation by resisting castration-induced apoptosis and are further enriched in the prostate of castrated mice; and (5) they are phenotypically heterogeneous with only a subset expressing previously reported progenitor markers such as CK19 and SCA-1.

Studying prostatic luminal cells has been challenging due to their low stemness and the current infeasibility of efficient culturing in vitro. Prior studies have mainly employed known SC-related surface markers identified in other tissues to dissect the epithelial biology of the prostate. Currently, there is no well-accepted definition for prostate luminal progenitors (Zhang et al., 2017), and only a few studies have reported putative phenotypic markers of stem-like luminal cells in humans (i.e., CD38low; Liu et al., 2016) and in regressed mouse prostates (e.g., NKX3.1+ [Wang et al., 2009], SCA-1+ [Kwon et al., 2016], and BMI1+ [Yoo et al., 2016]). The slow-cycling luminal progenitor (i.e., L-GFP+) cells, identified here in a marker-independent manner, only partially overlap with a few of...
previously reported progenitor markers. In fact, most of the markers we analyzed, including CD117, CD14, CD133, STAT3, NOTCH3, CD24, HOXB9, and SOX2, seem to be expressed at too low levels to be reliably detected in the normal mouse prostate. Nevertheless, luminal LRCs appear to generally express low NOTCH1 and high CK19 levels, and a subset of LRCs stains positive for SCA-1. Consistent with a recent report (Yoo et al., 2016) showing that BMI1+ luminal cells are rarely seen in normal prostate, our LRCs do not co-localize with BMI1+ cells. These observations raise a question of whether the mouse prostate, under homeostatic conditions, might have distinct luminal progenitor subsets at varying proliferative states, like the hematopoietic system. For example, our LRCs luminal progenitor subsets at varying proliferative states, might have distinct activities and the luminal cell layer is self-sustaining (Wang et al., 2013; Zhang et al., 2016a). Second, inflammation to generate aggressive tumors (Wang et al., 2014b). This raises the possibility of luminal progenitors functioning as preferred cells-of-origin for CRPC. Our current findings that luminal LRCs preferentially survive androgen ablation provide support to this possibility. Third, stem-like luminal cells may function as tumor-propagating cells (Stoyanova et al., 2013), in line with mouse genetic studies (Abou-Kheir et al., 2010; Korsten et al., 2009) and with previous report showing that aggressive breast basal-like cancers actually originate from luminal progenitors rather than from basal SCs (Molyneux et al., 2010). Fourth, while this manuscript was under revision, a study identified a rare castration-resistant luminal progenitor cell population with a Lin−/SCA-1+/CD49fmed phenotype highly enriched in Pten-null prostate tumors (Sackmann Sala et al., 2017). Finally, the luminal progenitor gene expression profile can be linked to advanced and aggressive PCa subtypes (Liu et al., 2016; Sackmann Sala et al., 2017; Zhang et al., 2017; this study). This notion is further strengthened by the molecular resemblance of our luminal LRCs to recently reported human CD38low prostatic luminal progenitors (Liu et al., 2016), which has been shown to be associated with inflammation and PCa progression with poor outcome. In aggregate, our study presents a comprehensive biological and molecular characterization of slow-cycling prostate luminal progenitors. Further studies on the functional roles of luminal LRCs in tumorigenesis, and the molecular mechanisms that regulate their homeostasis and dormancy, may yield new predictive markers and therapeutic targets for aggressive CRPC.

EXPERIMENTAL PROCEDURES

General procedures in producing and propagating transgenic animals have been described previously (Suraneni et al., 2010). All animal work was performed under the University of Texas MD Anderson Cancer Center and Roswell Park Cancer Institute IACUC approved protocols. Immunostaining was performed on either paraffin-embedded or OCT frozen sections. The mouse prostate tissues were enzymatically digested with collagenase IA and Dispase, followed by FACS analysis using the BD Aria (BD Biosciences).

Additional experimental procedures are detailed in the Supplemental Experimental Procedures.

ACCESSION NUMBERS

The accession number for the RNA-seq data reported in this paper is GEO: GSE98760.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and three tables and can be found with this article online at https://doi.org/10.1016/j.stemcr.2017.11.016.
AUTHOR CONTRIBUTIONS

D.Z., C.J., and D.G.T. conceived and designed the study. D.Z., S.G., C.J., and A.T. performed the experiments; D.Z., Y.L., and J.S. performed RNA-seq and bioinformatics analysis. D.Z. and D.G.T. analyzed and interpreted the data. D.Z. and D.G.T. wrote the paper. All authors read and approved the final manuscript.

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Supplemental Information

Histone 2B-GFP Label-Retaining Prostate Luminal Cells Possess Progenitor Cell Properties and Are Intrinsically Resistant to Castration

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Figure S1. Establishment of a bigenic mouse model to label slow-cycling cells (LRCs) in the prostatic epithelium, Related to Figure 1

(A) Strategy to generate Pb-tetVP16-GFP bigenic mouse model (see Supplemental Experimental Procedures for detail).
(B) PCR genotyping of single and bigenic Pb-tetVP16-GFP mice.
(C) IF staining of CK5 and GFP in different prostate lobes dissected from unchased adult bigenic mouse identifying GFP+ basal cells and nonepithelial stromal cells. Boxed regions are enlarged. Scale bar, 100 μm.
**Figure S2. Phenotypic characterization and quiescence of prostate LRCs, Related to Figure 2**

(A) Double IF staining of CK5 or CK8 and GFP in different prostate lobes harvested from bigenic mice chased for 0 - 18 weeks. Arrows and dashed arrows indicate positive and non-specific GFP staining, respectively.

(B) Double IF of Ki67 and GFP in different prostate lobes of bigenic mice chased for 0 week and 12 weeks. Representative low-magnification images (left) and quantification data (right) are shown. Arrows and dashed arrows indicate Ki67+GFP- single positive and Ki67+GFP+ double positive cells, respectively.
Figure S3. LRCs can regenerate prostate glands in vivo, Related to Figure 3
(A) Sorting strategy to isolate lineage-negative GFP⁺ and GFP⁻ cells from disassociated bulk prostatic cells.
(B) IF of GFP in recombinants showing that both LRCs and non-LRCs can regenerate prostate tissues in vivo. Note that as we used a Tet-off system, the GFP⁺ cells isolated from 12-week chased animals (non-LRCs) would regain the GFP expression during the tissue regeneration assays.
Figure S4. Luminal LRCs, but not basal LRCs, exhibit preferential stem/progenitor activities compared to non-LRCs, Related to Figure 4

(A) FACS strategy to isolate GFP⁺ and GFP⁻ epithelial cells form both basal and luminal cell populations.
(B-D) Basal GFP⁺ LRCs demonstrate similar sphere-forming efficiency to basal GFP⁻ non-LRCs. Freshly purified Lin⁻ prostate basal GFP⁺/GFP⁻ cells (B-GFP⁺/B-GFP⁻) from 12 week-chased animals were used in colony formation (B) and limiting dilution sphere (C) assays. Note that B-GFP⁺ cells did generate larger spheres than B-GFP⁻ cells (D). Results shown were representative data of at least 2 independent experiments. The P value was calculated using Student’s t-test. For (C) and (D), data represent mean ± SD of values obtained from one experiment.

(E, F) Freshly purified luminal GFP⁺ and GFP⁻ cells from unchased adult animals exhibit similar stem/progenitor cell activities. Shown are 2D colony formation (E) and 3D organoid (F) assays. Pictures in E (left panel) confirmed the epithelial identify of the cells cultured in the indicated medium. Results shown were representative data of at least 2 independent experiments.

(G) Luminal LRCs isolated from 12 week-chased bigenic mice generated organoids containing both basal and luminal cells. Shown are H&E staining and IF analysis of p63, CK5 and CK8 in organoids.

(H) Semi-quantitative RT-PCR of PbSn mRNA in pelvic urogenital sinus (UGS) isolated from newborn and 1-week old mice.

(I, J) H&E staining and IF analysis of PbSn in pelvic UGS of newborn (I) and 1-week old WT mice (J) showing the minimal and considerable expression of PbSn at protein level in newborn and 1-week old mice, respectively.

(K) Double IF analysis of GFP and PbSn or CK8 in pelvic UGS of 1-week old bigenic mice. Boxed regions are enlarged.
|                | L_GFP-         | L_GFP+         |
|----------------|----------------|----------------|
| total number of fragments | 73869998     | 60259391       |
| fragments with both ends mapped | 64184312     | 69737172       |
| fragments with one end mapped   | 4332527      | 4982840        |
| tags that are not mapped        | 14630277     | 16061598       |
| total mapped rate               | 90%           | 89%            |
Figure S5. Luminal LRCs express unique progenitor cell gene signatures linked to CRPC, Related to Figure 5

(A) RNA-seq statistics.
(B) The MA plot showing appropriate normalization of our RNA-seq data.
(C) GSEA results showing the enrichment of indicated gene signatures in luminal GFP+ cells.
(D) Double IF staining of GFP and AR in prostates isolated from 12-week chased bigenic animals. Solid arrows point to GFP+/ARhi cells and dashed arrows to GFP+/ARlow cells.
(E) qRT-PCR validation of RNA-seq data. Two independently prepared sample pairs (luminal GFP+ and GFP- cells) were used in qRT-PCR analysis of several representative genes in ‘luminal’, ‘basal’, and ‘SC’ categories. Overall, the trend of gene expression was consistent between these two measurements with more dramatic differences detected by qPCR, which is reasonable as qPCR utilized Actb as normalization and tended to be more quantitative while RNA-seq did not use housekeeping gene expression for normalization. Note that both Actb and Gapdh mRNA expression levels were not evenly distributed in the L-GFP+ and L-GFP- cell populations in RNA-seq analysis (right boxed). Gene expression values in L-GFP+ cells detected by qPCR or RNA-seq were arbitrarily set as 1 for the purpose of comparison.
(F) Ingenuity pathway analysis (IPA) of genes preferentially upregulated in luminal GFP+ cells.
(G) GSEA results showing the enrichment of indicated gene signatures in luminal GFP+ cells.
(H) Heatmap of representative cell-cycle regulators in luminal GFP+ and GFP- cells. The relative gene expression values (i.e., fold change, FC) in L-GFP+ over GFP- (arbitrarily set as 1) were used.
(I, J) GSEA results showing the enrichment of indicated gene signatures in luminal GFP+ cells. Note that an FDR < 0.25 is statistically significant for GSEA analysis.
Figure S6. Luminal LRCs mark androgen-independent cells and resist castration-induced apoptosis in vivo, Related to Figure 6

(A) Representative IF images and quantification of CK8 and GFP in different prostate lobes harvested from bigenic male mice castrated at 6-8 weeks and chased for 4, 10 and 12 weeks.

(B) IF of cleaved Caspase 3 and GFP in prostate tissues harvested from castrated mice. Different experimental settings, i.e., mice chased for 12 weeks were castrated and VP lobes harvested 3 days (top) or 2 weeks (bottom) later, or castrated at 13 weeks of chase and harvested 1 week later (middle panels) were employed to show that GFP+ luminal LRCs were apoptosis-negative.

(C) IF of Ki67 in prostate tissues harvested from indicated castrated mice.

(D) IF of GFP and Ki67 in prostate tissues harvested from indicated castrated mice.
Figure S7. Phenotypic heterogeneity of luminal progenitor cells, Related to Figure 7
(A) Double IF staining of GFP and the indicated markers in frozen sections of prostate glands harvested from bigenic mice chased for 12 weeks. In the middle (right panel), the dashed boxed region is enlarged.
(B) Double IF of GFP and Bmi1 (left panel) or Sca-1 (middle panel) in FFPE sections obtained from bigenic mice chased for 12 weeks, and of GFP and cleaved Caspase 3 (CC3) in frozen tissue sections of prostate isolated from adult bigenic mice castrated for 1 week (right panel).
Table S1. Supporting data for quantitation of GFP$^+$ cell properties, Related to Figure 2, 4, 5, 6, S2, S6

Table S2. List of the differentially expressed genes (DEGs) in luminal GFP$^-$ and GFP$^+$ cells (comparative to each other) isolated from 12-week chased bigenic mice in the RNA-Seq analysis, Related to Figure 5 and S5
| Antibody | Supplier | Catalog. No | Species | Dilution |
|----------|----------|-------------|---------|----------|
| AR       | EMD Millipore | 06-680 | Rabbit | 1:750   |
| Ck8      | Developmental Studies Hybridoma Bank | TROMA-1 | Rat | 1:50  |
| Ck5      | Covance | PRB-1609 | Rabbit | 1:500  |
| p63      | Cell Signaling | 4892S | Rabbit | 1:500  |
| Ki67     | Leica Biosystems | NCL-Ki67p | Rabbit | 1:500  |
| Sox2     | Cell Signaling | 14962 | Rabbit | 1:150  |
| GFP      | Sigma | G6539 | Mouse | 1:500  |
| GFP      | Life Technologies | A-11122 | Rabbit | 1:1000  |
| Ck19     | Abcam | ab52625 | Rabbit | 1:200  |
| Notch1   | Abcam | ab52627 | Rabbit | 1:100  |
| Stat3    | Abcam | ab76315 | Rabbit | 1:150  |
| Sca1     | Abcam | ab109211 | Rabbit | 1:100  |
| Cd117/c-Kit | Biolegend | 105801 | Rat | 1:150  |
| Cd14     | Biolegend | 150101 | Rat | 1:150  |
| Cd133    | Biolegend | 141201 | Rat | 1:100  |
| Notch3   | R&D System | AF1308 | Goat | 1:100  |
| Cd24     | Santa cruz | sc-19585 | Mouse | 1:150  |
| Bmi1     | Millipore | 05-637 | Mouse | 1:100  |
| Hoxb9    | Assay Biotech | C10404 | Rabbit | 1:150 |
| Pbsn (M-18) | Santa Cruz | sc-17124 | Goat | 1:100  |
| Casp3-cleaved | Cell Signaling | 9664 | Rabbit | 1:150 |

| Primer Name | Sequence (Forward 5'-3') | Sequence (Reverse 5'-3') |
|-------------|--------------------------|--------------------------|
| mGapdh      | ATCACTGCCACCCCAGAAGAC    | AGGTTTCTCCAGCGGCAC       |
| mActb       | AGCAAGCAGGAGTACGATGAG    | AAGCCATGCAATGTTGTC       |
| mPbsn       | TGCACAGTATGAAGGAGCAT     | GTCCGTGCCATGATCAGCT      |
| mKrt5       | GGCTCTTGGTTGGGTAGCG      | AGGCTCTTCTTAGCTTCTAGAG   |
| mKrt14      | GGAGATGATCGGCAGTGTTGCG   | GAAGATGAAAGTGGGCGT       |
| mTrp63      | GCTGCCTGGAAGAAATGACG     | TTGCTGTTTCTGATGCTGTC     |
| mKrt8       | ACACCTTACGCCCAGCAC       | TCTCCCCGTAGCCCTGA        |
| mKrt18      | GAGACGCCACCCAGCACCT      | GCTCCATCGTCCTGTAGTAC     |
| mAr         | GACCTGCTAATCAAGTCCTCCAT  | ATTAGGGTTTCCAAATCCTAC    |
| mBmi1       | TTATCAGCCATCAGTTTTTGT    | ACAGCAATGTGTGTAAGAATGAT  |
| mBcl2       | CTACCGTCGTAGTCTGCGAG     | ACACACATGACCCACCAGG     |
| mAldh1a1    | TCAACAGCAGTCCGCAATAGAAGAT| AAAACAGCAGCTATGCTGTGTAAGA|
| mAldh3a1    | AGACATCAAGCGGTTGGAAGT    | AGACCTCCAGCGACGAGC      |
| mLy6a       | CAGCAGTTATTGTTGATATCTCA  | CCTCCATTGGGAACGTCCTCAC  |
| mNKX3-1     | TCCACCACCTGCTGCTATCAG    | AACAGGACAGAAGGAAATC     |
| mVIM        | AGCTGCTAATCACCGAGACACTA | CAGAGGAAGTGGACTCCAGGTTAG|
| mSetdb1     | TCAACACACATGGCCTCCGT     | CACCAAGGAGGAGCAGAGA     |
| mKdm1a      | GTATCTCGTGTCGCTGTCG      | GCCCACTACACAGAGCACCAC   |
| mCtnmb1     | CCACCCTGGTGCAGTACATC     | ATTTACAGGTCAGTACAACCCAG |

Note: “m” means mouse.
Supplemental Experimental Procedures

Generation of Prostate-Specific Pb-tetVP16-GFP Transgenic Mice

General procedures in producing and propagating transgenic animals have been previously described (Suraneni et al., 2010). Briefly, we adopted a modified tetracycline-repressible histone H2B-GFP (Tet-off) system previously used to assess slow-cycling stem cells in the skin (Tumbar et al., 2004) to monitor GFP dynamics as readout of cell cycle frequency in prostate epithelial cells. Transgene constructs (Figure S1A) were prepared by placing tetR-VP16 (pTET-Off, BD Biosciences Clontech) under the control of prostate specific and androgen responsive composite probasin promoter ARR2PB (Zhang et al., 2000) with an intervening rabbit b-globin second intron sequence to augment transgene expression and followed by b-globin-SV40 hybrid polyA sequences for transcript stability (Chen et al., 2009). This ARR2PB-tetR-VP16 line were bred with tetO-HIST1H2B/J/GFP (H2B-GFP) mice (stock # 005104, Jackson Laboratory), in which expression of H2B-GFP is under the control of the TET response element (TRE), to generate Pb-tetVP16-GFP bigenic mice (Figure S1A). To analyze the decay of H2B-GFP signal, young adult bigenic male mice were fed DOX chow (2 g/kg body weight, Teklad) starting at 6 weeks and were kept on DOX food for entire chase period (6-18 weeks).

Colony and Sphere Formation Assays

For colony formation assays (Zhang et al., 2011; Zhang et al., 2016), we plated freshly purified prostate epithelial cells at a low density (i.e., 1,000-10,000 cells/well) in 6-well plate precoated with PureCol (Advanced BioMatrix, San Diego, CA), and let cells grow for 7-10 days before the visualization of the culture by crystal violet staining. For sphere-formation assays (Liu et al., 2011), cells were suspended in 1:1 Matrigel (BD Biosciences, San Jose, CA)/medium in a total volume of 100 µl. The mixtures were then plated around the rims of wells in a 12-well plate and allowed to solidify in 37°C incubator for 25 minutes, followed by addition of 1 ml of warm medium. Usually 7-9 days after plating, spheres with a diameter over 50 µm were counted. PrEGM and modified WIT medium capable of propagating prostatic stem-like cells (inducing luminal progenitors) was used for 2D culture (Zhang et al., 2017) whereas both WIT and recently established prostate organoid medium (Karthauser et al., 2014) were used for 3D sphere/organoid cultures. For all experiments, we ran a minimum of triplicate wells for each condition and repeated experiments 2-4 times.

Histology, Immunofluorescence (IF) Staining, and Microscopy

Hematoxylin and eosin (H&E) and IF staining was performed on either 5-µm paraffin-embedded or OCT frozen sections. Basic IF procedures have been described previously (Jeter et al., 2009). The coverslips or the tissue slides were blocked with Background Sniper (Biocare Medical, Concord, CA) for 30 minutes, followed by primary antibody incubation overnight at 4°C. Primary antibodies and dilutions used are listed in Table S3. Slides were then incubated with secondary antibodies (diluted 1:700 in antibody diluent (Dako, Carpinteria, CA)) labeled with AlexaFluor 488 or 594 (Invitrogen/Molecular Probes, Grand Island, NY). After washing (3X) with PBS, sections were counterstained with 4,6-diamidino-2-phenylindole (DAPI) (Sigma-Aldrich, St. Louis, MO) and mounted with ProLong® Gold Antifade Mountant (Life Technologies, Grand Island, NY). For quantification of AR staining intensity in luminal GFP+ cells, a serial dilution of AR antibody from 1:100 to 1:750 was tested and 1:750 dilutions was selected. H&E and IF images were captured by Olympus IX71, and Keyence BZ-X700 and Zeiss LSM510 META confocal microscope, respectively.

Mouse Prostate Tissue Disassociation and Flow Cytometry

Prostate tissues were dissected and minced to small clumps, followed by enzymatic dissociation with collagenase IA (200 U/ml) and Dispase (1 U/ml) in DMEM media with 10% FBS and 10 µM of p160ROCK inhibitor Y-27632 dihydrochloride (Selleckchem, Houston, TX) for 90 min. Dissociated tissue was then treated with 0.2% Trypsin for 10 min followed by 35 min DNase I (0.1 mg/mL) digestion in 37°C. After passing sequentially through 21- and 23- gauge needles, cells were filtered by a 40-µm cell strainer to obtain single-cell suspensions. A mouse lineage cell depletion (order # 130-090-858, Mylenyi Biotech) step was applied according to manufacturer’s instruction. The resultant cells were stained for 30 min at 4°C with PE-CD49f and APC-Sca-1 (eBioscience, San Diego, CA). FACS was performed by using the BD Aria or Fusion (BD Biosciences, San Jose, CA). PI (Propidium iodide) or 7-AAD was added prior to FACS analysis to separate viable from dead cells. As described previously (Valdez et al., 2012), three major populations were defined on the FACS profile as basal (Sca-1+CD49f+), luminal (Sca-1+CD49flo) and stromal (Sca-1+CD49fhi) cells. For the majority of the experiments, GFP+ and GFP- cells were specifically sorted out from the luminal population.

Prostate Tissue Regeneration Assays

Tissue regeneration assays were performed as described (Xin et al., 2003). Briefly, varying number of freshly dissociated prostatic cells or FACS-purified cells were combined with 1.0–2.0 × 10^5 mouse or rat urogenital sinus mesenchyme (UGM)
in collagen and injected subcutaneously or under the kidney capsule. The outgrowths were harvested for analysis from the experimental mice 2–3 months later.

**RNA Isolation and Quantitative RT-PCR (qPCR)**

Total RNA was isolated from cells using the RNeasy mini kit (Qiagen, Valencia, CA). The first-strand cDNA synthesis was achieved by reverse transcription of RNA using random hexamers and SuperScript III Reverse Transcriptase (Invitrogen, Grand Island, NY). Quantitative RT-PCR was performed using the iQ™ SYBR® Green supermix (BioRad, Hercules, CA) on a 7900HT Fast Real-Time PCR System (ABI, Applied Biosystems, Foster City, CA). The primers used in this study are listed in Table S3. The housekeeping gene Gapdh or Actb was used as internal control for gene expression normalization.

**RNA Sequencing (RNA-seq) and Bioinformatics**

For RNA-Seq analysis, freshly purified GFP^+^/GFP^-^ luminal cells were subjected to total RNA extraction using an RNeasy mini kit (Qiagen, Valencia, CA). cDNA libraries were constructed using the TruSeq Stranded Total RNA Preparation Kit (Illumina, cat#: RS-122-2301), which contained Ribo-Zero™ Gold to deplete rRNA. We only amplified our libraries for 10 PCR cycles (instead of 15 suggested by manufacturer) to minimize amplification-induced noise. Purified libraries were quantified using a Kapa library quantification kit (KAPA Biosystems, Wilmington, MA) and then loaded onto a cBot (Illumina, San Diego, CA) at a final concentration of 10 pM to perform cluster generation, followed by 2 x 75 bp sequencing on a HiSeq 2000 (Illumina, San Diego, CA). Two libraries were loaded per lane, yielding an average of 77 M pairs of reads per sample. We mapped the sequencing reads to the reference mouse genome (mm10) using TopHat (version 2.0.10) (Kim et al., 2013) and Bowtie 2 (version 2.1.0) (Langmead and Salzberg, 2012). The number of fragments in each known gene from GENCODE Release M7 (Mudge and Harrow, 2015) was enumerated using htseq-count from HTSeq package (version 0.6.0) (Anders et al., 2015). Genes with fewer than 10 fragments in all samples were removed before differential expression analysis. Differential expression between conditions was statistically assessed by R/Bioconductor package DESeq (version 1.16.0) (Anders and Huber, 2010). Due to the rarity of sorted cells, one sample pooled from multiple sorting was collected for RNA-seq analysis for each cell type. To call differentially expressed genes (DEGs), we used fold change (FC) ≥ 6 and normalized read counts ≥20 for bulk luminal GFP^+^ cells, and FC ≥4 and normalized read counts ≥10 for rare luminal GFP^+^ LRCs. These criteria generated a manageable list of 929 genes preferentially expressed in GFP^+^ cells and 1303 genes in GFP^-^ cells (Table S2).

For Gene Ontology analysis, IPA (Qiagen, Valencia, CA) and DAVID version 6.7 (Huang da et al., 2009) were used with gene symbols. GSEA (the pre-rank function) was carried out by using the curated gene sets (C2) of the Molecular Signature Database (MSigDB) version 4.0 provided by the Broad Institute (http://www.broad.mit.edu/gsea/) (Subramanian et al., 2005). Note that the list of DEGs and entire detectable genes derived from each sample were used for IPA/GO and GSEA analysis, respectively. In particular, we followed the standard procedure as described by GSEA user guide (http://www.broadinstitute.org/gsea/doc/GSEAUserGuideFrame.html). The FDR for GSEA is the estimated probability that a gene set with a given NES (normalized enrichment score) represents a false positive finding, and an FDR <0.25 is statistically significant for GSEA analysis. To reconcile the difference between mouse and human species, the “gene_symbol.chip” function was employed to collapse dataset to gene symbols when performing GSEA. The RNA-Seq data reported in this paper is available in the Gene Expression Omnibus (GEO) database under accession number GSE98760.
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