Proteomic Comparison and MRM-Based Comparative Analysis of Metabolites Reveal Metabolic Shift in Human Prostate Cancer Cell Lines

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ABSTRACT: One of the major challenges in prostate cancer therapy remains the development of effective treatments for castration-resistant prostate cancer (CRPC), as the underlying mechanisms for its progression remain elusive. Previous studies showed that androgen receptor (AR) is crucially involved in regulation of metabolism in prostate cancer (PCa) cells throughout the transition from early stage, androgen-sensitive PCa to androgen-independent CRPC. AR achieves such metabolic rewiring directly either via its transcriptional activity or via interactions with AMP-activated protein kinase (AMPK). However, due to the heterogeneous expression and activity status of AR in PCa cells, it remains a challenge to investigate the links between AR status and metabolic alterations. To this end, we compared the proteomes of three pairs of androgen-sensitive (AS) and androgen-independent (AI) PCa cell lines, namely, PC3-AR+/PC3, 22Rv1/Du145, and LNCaP/C42B, using an iTRAQ labeling approach. Our results revealed that most of the differentially expressed proteins between each pair function in metabolism, indicating a metabolic shift between AS and AI cells, as further validated by multiple reaction monitoring (MRM)-based quantification of nucleotides and relative comparison of fatty acids between these cell lines. Furthermore, increased adenylate kinase isoenzyme 1 (AK1) in AS relative to AI cells may result in activation of AMPK, representing a major regulatory factor involved in the observed metabolic shift in PCa cells.

KEYWORDS: quantitative proteomics, metabolite, metabolic shift, androgen receptor, AMPK, CRPC

INTRODUCTION

Prostate cancer (PCa) is the most prevalent cancer among males.1 PCa patients are initially treated primarily with androgen-deprivation therapy (ADT). However, most cases of advanced prostate cancer progress to castration-resistant prostate cancer (CRPC) after a period of ADT. Due to the limited use of chemotherapy in the management of CRPC, there remains an urgent need for effective therapy that delays or prevents the landmark events that characterize the morbidity associated with this cancer.2

PCa is very heterogeneous in terms of grade, genetics, ploidy, and oncogene/tumor suppressor gene expression, and its biological, hormonal, and molecular characteristics are extremely complex.3,4 Previous studies on clinical samples and xenograft models of PCa indicate that abnormal activation of androgen signaling plays a pivotal role in the progression of CRPC.5 As a result, the sensitivities of different PCa cell lines to androgen are heterogeneous. The differing androgen sensitiv-
ities of PCa cell lines are well-documented (http://capcellines.ca/) and represent the multiple stages of PCa progression.\(^6\) Cai et al.\(^6\) suggested that, in CRPC, partial restoration of the transcriptional activity of the androgen receptor (AR) allowed active expression of the genes that mediate androgen biosynthesis and maintained intracellular androgen level as well as AR activity. Recently, abiraterone acetate, an androgen biosynthesis inhibitor, was shown to improve overall survival in patients with CRPC after chemotherapy and also showed a trend toward improved overall survival in patients who had not received previous chemotherapy.\(^7\)

Advanced CRPC patients treated with abiraterone acetate generally relapse within a year, and AR appears to be active in the relapsed tumors.\(^8\) AR signaling has been shown to cross-talk with alternative survival pathways, such as PI3K/AKT and MAPK pathways.\(^9\) This may partially explain the resistance to AR-targeted therapies in CRPC. However, interactions between AR signaling and these survival pathways are cell-type- and context-dependent.\(^10\) Hence, conclusions based on one PCa cell line could not be directly transferred to all other PCa cell lines or applied to clinical situations.

Here, we set out to obtain a systematic and unbiased understanding of the mechanisms underlying CRPC progression. To this end, we used the iTRAQ labeling approach and performed pairwise comparisons between three androgen-sensitive (AS) and three androgen-independent (AI) PCa cell lines. Also, multiple reaction monitoring (MRM) analyses were applied to targeted metabolites in order to validate the metabolic changes assumed. Our results indicate that a metabolic shift and distinct forms of cell adhesion existed between AS and AI PCa cells.

### Experimental Section

#### Cell Lines and Reagents

Prostate cancer cell lines PC3, Du145, C42B, 22Rv1, and LNCaP were purchased from the American Type Culture Collection (USA). LNCaP cells were maintained in RPMI-1640 supplemented with 10% fetal bovine serum (FBS), 100 U/mL penicillin, and 100 mg/mL streptomycin; the other cells were maintained in DMEM supplemented with 10% FBS, 100 U/mL penicillin, 100 mg/mL streptomycin, and 2 mM glutamine. The PC3-AR\(^+\) cell line, constructed by stable expressing of GFP-tagged AR in PC3 cells, was kindly provided by Prof. Gao Wei-qiang. Cell lines were cultured at 37 °C in a humidified atmosphere containing 5.0% CO\(_2\).

Forty two methyl ester standards of fatty acids and 24 kinds of fatty acids were purchased from Cayman (Michigan, USA) and Sigma (Missouri, USA) as listed in Supporting Information Table S1. Eight standards for nucleotides and acetyl-CoA were purchased from Sigma and are listed in Table S1, and commonly used chemical reagents were purchased from Sigma. All chromatographic reagents were from Avantor Performance Materials (Pennsylvania, USA). Trimethylsilyl diazomethane was purchased from Acros Organics (Massachusetts, USA).

#### Protein Extraction and Denaturing

Cells were washed with cold phosphate-buffered saline (PBS) twice and harvested by scraping in ice-cold lysis buffer (8 M urea, 100 mM Tris-HCl pH 8.5, and protease inhibitor cocktail (Roche, Basel, Switzerland)). Cells were sonicated on ice, and cell lysates were centrifuged at 16 000g for 15 min at 4 °C to remove cell debris. Supernatants were collected and stored at −80 °C until use. Cell lysates were denatured according to the FASP procedure with minor modifications.\(^11\) Briefly, the cell lysates were transferred to Amicon-0.5 ultrafiltration units, and the buffer was exchanged with UA buffer (8 M urea, 0.1 M Tris, pH 8.5) three times. Protein samples were reduced with 10 mM dithiothreitol at 37 °C for 1 h and then alkylated with 50 mM iodoacetamide at room temperature for 1 h (in the dark). The buffer of the denatured protein samples was then exchanged with UA buffer three times and exchanged with 50 mM tetraethylammonium bromide (TEAB) twice. Proteins were recovered by reverse centrifugation at 1000g for 2 min and resolved in 50 mM TEAB. Protein concentrations were determined in triplicate using the bicinchoninic acid method (Thermo Scientific Pierce, Massachusetts, USA) according to the manufacturer's instructions.

#### Liquid Chromatography

The labeled peptide samples were pooled and desalted using Sep-Pak Vac 1 cm\(^3\) (100 mg) C18 SPE cartridges (Waters, Massachusetts, USA). Briefly, the column was washed with buffer B (80% ACN, 50 mM NH\(_4\)HCO\(_3\)) three times and 1 mL each time, then washed with buffer A (50 mM NH\(_4\)HCO\(_3\)) three times and 1 mL each time. Samples were loaded onto the column three times; the column was washed with buffer A three times and 1 mL each time, then eluted with 0.5 mL of 40% ACN, 50 mM NH\(_4\)HCO\(_3\) and 0.5 mL 60% ACN, 50 mM NH\(_4\)HCO\(_3\) sequentially. The combined elute was concentrated in a CentriVap benchtop centrifugal vacuum concentrator (Labconco, Missouri, USA) at 30 °C. The volume was taken to 33 μL with solution A used in peptide fractionation. Thirty microliters of the sample was loaded onto a YMC-Triart C18 basic reversed-phase liquid chromatography column (250 × 4.6 mm, 5 μm particles) (cat. no. TA12S05-2546WT, YMC, Kyoto, Japan). Peptides were separated in a binary buffer system of 2% ACN, pH 10 (pH adjusted with ammonium hydroxide, solution A), and 98% ACN (without pH adjustment, solution B) in an Ultimate-3000 LC system (Thermo Scientific, Massachusetts, USA). The gradient of buffer B was set as follows: 0–5% for 1 min, 5–8% for 5 min, 8–18% for 35 min, 18–32% for 22 min, 32–95% for 2 min, and 95% for 7 min. Eluates were collected every 84 s, and a total number of 50 fractions were obtained. The first five fractions were discarded because most of the excess labeling reagent was eluted during these fractions. The remaining 45 fractions were combined into 15 fractions by merging fraction 1, 16, 31; fraction 2, 17, 32; and so on. Then, all fractions were dried in a vacuum concentrator and stored at −20 °C until further analysis.

#### LC-MS/MS Analysis

For each fraction, an 8 μL sample was injected under trapping conditions using an ekspert nanoLC400 system (AB Sciex, Massachusetts, USA). The gradient of buffer B was set as follows: 0–1% for 1 min, 1–10% for 2 min, 10–90% for 3 min, and 90% for 5 min. Eluates were collected every 84 s, and a total number of 50 fractions were obtained. The first five fractions were discarded because most of the excess labeling reagent was eluted during these fractions. The remaining 45 fractions were combined into 15 fractions by merging fraction 1, 16, 31; fraction 2, 17, 32; and so on. Then, all fractions were dried in a vacuum concentrator and stored at −20 °C until further analysis.
Protein Identification and Quantification

The LC-MS/MS data for the identification and quantification of the global proteome were analyzed using ProteinPilot Software 4.5 (AB Sciex, Massachusetts, USA) against the Uniprot KB human database (released on June, 2013) with 88 771 entries, to which 175 commonly observed contaminants and all the reverse sequences were added. iTRAQ 8-plex cations. A 95% confidence (unused protein score >1.3) was used for protein identification. The PDST_V3.708pE_-TEMPLATE (AB Sciex, Massachusetts, USA) was used to calculate p values for each ratio. Ratio channels of 114/113, 116/115, 117/118, and 119/121 were assigned as "target". Ratio channels of 117/119 and 118/121 (technical replicates) were assigned as "decoy" (Figure 1). By assigning each ratio channel as either "target" or "decoy", the PDST estimates target-decoy global false discovery rate (FDR) for the ratios according to their p values. The p value limit was set on the basis of target-decoy FDR analysis. Principally, a protein with a quantitative ratio differing from 1 and the ratio’s p value less than 0.01 is declared as differentially expressed between a pair of AS and AI cells.

Method Validation for MRM Analysis

Method performance was evaluated for 42 fatty acid methyl ester (FAME) standards, eight nucleotides, and acetyl-CoA (Supporting Information, Table S1), with respect to linearity and reproducibility. For each FAME standard, ≥0.1 mg/mL stock solution was prepared in methanol and stored at −20 °C. From these stock solutions, mixtures of 42 FAMES dissolved in hexane were prepared at 0.05, 0.1, 0.5, 1, 5, 10, 50, and 100 ng/mL and analyzed using GC-MS. For each nucleotide and acetyl-CoA standard, 1 mM stock solution was prepared in water and stored at −20 °C. From these stock solutions, mixtures of the nine metabolites dissolved in water were prepared at 0.17, 0.42, 0.83, 1.67, 8.33, and 83.3 μM and analyzed using LC-MS. Linearity was evaluated using the linear regression of peak areas with respect to concentration. In addition, FAME standards at 0.5 ng/mL were run two times to determine inter-run quantitative reproducibility, as measured by the relative standard deviation (RSD).

MRM-Based Relative Quantification of Fatty Acids

Fatty acids were extracted from PC3, Du145, 22Rv1, LNCaP, and C42B using a liquid–liquid extraction method. Briefly, cells were grown to confluence in 6 cm dishes under conditions same as those in the proteomic experiment. After the medium was removed, cells were washed with 5 mL of cold PBS before 300 μL of 0.9% KCl solution was added, and cells were then detached with a cell scraper. Deuterium-labeled C17:0 fatty acid was added as an internal standard. Cells were transferred to an Eppendorf Safe-Lock 2.0 mL tube (cat. no. 0030 120.094, Eppendorf, Hamburg, Germany). After adding 300 μL of methanol and the antioxidant reagent butylated hydroxytoluene (0.01% final concentration), cells were incubated overnight at −20 °C. Then 1 mL of methyl tert-butyl ether (MTBE) was added, and the samples were vortexed thoroughly on a Vortex-Genie 2 mixer (cat. no. SI-0236, Scientific Industries, New York, USA) for 1 h at room temperature for phase separation. After centrifugation at 18 000g for 5 min at 4 °C, the upper phase was collected and divided into two parts. One part (200 μL) was used for analysis of phospholipid-derived fatty acids (FFs). The other part was used for analysis of free fatty acids (FFAs). Both parts were dried under vacuum. FFs were derivatized according to the analytical procedure on the Web site of Avanti Polar Lipids company (http://www.avantilipids.com/index.php?option=com_content&view=article&id=1693&Itemid=412). Samples were dissolved in 1 mL of hexane. Fifty microliters of sodium methoxide (1 M) was added and vortexed thoroughly. Samples were incubated at room temperature for 5 min and centrifuged at 1600 rpm at 20 °C for 5 min. The upper phase was transferred to another Safe-Lock tube and vacuum-dried and then centrifuged in 50 μL of hexane for analysis. For FF derivatization, trimethylsilyl (TMS)–diazomethane was used. Samples were resolved in methyl esterification buffer (500 μL of MTBE/150 μL of...
MeOH/6.5 μL of 1 M HCl) in a Safe-Lock tube. Fifty microliters of TMS–diazomethane (2 M in hexane) was added and mixed at a mild speed on a wheel for 30 min at room temperature. Six microliters of glacial acetic acid was added to quench the reaction. After 1 min, 125 μL of water was added and mixed thoroughly. Samples were centrifuged at 1500g for 3 min for phase separation. The upper phase was collected and dried under vacuum. Samples were then dissolved in 50 μL of hexane for analysis. Derivatized FFs and PFs were separated using Agilent 6890N gas chromatograph (Agilent, California, USA) equipped with a J&W HP-88 column (100 m × 0.25 mm i.d. × 0.2 μm film thickness) (cat. no. 112-88A7E, Agilent, California, USA). The flow of helium carrier gas was set to 1 mL/min (20 cm/s); the injector temperature was set to 260 °C, and the temperature of the sample transfer line was set to 240 °C. Samples were analyzed in splitless mode, and the gradient started from 50 °C, held for 1 min, then increased by 25 °C/min to 175 °C, further by 1 °C/min to 216 and 4 °C/min to 240 °C, then held for 10 min. Mass spectra were collected on an Agilent 7000B mass selective detector (Agilent, California, USA), using a chemical ionization source in the positive mode. Each cell line was analyzed in biological triplicate. A washing run was added with every three samples. In order to eliminate the influence of fatty acid contaminations, three solvent blanks were treated in parallel with samples, and the mean values of the peak areas of solvent blanks were subtracted from each sample. In addition, a mixture of 24 fatty acid standards (Table S3) dissolved in methanol was prepared at 5 ng/mL, derivatized with TMS–diazomethane, and used for recovery rate evaluation.

**MRM Quantitation of Nucleotides and Acetyl-CoA**

Nucleotides and acetyl-CoA were extracted from five cell lines (PC3, Du145, 22Rv1, LNCaP, and C42B), according to a method published earlier. Briefly, cells were grown to confluence (~10^7 cells) in 10 cm dishes under conditions that were the same as those in the proteomic experiment, and cells were washed twice with ice-cold PBS. Then 100 μL of ice-cold water was added into the 10 cm dish, which was then flash frozen at ~80 °C for about 30 s. Cells were detached with a cell scraper on ice and transferred into 1.5 mL Eppendorf tubes. Cell suspensions were lysed by two rapid freeze–thaw cycles at ~80 °C, followed by 30 s of sonication on ice. The volume was made up to 400 μL with ice-cold water, and each sample was then divided equally into two Eppendorf tubes. Five hundred microliters of cold methanol and 250 μL of chloroform were added to the homogenates and vortexed. For safety considerations, addition of chloroform as well as other toxic reagents and sample transfer were always performed in a fume hood. Samples were mixed for 10 min at 4 °C and centrifuged for 5 min at 2000g at 4 °C to remove debris and precipitate protein. The pellet was dissolved in 8 M urea, 0.1 M Tris, pH 8.5, and protein content was quantified using a BCA kit. The supernatant was transferred to a new tube, and 250 μL of chloroform and 250 μL of water were added. Samples were vortexed for 60 s, kept on ice for 10 min for phase separation, and centrifuged for 10 min at 2000g at 4 °C. The resulting upper polar and lower nonpolar fractions were separately transferred to Eppendorf tubes, vacuum-dried, and combined after being dissolved in 30 μL of LC buffer A (5 mM ammonium acetate/0.1% FA). A combined volume of a ~60 μL sample was obtained and centrifuged, and a 5 μL centrifuged sample was analyzed on an Agilent 6460 LC-Triple Quadruple MS (Agilent, California, USA) with an Acclaim 120 C18 RP column (3 × 150 mm, 3 μm particles) (cat. no. 063691, DIONEX, California, USA). Metabolites were eluted from the column with buffer B (ACN/5 mM ammonium acetate/0.1% FA), and buffer A was 5 mM ammonium acetate/0.1% FA. The gradient of buffer B started at 2% and was kept for 1 min and then increased from 2 to 30% in 9 min and from 30 to 80% in 5 min. Then, the gradient of buffer B was decreased to 2% in 2 min and kept for 13 min. The total LC time was 30 min, with a flow rate of 0.5 mL/min. Gas temperature was set at 325 °C, and gas flow was 7 L/min. Sheath gas temperature was set at 325 °C, and sheath gas flow was 11 L/min. Capillary voltage was 3500 V, and the nozzle voltage was 500 V. Delta EMV(+) was 200. Each cell line was analyzed in biological triplicate.

**Mitochondrial Staining and Activity Analysis**

Mitochondrial morphology imaging was conducted with Mitotracker Green (Life Technologies, Massachusetts, USA) as described by Chen et al. Cells were seeded in 35 mm Petri dishes, and after reaching 50–70% confluence, the cells were stained with 50 nM Mitotracker Green diluted in the corresponding culture media for 20 min at 37 °C. After incubation, cells were washed twice with PBS, stained with Hoechst 33342 (Life Technologies, Massachusetts, USA) for 5 min, and washed three times with PBS. Cell images were obtained with an Olympus FV1000 laser scanning confocal microscope (OLYMPUS, Tokyo, Japan).

**Data Analysis**

Enrichment analysis for gene ontology (GO) terms and protein interaction network construction was performed using STRING. The p values were corrected for multiple testing using the method by Benjamini and Hochberg, and the top 10 significant categories were presented according to their p values. Pathway mapping was executed using the Reactome database. The Matlab software (version 2010b, MathWorks) was used for hierarchical clustering analysis. Proteins were clustered using unsupervised hierarchical average linkage clustering and a Euclidian distance metric, based on their logarithm-transformed ratios.

For fatty acids, peak areas were integrated with MassHunter software (Agilent, California, USA). After blank subtraction, internal standard correction (this correction was only applied for FFs), and protein content correction, the values were log-transformed and z-score-normalized and then clustered using unsupervised hierarchical average linkage clustering and a Euclidian distance metric. Fatty acids with a signal-to-noise ratio less than 10 in any sample were excluded for further analysis. For nucleotides and acetyl-CoA, the concentrations were calculated directly via the standard curves built with MassHunter software. After correction for protein content, the contents of nucleotides and acetyl-CoA among different...
samples were compared using the student’s t-test with a p value of <0.05 as the significance level.

**Safety Considerations**

All procedures involving toxic reagents should be performed in a fume hood, and the waste produced must be recovered under specific recovery processing. The TMS-diazomethane was originally developed as a less toxic substitute for the compound diazomethane. However, inhalation of TMS-diazomethane may cause lung damage or central nervous system depression and should therefore be handled with the appropriate safety procedures in place, that is, inside a fume hood, with adequate personal safety equipment. Excess TMS-diazomethane is to be neutralized using acetic acid. Care must be taken if handled in large volumes, as it is highly volatile and produces nitrogen gas.

**RESULTS AND DISCUSSION**

**Differentially Expressed Proteins between AS and AI Cells**

In order to identify the differentially expressed proteins between the three pairs of PCa cells, we performed 8-plex iTRAQ labeling and used a target-decoy global FDR method to set a cutoff p value for the ratios of quantified proteins. A total number of 4953 proteins were identified with 1% global FDR at the protein level, among which 4847 proteins were quantified (Table S2). When setting the cutoff p value below 0.01, the target-decoy global FDR for quantification was 2%. In total, 528 proteins showed at least one quantitative ratio with a p value less than 0.01 (Figure S1). By applying hierarchical clustering to the quantitative ratios of the 528 proteins, we found that the changes of these differentially expressed proteins were heterogeneous among the three pairs of AS and AI cell lines (Figure 2A). Among the 528 proteins, 191 were differentially expressed in at least two pairs, with 72 proteins increased, 76 decreased, and 43 proteins changed in both directions in AI relative to AS cells (Figure 2B). The other 337 proteins were found to be differentially expressed in only one pair (Table S4).

Only 19 proteins among the differentially expressed proteins were changed in the same direction in all of the three pairs, among which 10 were increased and 9 were decreased in AI relative to AS cells (Figure 2B and Table 1). Two of the 19 proteins have been shown previously to be critically involved in cancer development. They are major vault protein (MVP) and poly(ADP-ribose)polymerase 1 (PARP-1). MVP was increased, and PARP-1 was decreased in AI relative to AS cells. MVP has been reported to promote cancer progression by mediating epidermal growth factor receptor (EGFR)/PI3K-mediated migration and survival pathways in human glioblastoma.15

Intriguing, abnormal activation of the PI3K pathway is a

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**Figure 2. Dynamic changes of differentially expressed proteins in different pairs of PCa cells and gene ontology annotations of these proteins. (A) Hierarchical clustering based on the 10-base logarithm-transferred ratios of the 528 differentially expressed proteins. By setting the distance cutoff, four clusters showing in different colors were identified. (B) Venn diagram depicts the distribution of these differentially expressed proteins in the three pairs, and the histogram at the bottom represents the regulation directions of the 191 proteins that change in at least two pairs of PCa cells, for which a is PC3-AR+/PC3, b is 22Rv1/Du145, and c is LNCaP/C42B. (C) Enrichment analysis of gene ontology annotations for the 191 differentially expressed proteins that changed in at least two pairs of PCa cells, GOBP for biological process, and GOCC for cellular location.**
common event during PCa progression. Here, we also found higher expression of EGFR in AI relative to AS cells (Table S4). Thus, we speculate that MVP plays a similar role in PCa due to its increased expression in AI relative to AS cells (Table S4). Previously, it was shown that during neoplastic transformation both quantitative increase and isoform alterations of PFK occurred secondary to altered gene expression. In particular, PFK controls the entry of glucose metabolites into the downstream steps of glycolysis.

Differentially Expressed Proteins Mainly Function in Central Metabolism

To investigate the functions of these differentially expressed proteins as well as their roles in CRPC progression, the 191 proteins whose expressions changed in at least two pairs of AS and AI PCa cells were chosen for GO annotation enrichment analysis. The result showed that most of these proteins were involved in cell metabolism and located in exosomes (Figure 2C). Exosomes play a pivotal role in cancers, we thus searched the exosome database Exocarta and Vesiclepedia and classified these exosome-located proteins based on their reported origins (Figure S2). These proteins likely play important roles in PCa progression.

Our results showed that nearly all of the enzymes involved in the TCA cycle showed a decrease in expression, while glycolytic enzymes including ATP-dependent 6-phosphofructokinase, platelet-type (PFKP), glyceroldehyde-3-phosphate dehydrogenase (GAPDH), α-enolase (ENO1), fructose−phosphatase aldolase A (ALDOA), and the rate-limiting enzyme LDHA showed increased expressions in AI relative to AS cells (Figure S3 and Table S4). Specifically, we observed that different isoforms of PFK changed in both directions. PFKP was increased while PFKM was decreased in AI relative to AS cells (Table S4). Previously, it was shown that during neoplastic transformation both quantitative increase and isoform alterations of PFK occurred secondary to altered gene expression. In particular, PFK controls the entry of glucose metabolites into the downstream steps of glycolysis. Recently, Ros et al. found that the glycolytic enzyme PFKFB4 is essential for PCa cell survival, confirming the importance of glycolysis regulation in PCa.

### Table 1. Proteins Showing Consistently Expressional Changes in PCa Cells

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</tbody>
</table>

Since glucose is mainly utilized during glycolysis, glutamine-driven TCA turning powers oxidative phosphorylation and accounts for the majority of ATP production in most cancer cells. A recent report showed that PC3 cells use glutamine as their main energy source and revealed a key role of glutaminase (GLS1) in the growth of PCa cells. In agreement with these studies, we also observed increased levels of GLS1 in C42B relative to LNCaP and in Du145 relative to 22Rv1, and its increase in PC3 relative to PC3-AR− was not significant (Table S4). However, glutamate dehydrogenase 1 (GLUD1) was decreased in C42B relative to LNCaP and in Du145 relative to 22Rv1 (Table S4), suggesting a more complex regulation of glutamine metabolism in these cells. Recently, GLUD1-dependent reductive glutamine metabolism was shown to increase in PCa cells when glucose oxidation was decreased by metformin. The same study observed a more obvious effect of metformin in the LNCaP cells, indicating that AS cells were dependent more on reductive glutamine metabolism than were AI cells. In support of this observation, another key enzyme for reductive glutamine metabolism, isocitrate dehydrogenase [NADP] cytoplasmic (IDH1), was also shown to be increased.
in AS relative to AI cells in our results (Table S4). In addition, we found the increase of the mitochondrial aspirate—glutamate carrier SLC25A13 in AS relative to AI cells (Table S4). These results, together with the information mentioned above, likely reflect that glutamine enters multiple metabolic pathways for distinct utilities in these PCa cells (Figure 5B).

Functional Changes of Mitochondria Occur between AS and AI Cells

To investigate whether the decrease of enzymes in the TCA cycle in AI cells was caused by loss of mitochondrial mass, staining of mitochondria and mitochondrial activity analysis were conducted in each of the five cell lines (Figure 3). We excluded PC3-AR+ cells because they showed distinct changes of the described TCA enzymes relative to PC3 when compared to the other two pairs. We did not observe a significant loss of mitochondrial mass in the AI PCa cells from the staining result, although the morphology and distribution of mitochondria varied considerably (Figure 3A). Moreover, expressions of the mitochondrial electron transport chain (ETC) components, including electron transfer flavoprotein (ETF) and ATP synthase, were increased in AS relative to AI cells except for those in the PC3-AR+/PC3 pair in our results (no change of ETF and decrease of ATP synthase, Figure S4). However, no changes were observed in the other ETC components as well as many quantified structural proteins of mitochondria in our results, though it may be partially explained by a lack of sufficient quantitative peptides. Therefore, functional alterations rather than mass loss of mitochondria could eventually lead to such a complex change of proteins located in this organelle among different pairs of PCa cells. To further prove this, two mitochondrial proteins showed opposite changes in our results. They are the Lon protease homologue mitochondrial LONP1 that increased and calcium-binding mitochondrial carrier protein SCaMC-1 (SLC25A24) that decreased in AS relative to AI cells (Table S4).

Figure 3. Analysis of mitochondrial morphology and function in PCa cells. (A) Mitochondria were stained with MitoTracker Green, and nuclei were stained with Hoche 33342. (B) Mitochondria metabolic activity analysis. PC3, Du145, and C42B cells were plated at 15 000 cells/well, and 22Rv1 and LNCaP cells were plated at 40 000 cells/well in XF24 V7 cell culture plates 24–48 h prior to the assays. For OCR assay, oligomycin (1 μM), FCCP (0.5 μM), and rotenone (1 μM) with antimycin (1 μM) were added at time points indicated by arrows. For the ECAR assay, glucose (10 mM), oligomycin (1 μM), and 2-DG (100 mM) were added at time points indicated by arrows. Each data point represents mean ± SD (n = 3–4). Medium used for different assays was described in the experimental method. For OCR response, data were presented as percentages of baseline values (measurement of 3).
relative to AI cells (Table S4). LONP1 plays a key role in metabolic reprogramming by remodeling OXPHOS complexes. SLC25A24 regulates the contents of adenosine nucleotides in the mitochondrial matrix and increases the antiapoptotic activity of mitochondria. Both of them are involved in timely response of mitochondria to cellular metabolic dynamics. An even more complex change was observed for dynamin-like 120 kDa protein, mitochondrial OPA1, which was increased in PC3-AR+ relative to PC3 and in LNCaP relative to C42B, but was decreased in 22Rv1 relative to Du145 (Table S4). OPA1 is the key effector for fusion of the mitochondrial inner membrane. Importantly, the rate of mitochondrial fusion responds to changes in metabolism. Together, these results reveal that metabolism-related functional alteration rather than mass loss of mitochondria occurs between AS and AI cells.

**High Rate of Glycolysis Serves the High Demand for a Biosynthetic Precursor in AI Cells**

Cancer cells are known to rewire their metabolism and energy production networks to support rapid proliferation. In contrast to resting cells, proliferating cancer cells are in much greater need of building blocks for reductive biosynthetic reactions.

To further investigate the metabolic signatures of PCA cells, we compared the contents of several metabolites in PCA cells by the MRM approach. In Table S1, we summarize the results of MRM method validation for all targeted metabolites in this study. All metabolites gave a linear response ($R^2 > 0.97$). As shown in Figure 4 and Table S5, the contents of ADP, ATP, AMP, and cAMP in PCA cells are shown as mean $+ SD$ ($n = 3$) in (a), and their ratios to ATP are shown as mean $+ SD$ ($n = 3$) in (b). The cellular content of acetyl-CoA is shown as mean $+ SD$ ($n = 3$) in (c). The cellular content of NAD and NADH is shown as mean $+ SD$ ($n = 3$) in (d), and the cellular content of NADP and NADPH is shown as mean $+ SD$ ($n = 3$) in (e). The ratios of NADH/NAD, as well as NADPH/NADP, are shown as mean $+ SD$ ($n = 3$) in (f). The letters above the columns denote significant differences ($p < 0.05$) for comparison, among which “a” represents comparison between C42B and LNCaP and “b” represents comparison between Du145 and 22Rv1.
AMPK activation. AMPK is a core regulator of cellular metabolism. A previous study using human PCa specimens showed that AMPK is highly activated in 40% of the cases, and it serves as a downstream effector of AR-induced PCa cell proliferation.36 In androgen-treated PCa cells, the androgen-mediated AMPK-PGC-1α signaling cascade is responsible not only for increasing the rate of glycolysis but also for increasing glucose and fatty acid oxidation.37 Furthermore, Ca2+/calmodulin-dependent protein kinase kinase β (CaMKKβ), an AMPK kinase, was found to be highly expressed in normal prostate tissue and elevated in prostate cancer.38 One recent study reported that a novel AMPK activator inhibited the growth of PCa cells.39 It is worth mentioning that the study includes different AR phenotypes, and the basal level of AMPK activation and the functions of AMPK activator are different in these PCa cells. However, the role of AMPK in prostate cancer is controversial.40,41 The timing, level, and cellular context of AMPK activation may have profound effects on the functional consequences of AMPK signaling.42 A more complicated negative feedback loop exists between AR and AMPK in PCa cells.43

In our proteomic results, an increased expression of AK1 in AS relative to AI cells was observed (Table S4). The highly reversible adenylosuccinate kinase reaction can be partially displaced toward synthesis of AMP and cause a large rise in AMP,44 representing an alternative route for AMPK activation in AS cells (Figure 6). Further studies are required to confirm these results.

Simultaneous Increase of Fatty Acid Synthesis and Fatty Acid Oxidation in AS Cells

A metabolic shift, in which PCa cells use alternative enzymes and pathways to facilitate the production of fatty acids, is proposed to contribute to the well-established lipogenic phenotype of PCa.45 In agreement with this, we observed an increase of fatty acid synthase (FASN) in AS relative to AI cells (Table S4). Importantly, FASN has been reported to be a candidate oncogene in PCa.46 However, changes of other enzymes in fatty acid synthesis, such as ATP-citrate synthase (ACLY) and acetyl-CoA carboxylase 1 (ACC), were complex among the different pairs (Table S4). ACLY was decreased in LNCaP relative to C42B but increased in 22Rv1 relative to Du145. ACC was only increased in 22Rv1 relative to Du145 and showed no difference in the other two pairs. In agreement with the changes observed for ACLY and ACC, the content of acetyl-CoA was higher in C42B cells compared to LNCaP as well as to those in the other PCa cells, but showed no difference between Du145 and 22Rv1 (Figure 4c), suggesting that acetyl-CoA accumulated in C42B but was consumed in 22Rv1 through fatty acid synthesis. Another function of acetyl-CoA is to serve as substrate for cholesterol synthesis. We observed that squalene synthase (FDFT1), the key enzyme for cholesterol synthesis, was present at an elevated level in AS relative to AI cells (Table S4), a fact that is in agreement with the increased de novo synthesis of cholesterol in PCa cells.47 Recently, AI cells were shown to bear aberrant accumulation of esterified cholesterol in lipid droplets, which arises from an
uptake of exogenous lipoproteins and is independent of androgen signaling. Thus, AI cells appear to obtain cholesterol through exogenous uptake rather than de novo synthesis, which is in contrast to AS cells.

After synthesis, fatty acids can be oxidized for energy supply or converted into the carbon skeleton of triglyceride for lipid storage as well as for phospholipid synthesis. We observed that carnitine O-palmitoyltransferase (CPT) changed in the isoform context among different pairs of PCa cells (Table S4). CPT1A and peroxisomal carnitine O-octanoyltransferase (CROT), which are located on the outer membrane of mitochondria and peroxisomes, respectively, were found to be increased in LNCaP relative to C42B, while showing no difference in the other pairs. CPT2, which locates on the inner membrane of mitochondria, was increased in LNCaP relative to C42B and in 22Rv1 relative to Du145. One reason for this observation could be that oxidation for different fatty acids in mammalian cells is compartmentalized according to the lengths of their acyl chains. In comparison with AI cells, AS cells overexpressed several mitochondrial enzymes that are responsible for fatty acid β-oxidation, including long-chain fatty acid–CoA ligase 3 (ACSL3), enoyl-CoA hydratase (ECHS1), Delta(3,5)-Delta(2,4)-dieneoyl-CoA isomerase, mitochondrial (ECH1), trifunctional enzyme subunit alpha (HADHA), and acetyl-CoA acetyltransferase (ACAT1) (Table S4). Consistent with the increased levels of these enzymes, most of the 17 FFs reliably detected by MRM were found to be increased in AS relative to AI cells (Figure 5A, blue cluster). Together, these results indicate that fatty acid β-oxidation in mitochondria is increased in AS cells. This may result from the androgen-mediated AMPK-PGC-1α signaling cascade, which directly causes the increase of fatty acid β-oxidation. In order to investigate the fates of fatty acids in PCa cells, we also compared the changes of PFs in different PCa cells relatively using MRM. Consistent with the complex changes of lipogenic enzymes observed here (data not shown), the contents of different PFs showed complex changes between AS and AI cells (Figure 5A). Our results were consistent with a recent MS study on phospholipid composition in PCa cells. That study reported that AS cells contain more polyunsaturated phospholipids but less saturated and monounsaturated phospholipids than AI cells. Moreover, we observed that the contents of short-chain PFs as well as FFs were higher in AS relative to AI cells, indicating that synthesized FFs were partially converted to short-chain PFs in AS cells. Long-chain PFs, especially the monounsaturated counterparts, displayed changes in opposing directions in our results, suggesting preferential uptake from the environment rather than de novo synthesis in AI PCa cells.

Figure 6. Proposed model of the AR signaling network and its role in cellular metabolism regulation. Different status of AR activity regulates the transcription of a distinct set of androgen-responsive genes (ARGs) and results in the two types of PCa cells, that is, AS and AI (also called CRPC cells). A major role of AR is its inhibitory effect on in situ androgen synthesis, which decreases intracellular androgen level directly and forms a negative feedback loop in AR activity regulation. This negative feedback loop suppresses AR signaling at high androgen levels but allows increased AR and androgen synthesis in CRPC. ADT induces the change of intracellular androgen level, switching AR to partial activation, and finally contributes to androgen independence. AR activation will increase the expression of CaMKKβ and activate AMPK, and the latter will conversely inhibit AR activation. As a central regulator of cellular metabolism, AMPK mediates metabolic shift in PCa cells. Different ARGs regulated by distinct AR activity status also contribute to this shift. AK1 can convert ADP to AMP and increase the AMP/ATP ratio, leading to AMPK activation. Different metabolic pathways highlighted in the colored octagonal boxes are dominant in distinct PCa cells, and the related enzymes that are differentially expressed in PCa cells are highlighted in the colored boxes above each pathway.
Thus, we suggest that a simultaneous increase in synthesis and β-oxidation of fatty acids occurs in AS cells, which may serve as a major energy supply (Figure SB). In agreement with our hypothesis, a previous study showed that, in PCa cells, androgen could induce the activation of both lipogenesis and fatty acid degradation in a coordinated fashion.51 Further studies will be required to clarify the regulation of lipid metabolism in CRPC cells.

**AI PCa Cells Favor Cell–ECM Focal Adhesion over Cell–Cell Adhesion**

Another category of proteins that was observed to show changes in expression between AS and AI is that of factors associated with cell adhesion. These factors included tubulin β-6, talin-1, Fermitin family homologue 2, IQGAP1, fascin, actin-related protein 3, CAP1, TWF2, coronin, and profilin-1. All of these proteins were found to be present at elevated levels in AI relative to AS cells. In addition, α-actinin-1 and isoform 1 of vinculin changed in the same direction between AS and AI cells (Table S4). AI cells also expressed more integrin β1 and less E-cadherin relative to AS PCa cells (Table S4). It is well-established that integrin β1 mainly mediates cell adhesion to extracellular matrix (ECM) components, whereas E-cadherin mainly mediates epithelial cell–cell adhesion.52 According to the model of adhesion dynamics proposed by Parsons et al.,53 we conclude that AI PCa cells favor focal adhesion to ECM over cell–cell adhesion based on the changes of these adhesion-related molecules observed here. Adhesion-mediated signaling through integrin β1 and integrin-linked kinase has a critical role in caveolae formation.54 In addition to caveolae formation, caveolin-1 plays many other roles in CRPC progression.10 Consistently, we observed that caveolin-1 was increased in AI relative to AS cells, though the p values were above 0.01 (data not shown), which may be a result of insufficient numbers of peptides for quantification. Another key cell–ECM adhesion molecule, CD44, was also increased in AI relative to AS cells (Table S4). Previously, CD44 was shown to connect to the actin cytoskeleton, bind to growth factors, and enhance signaling through cis interactions with EGFR.52 Whether CD44 links to EGFR for PCa cell adhesion needs further investigation.

Overall, AI cells favor integrin-β1-mediated cell adhesion to ECM over E-cadherin-mediated cell–cell adhesion. Like the metabolic enzymes, most of these adhesion-related molecules also locate in the exosome (Figure S2). Therefore, elucidating the internal relations between cell adhesion and metabolism may finally uncover the role of exosomes in PCa.

**CONCLUSION**

An increasing number of studies reveal the heterogeneity of PCa as a problem for the identification of common features in its biological properties. In this study, we took this heterogeneity of PCa into consideration and compared the proteomes of three pairs of PCa cells. We identified 19 differentially expressed proteins that changed in the same direction in all three pairs, among which four proteins have been reported to be the products of androgen-responsive genes. Our results suggest that the 19 proteins are pivotal in CRPC progression. Furthermore, isozymic alteration of PKF, together with the resulting aberrant glycolysis, may accompany CRPC progression, based on the established role of glycolysis in rapid proliferating cancer cells. Apart from altered glycolysis, glutamine and fatty acid metabolisms are also different between AS and AI cells according to our proteomic results as well as MRM analysis of fatty acids, nucleotides, and acetyl-CoA. These changes suggest a metabolic shift between AS and AI cells and reveal a central place of mitochondria in this shift. Increased glycogen mobilization is likely to represent an additional sign for the metabolic shift, based on our observations that two isoforms of glycogen phosphorylase, namely, PYGL and PYGB, were elevated in AI relative to AS cells (Table 1 and Table S4). The simultaneous increases of AMP and AK1 likely contribute to AMPK activation in AS cells, suggesting a potential mechanism underlying the metabolic shift (Figure 6). Furthermore, integrin-β1-mediated cell adhesion to ECM was preferentially utilized in AI relative to AS cells. Because many differentially expressed metabolic enzymes as well as adhesion molecules observed in our study are located in exosomes, the synergistic action between cell adhesion and metabolism in exosomes confirmatively contributes to CRPC progression and deserves further investigations.


(20) Hosseini-Beheshti, E.; Pham, S.; Adomat, H.; Li, N.; Tomlinson Guns, E. S. Exosomes as biomarker enriched microvesicles: character-


