Stable Isotope Labeling with Amino Acids (SILAC)-Based Proteomics of Primary Human Kidney Cells Reveals a Novel Link between Male Sex Hormones and Impaired Energy Metabolism in Diabetic Kidney Disease*

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Male sex predisposes to many kidney diseases. Considering that androgens exert deleterious effects in a variety of cell types within the kidney, we hypothesized that dihydrotestosterone (DHT) would alter the biology of the renal tubular cell by inducing changes in the proteome. We employed stable isotope labeling with amino acids (SILAC) in an indirect spike-in fashion to accurately quantify the proteome in DHT- and 17β-estradiol (EST)-treated human proximal tubular epithelial cells (PTEC). Of the 5043 quantified proteins, 76 were differentially regulated. Biological processes related to energy metabolism were significantly enriched among DHT-regulated proteins. SILAC ratios of 3 candidates representing glycolysis, N-acetylglucosamine metabolism and fatty acid β-oxidation, namely glucose-6-phosphate isomerase (GPI), glucosamine-6-phosphate-N-acetyltransferase 1 (GNPNAT1), and mitochondrial trifunctional protein subunit alpha (HADHA), were verified in vitro. In vivo, renal GPI and HADHA protein expression was significantly increased in males. Furthermore, male sex was associated with significantly higher GPI, GNPNAT1, and HADHA kidney protein expression in two different murine models of diabetes. Enrichment analysis revealed a link between our DHT-regulated proteins and oxidative stress within the diabetic kidney. This finding was validated in vivo, as we observed increased oxidative stress levels in control and diabetic male kidneys, compared with females. This in depth quantitative proteomics study of human primary PTEC response to sex hormone administration suggests that male sex hormone stimulation results in perturbed energy metabolism in kidney cells, and that this perturbation results in increased oxidative stress in the renal cortex. The proteome-level changes associated with androgens may play a crucial role in the development of structural and functional changes in the diseased kidney. With our findings, we propose a possible link between diabetic and non-diabetic kidney disease progression and male sex hormone levels. Data are available via ProteomeXchange ([https://www.ebi.ac.uk/pride/archive/](https://www.ebi.ac.uk/pride/archive/)) with identifier PXD003811.

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Chronic Kidney Disease (CKD) often results in irreversible deterioration of renal function that can progress to renal failure (1). Sex plays a relevant role in the progression and severity of many kidney diseases (2). At a clinical level, it is

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Note: The abbreviations used are: CKD, chronic kidney disease; AR, androgen receptor; ARE, androgen response element; DHT, dihydrotestosterone; DKD, diabetic kidney disease; ERα/ERβ, estrogen receptor alpha/estrogen receptor beta; ERE, estrogen response element; EST, 17β-estradiol; FAO, fatty acid beta-oxidation; GNPNAT1, glucosamine-6-phosphate-N-acetyltransferase 1; GPER, G protein coupled estrogen receptor 1; GPI, glucose-6-phosphate isomerase; HADHA, mitochondrial trifunctional protein subunit alpha; HBP, hexosamine biosynthetic pathway; HK-2, immortalized human kidney cells; LC-MS/MS, liquid chromatography – tandem mass spectrometry; PTEC, proximal tubular epithelial cells; ROS, reactive oxygen species; SILAC, stable isotope labeling with amino acids in cell culture; TCA, tricarboxylic acid.
generally accepted that male sex is a risk factor for CKD (3). In addition, sex differences impact the progression of renal disease in animal models of non-diabetic (4) and diabetic nephropathy (5–7).

Experimental evidence suggests that androgens play a detrimental role in kidney disease. Testosterone promoted apoptosis in immortalized and primary human proximal tubule cells (8). In addition, preincubation with the antiandrogen flutamide prevented the apoptotic effects of testosterone, indicating that deleterious effects of androgens are mediated by testosterone conversion to DHT and binding to androgen receptor (AR). In turn, EST significantly attenuated the fibrotic effect of TGFβ-1 in mesangial cells (9). The counter-regulatory effects of androgens and estrogens have also been observed in podocytes, where EST prevented the testosterone-induced increase in the percentage of TUNEL-positive cells. In this sense, both estrogen receptor alpha (ERα) deficiency and testosterone administration were associated with podocyte loss and augmented apoptosis in vivo (10). In type 1 diabetic castrated male rats, low dose of DHT attenuated, whereas high dose accentuated, the severity of several hallmarks of kidney disease such as glomerulosclerosis and tubulointerstitial fibrosis (11).

Androgens and estrogens have shown to trigger genomic and non-genomic events within the renal cortex (12) and in renal tubular cells in culture (13, 14). Genomic actions of sex hormones are mediated by primary interactions with their specific nuclear receptors (AR for androgens and ERα/ERβ for estrogens) (15). In addition, estrogens promote rapid extranuclear signaling upon binding to ERα, ERβ or G protein coupled estrogen receptor 1 (GPER) localized on the cell membrane (16), promoting several downstream events such as changes in cAMP (17, 18), recruitment and activation of the MAPK/ERK signaling cascade (19), and induction of PI3K/AKT signaling to activate eNOS (20). Androgens can also stimulate MAPK/ERK (21) and AKT pathways (22) after interacting with the AR or the G-coupled receptor GPRC6A on the cell membrane (23, 24).

Taken together, in vitro and in vivo data suggest that androgen signaling pathways in the renal cortex play a harmful effect on the progression of CKD. However, the specific mechanisms by which androgens detrimentally regulate kidney cells over estrogens remain unclear. To enhance our understanding of sex effects on CKD progression, we aimed to perform an in-depth analysis of the sex hormone-regulated proteome in human proximal tubular epithelial cells (PTEC) after stimulation with DHT or EST. We hypothesized that male sex hormones induce changes in the proteome of the renal tubular cell in a more detrimental manner than estrogens.

Quantitative proteomics is widely considered as an invaluable approach to explore biological systems in vitro and in vivo (25–27). To achieve our aim, we carried out a quantitative strategy by using stable isotope labeling of amino acids in cell culture (SILAC) in combination with a high-resolution mass analyzer. To quantify the proteome of a single cell line, the SILAC experiment can be performed in a classical or in a spike-in format (28). One of the advantages of spike-in SILAC is that the experimental cells are maintained in their normal state and are not affected by the special media required for SILAC or by the use of dialyzed serum. Furthermore, the same standard can be used for multiple samples and similar cell lines, and there is therefore no need to label all of them (29). Single cell line spike-in standard has been previously and efficiently used when multiple samples with high similarity were studied, or in cases where growing the cells in SILAC media was challenging (30).

Because SILAC-labeling of primary PTEC is not trivial and the several passages needed for the full incorporation of the labeled amino acids may lead to cell culture-induced loss of differentiation, we have employed spike-in SILAC to accurately quantify their proteome after sex hormone incubation. In particular, we SILAC-labeled immortalized human kidney HK-2 cells, and used their “heavy” proteome as internal standard to quantify the treated proteome from PTEC. Both PTEC and HK-2 are epithelial cells and originate from human renal proximal tubule. Thus, we reasoned that HK-2 proteome would allow us to accurately quantify a large proportion of PTEC proteome. In addition, HK-2 cells have been reported to show excellent labeling efficiency (31). However, HK-2 cells are immortalized cells and may differ from tubular cells in vivo in several important aspects, such as the changes in their proteome in response to certain stimuli (32, 33). We thus selected the primary PTEC cells for stimulation experiments, which were previously shown to respond to sex hormones (8, 14). Among all different cell types in the kidney, we performed our study in proximal tubular cells because tubules constitute most of the renal parenchymal mass, and tubular atrophy with interstitial fibrosis carries prognostic significance, and represents the common final pathway of most causes of CKD (36, 37).

To our knowledge, the effect of sex hormones on the proteome of human kidney cells has not been previously explored. In addition, we are the first to use a spike-in SILAC quantitative proteomic approach between two different renal cell lines. In this discovery-based study, we provide a detailed portrait of the key biological processes impaired by DHT and not EST. By performing bioinformatics analyses and the corresponding verification and validation experiments, we demonstrate: (1) a sex-specific regulation of three candidate proteins related to energy metabolism; and (2) an association between male sex and increased oxidative stress levels within the kidney. In this article we suggest for the first time that male sex hormones induce perturbations in the metabolism of the tubular cell that may ultimately lead to impaired oxidative stress and hypertrophy in the kidney and consequently increase the susceptibility to renal disease progression in males, especially in the context of diabetes. With our findings, we provide a novel link between male sex hormones and impaired energy metabolism in the kidney that may shed new
light on how to explore the mechanisms involved in the more rapid progression of CKD ascribed to male sex.

**EXPERIMENTAL PROCEDURES**

**Experimental Design and Statistical Rationale**—To cover the intrinsic variability associated to cell culture across passages, sex hormone stimulation experiments in PTEC were conducted in two different passages (passage 4 and 5). In each passage, incubation of PTEC with sex hormones or ethanol was performed in two separate experiments, aiming to obtain biological replicates that allowed us to achieve adequate reliability and consistency in further quantitative proteomics profiling.

**Cell Culture**—PTEC were purchased from Lonza Walkersville Inc (Walkersville, MD). They were cultured in custom-made Dulbecco’s modified Eagle’s medium (DMEM), and supplemented with 10% v/v dialyzed fetal bovine serum (FBS), 10 ng/ml EGF, 5 μg/ml transferrin, 5 μg/ml insulin, 0.05 μg hydrocortisone, 50 units/ml penicillin, and 50 μg/ml streptomycin, as previously described (38). Cells were serum starved for 18 h and treated with 100 nM DHT or EST (n = 4) or EST (n = 3) for 10 min (aiming to detect AKT and ERK phosphorylation as control experiments) or 8 h (aiming to activate both nongenomic and genomic signaling for proteomic analysis). Ethanol treated cells were used as controls (CONT, n = 4). After stimulation, cells were washed three times with PBS, harvested with trypsin, and snap-frozen at −80 °C until further analysis.

HK-2 cells used for SILAC labeling were provided by Dr. López Novoa (Instituto de Investigación Biomédica de Salamanca, Salamanca, Spain). They were cultured in DMEM/F12 (1:1) free of arginine, lysine, methionine and leucine (AthenaES, Baltimore, MD), and supplemented with 10% v/v dialyzed FBS, 50 units/ml penicillin, 50 μg/ml streptomycin, 2 mM glutamine, 5 μg/ml transferrin, 5 μg/μl insulin, 0.05 μg hydrocortisone, 1 mM T3 hormone, 10 ng/ml EGF, 147.5 mg/L heavy arginine (13C6), 91.25 mg/L heavy lysine (13C6), 20 mM iodoacetamide. Proteins were then alkylated with 20 mM iodoacetamide. Proteins were then digested with trypsin for 15 min at 15,000 rpm at 4 °C.

**Sample Preparation for Proteome Analysis**—Cell pellets from treated PTEC (light) and labeled HK-2 cells (heavy) were thawed on ice, resuspended in 200 μl of 0.1% w/v acid-labile detergent RapiGest SF (Waters, Milford, MA) in 25 mM ammonium bicarbonate, vortexed, and sonicated three times for 30 s. All lysates were centrifuged for 20 min at 15,000 × g at 4 °C. Total protein concentration was measured using a Coomassie (Bradford) protein assay reagent (Pierce, Waltham, MA). One hundred and fifty micrograms of protein from heavy HK-2 cells were spiked to 150 μg of each sample from light treated PTEC (1:1 mixing ratio) (Fig. 1). Proteins in detergent solution were denatured at 60 °C, and the disulfide bonds were reduced with 10 μm dithiothreitol. Following reduction, the samples were alkylated with 20 mM iodoacetamide. Proteins were then digested overnight at 37 °C with sequencing grade modified trypsin (Promega, Madison, WI). A trypsin/total protein ratio of 1:50 (w/w) was used. After digestion, RapiGest SF detergent was cleared with trifluoroacetic acid, 1% (v/v) final concentration, and samples were centrifuged for 15 min at 15,000 rpm at 4 °C.

**Proteome Analysis of DHT- and EST-stimulated PTEC Using Two-dimensional LC-MS/MS**—Upon removal of RapiGest, tryptic peptides were diluted to 500 μl with strong cationic exchange (SCX) mobile phase A (0.26 M formic acid in 5% v/v acetonitrile; pH 2) and loaded directly onto a 500 μl loop connected to a PolySULFOETHYL A column (2.1-mm inner diameter × 200 mm, 5 μm, 200 Å, The Nest Group Inc. (Southborough, MA)). The SCX chromatography and fractionation were performed on an HPLC system (Agilent 1100) using a 60 min two-step gradient. An elution buffer that contained all components of mobile phase A with the addition of 1 M ammonium formate was introduced at 10 min and increased to 20% at 30 min and then to 100% at 45 min. Fractions were collected every 3 min from the 20 min time point onward. The resulting 10 fractions (600 μl each) corresponding to chromatographic peaks of eluting peptides were collected. Peptides in each fraction were identified by LC-MS/MS as described previously (39). Briefly, peptides were extracted with OMIX C18 MB tips (Agilent Technologies, Lake Forest, CA), eluted in 3 μl of 65% v/v acetonitrile, diluted to 41 μl with 0.1% v/v formic acid in pure water, and loaded onto a 3.3 cm C18 pre-analytical column (IntegraFrit capillary, New Objective; inner diameter of 150 μm; 5 μm bead size; Agilent Pursuit C18, Agilent Technologies, Santa Clara, CA). Eluted peptides from the pre-analytical column were loaded onto a resolving analytical column with dimensions 15 cm × 75 μm ID (PicoTip emitter, 8 μm tip, New Objective Agilent Pursuit C18, 3 μm bead size). The trap and analytical columns were operated on the EASY-nLC1000 system (Thermo Fisher Scientific, San Jose, CA), and this liquid chromatography setup was coupled on line to a Q Exactive Plus hybrid quadrupole-Orbitrap mass spectrometer (Thermo Fisher Scientific, San Jose, CA) using a nano-ESI source (Proxeon Biosystems, Odense, Denmark). Each fraction was run using a 60 min gradient, and analyzed in a data-dependent mode in which a full MS1 scan acquisition from 400–1500 m/z with resolution 70,000 was acquired. This was followed by MS2 scan acquisition of the top 12 parent ions with resolution 17,500. The following parameters were enabled: monoisotopic precursor selection, charge state screening, and dynamic exclusion (enabled for 45 s), MS1 target value of 3e5 and maximum injection time (IT) of 100 ms, MS2 target value of 5e4 with maximum IT of 50 ms, isolation window of 1.6 Da, normalized collision energy (NCE) of 27, peptide match set to preferred, underfill ratio set to 2% (underfill ratio of 2e6), exclude isotopes set to ON. In addition, charge states of +1, >4, and unassigned charge states were not subjected to MS2 fragmentation. For protein identification and data analysis, Xcalibur software (version 3.0.63; Thermo Fisher) was utilized to generate RAW files of each MS run. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (1) partner repository (https://www.ebi.ac.uk/pride/archive/) with the data set identifier PXD003811.

**MS Data Analysis**—The MS files were processed with the MaxQuant software version 1.5.28.40 (40) and searched with Andromeda search engine (41) against the human UniProt database (42) (release 01_07_2015, 89,649 entries). The raw files from all biological replicates were analyzed simultaneously with MaxQuant (Fig. 1). To assess the false-positive rate, a reverse hit database was created by MaxQuant. The false discovery rates at the protein and peptide level were set to 1%. First search peptide tolerance was set to 20 ppm against a small “human-first-search” database for the purpose of mass recalibration and for main search precursor ion mass deviation of 4.5 ppm was allowed, while fragment mass deviation of 20 ppm was used. The minimum peptide length was set to 7 amino acids and strict specificity for trypsin cleavage was required, allowing up to two missed cleavage sites. Searches were performed with fixed carbamidomethylation of cysteines, variable oxidation of methionine and proline residues, and N-terminal acetylation. Multiplicity of 2 was used, selecting Arg (+6 Da) and Lys (+8 Da) as heavy labels. Re-
quantification and matching between runs was selected. Protein was identified with a minimum of one unique peptide. Quantification was performed using unmodified unique and razor peptides and a minimum of one counted ratio.

**SILAC Ratio Analysis**—The reverse and common contaminant hits were removed from MaxQuant output. Only proteins identified with at least one peptide uniquely assigned to the respective sequence were considered for the analysis. The distributions of all Log2 transformed H/L and H/L normalized ratios were normal (supplemental Fig. S1), thus the H/L normalized ratios of biological replicates were used to calculate DHT/CONT and EST/CONT ratios as described in Fig. 1. These ratios express the fold change between two different conditions and were calculated in all four experiments. Significance A of DHT/CONT and EST/CONT ratios was calculated for all the quantified proteins. A protein was considered to be significantly differentially regulated if its ratio was significant by significance A with $p < 0.01$ in at least two experiments. Proteins were eliminated if their ratios in the other two experiments showed significant changes in the opposite direction, defined as $\geq 2$-fold change in the opposite direction, or change in the opposite direction with $p < 0.05$. Only proteins with at least one unique peptide with two or more ratio counts were further pursued. In addition, if ratios of prototypic (unique) peptides of the same protein were changing in opposite directions, these proteins were also discarded for the validation study. A total of 76 proteins were found to be differentially regulated in at least one of the three comparisons (Table 1).

**Experimental Animals**—For the in vivo validation of the selected differentially expressed proteins found in SILAC experiments, we used 16-week-old C57BL/6 healthy female and male mice. To assess the effect of sex in the context of diabetes, we employed female and male streptozotocin (STZ)-induced mice. Diabetes was induced to 10-week-old mice following the High Dose STZ Induction Protocol for male streptozotocin (STZ)-induced mice. Diabetes was induced to 16-week-old C57BL/6 healthy female and male mice. To assess the differentially expressed proteins found in SILAC experiments, we used 16-week-old female and male diabetic Akita (Ins2WT/C96Y) mice. Five to eight animals were included in each experimental group. Mice were anesthetized with isoflurane and sacrificed by terminal surgery. Kidneys were removed, weighted, snap frozen, and kept at $-80^\circ$C until further analysis. Half of the right kidney was maintained in 10% formalin solution and paraffin embedded for histological studies. Mice were housed in ventilated cages with full access to chow and water at the Division of Comparative Medicine at the University of Toronto (control and Akita mice), or at the Animal Facility of Barcelona Biomedical Research Park (STZ-induced mice). All experiments were conducted under the guidelines of the University of Toronto Animal Care Committee and the Ethical Committee of Animal Experimentation of Barcelona Biomedical Research Park (CEEA-PRBB).

**Western Blot**—Proteins from sex hormone-treated cell pellets and mouse kidney cortex were solubilized in modified RIPA buffer (150 mM sodium chloride, 50 mM Tris-HCl (pH 7.4), 1 mM EDTA, 1% v/v Triton X-100, 1% w/v sodium deoxycholic acid, 0.1% v/v SDS) and extracted by sonication. Protein concentration was determined using a Coomassie (Bradford) protein assay reagent (Pierce, Waltham, MA). 15 μg of protein were loaded onto 12% acrylamide gels and separated by SDS-PAGE. Membranes were incubated with antibodies to pSer473 AKT (193H12, Cell Signaling, Danvers, MA), p44/42 ERK1/2 (9102, Cell Signaling), GPIX (PA5–26787, Thermo Scientific), HADHA (ab54447, Abcam, Cambridge, UK). The secondary antibody was anti-rabbit antibody developed in goat (sc-2004, Santa Cruz Biotechnology, Delaware Ave). Control for protein loading was performed by reblotting proteins for β-actin using a mouse monoclonal antibody (A1978, Sigma) and an anti-mouse secondary antibody (sc-3697, Santa Cruz Biotechnology). Following detection, bands were quantified by densitometry with Image J software. For these in vitro verification and in vivo validation experiments, three to six PTEC replicates per condition and five to ten animals per group were studied, respectively.

**Kidney Histology: Oxidative Stress and Immunolocalization Studies**—For all kidney histology studies, paraffin blocks were cut into 3 μm sections, deparaffinized in xylene and rehydrated through graded alcohols.

Oxidative stress is generally identified by indirect markers such as peroxynitrite production. For this reason, nitrotyrosine staining was used to evaluate superoxide and peroxynitrite levels in renal tissues as previously described (45). Deparaffinized samples were boiled in 10 mM sodium citrate solution (pH 6.0) for antigen retrieval. Kidney sections were stained with rabbit antinitrotyrosine antibody (1:500, Millipore, Billerica, MA). Fifteen microphotographs at $\times 40$ were taken for each sample, and brown-stained areas were quantified with ImageJ software. Data are expressed as percentage of positive area. All analyses were performed in a blinded fashion. For these experiments, six to eight animals per group were studied.

To localize top candidate proteins from our proteomic data set in the murine kidney, immunohistochemistry staining was also performed for HADHA, GPIX and GNPNAT1. Samples were boiled in 10 mM sodium citrate solution (pH 6.0) for GNPNAT1 staining, whereas antigen retrieval for HADHA and GPIX was performed in 10 mM Trizma® base, 0.5 M EDTA, 0.05% Tween buffer (pH 9.0). Sections were incubated with rabbit primary antibodies for HADHA (1:1000; ab54447, Abcam), GPIX (1:1000; PA5–26787, Thermo Scientific) or GNPNAT1 (1:500; HPA044647, Atlas Antibodies). HRP-conjugated anti-rabbit (EnVisionTM, Dako, Santa Clara, CA) was used as secondary antibody. Slides were counterstained with hematoxylin. For these experiments, three to six animals per group were studied.

**Statistical and Bioinformatics Analyses**—For the in vitro proteomics analysis, Perseus software (version 1.5.1.6) was used for calculation of significance A of protein ratios. Benjamini-Hochberg FDR procedure was used to calculate $p$ values and $p < 0.01$ was considered statistically significant. The corresponding gene IDs of the 76 sex hormone-regulated proteins were used to perform the gene ontology enrichment analyses using BINGO plugin in Cytoscape software (version 3.2.1) (46, 47). For the statistical analyses in BINGO, hypergeometric test with Benjamini-Hochberg correction was used and corrected $p < 0.05$ was considered significant. In addition, Enrichment Map plugin was used to assess the significantly enriched functional categories among our DHT-regulated proteins and other previous data sets reflecting the effect of diabetes (48) on kidney transcriptome. These data sets were publicly available in GEO and Nephromine online platforms. Hypergeometric test with FDR = 0.05 and Jaccard coefficient cutoff = 0.25 was set to build the enrichment map. Hypergeometric test cutoff of 0.05 was set for post-analysis involving comparison of our data set with the diabetic transcriptional data set above.

For the in vitro verification of SILAC ratios and the additional in vitro studies, densitometries of Western blot bands are reported as mean ± S.E. For the in vivo validation studies, values for physiological parameters, Western blot analysis and quantification of nitrotyrosine positive area are also reported as mean ± S.E. Statistical comparison between experimental groups was performed using a two-tailed Mann-Whitney U test. Spearman’s correlation coefficient was calculated when appropriate. $p$ values <0.05 were considered statistically significant.
RESULTS

Sex Hormone Stimulation of PTEC—As shown in Fig. 1, PTEC at the 4th and 5th passage were split into T25 flasks, deprived of serum for 18 h and treated with ethanol (CONT), DHT, or EST for 8 h. Experiments 1 and 2 were performed at passage 4, whereas experiments 3 and 4 were performed at passage 5. Cell lysates were collected and lysed as described under Experimental Procedures. As a control experiment, we determined in each passage levels of phosphorylated AKT and ERK after 10min stimulation with DHT or EST. Western blot analysis showed that both sex hormones led to phosphorylation of ERK (Fig. 2A,B) and AKT (Fig. 2A,2C) in comparison with the control cells. This control experiment demonstrated that DHT and EST engaged the androgen and estrogen receptor, respectively, leading to predictable signaling events in the cells. Although these phosphorylation events are two of many initiated by sex hormones, this was an important confirmation that PTEC exhibited a biological response to DHT and EST.

Spike-in SILAC Quantitative Analysis of Sex Hormone-treated PTEC Proteome—For accurate, in-depth characterization of the proteomes in these different conditions we employed the advantages of spike-in SILAC (29, 30). We thus labeled an immortalized proximal tubular cell line (HK-2) with heavy-SILAC labels, and used it in spike-in experiments for relative quantification. We first evaluated the efficiency of labeling of HK-2 cells. Although even five doublings demonstrated excellent labeling efficiency, we selected eight doublings, which showed the highest labeling efficiency (99.66%) (supplemental Table S1). Proteins from these cells were spiked into the lysate of each of the samples described above, acting as an internal heavy standard to quantify the light proteome of PTEC under each experimental condition. Mixed proteomes were then analyzed by online liquid chromatography mass spectrometry (LC MS/MS) (Fig. 1). EST-treated PTEC in experiment 1 ultimately showed poor protein recovery after peptide fractionation, and was thus eliminated from analysis. MS analysis identified 5111 proteins. Of those, 5043 were accurately quantified in at least one sample, and 3958 were accurately quantified in at least one replicate of each condition (supplemental Table S2). Log2 transformed ratios and their distributions were evaluated. Histograms showed normal distributions for the two ratios (Fig. 3B,3C). However, histograms for DHT/CONT and EST/CONT were modestly shifted to the right (Fig. 3B,3C), even after normalization. Spearman’s rank correlations were calculated between H/L normalized ratios of two different samples, considering all possible combinations among the 11 samples. As shown in supplemental Table S3, correlation coefficients between biological replicates from the same passage were higher than the coefficients between biological replicates from different passages. This observation was true not only in treated but also

Fig. 1. Experimental scheme. The figure shows a simplified workflow, including sex hormone treatment to PTEC, SILAC labeling of HK-2 cells, 1:1 mixing of labeled and nonlabeled proteins, protein digestion, SCX fractionation followed by LC-MS/MS, data analysis by MaxQuant, assignment of heavy/light protein ratios, calculation of DHT/CONT and EST/CONT ratios, selection of differentially regulated proteins, validation studies and bioinformatics analyses.

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in control cells, indicating an intrinsic variability of the PTEC proteome across passages. Similar findings had been reported previously (30). We decided to only consider those proteins that were differentially regulated in both passages, or that at least, did not change significantly in the opposite direction in different passages.

Based on significance A, a total of 76 proteins were differentially expressed between 2 conditions (Fig. 3A, Table I). The complete list of these proteins and their corresponding ratios in each experiment is in supplemental Table S4. Individual peptide ratios for all differentially regulated candidates were examined manually. Sixty proteins were found to
| Gene names   | DHT/CONT | EST/CONT |
|-------------|----------|----------|
|             | Median   | S.E.     | Significant | Median | S.E. | Significant |
| ALDH1B1     | 7.439±1  | 0.072    | YES (Upregulated) | 0.852  | 0.214 | NO          |
| ALDH4A1     | 6.806    | 2.960    | YES (Upregulated) | 1.575  | 0.444 | NO          |
| C14orf159   | 8.330    | 3.011    | YES (Upregulated) | 0.981  | 0.049 | NO          |
| C19orf25    | 5.837    | 1.241    | YES (Upregulated) | 0.495  | NaN   | NO          |
| CAD         | 3.704    | 1.034    | YES (Upregulated) | 1.100  | 14.303 | NO          |
| CALR        | 7.818    | 1.205    | YES (Upregulated) | 0.981  | 0.576 | NO          |
| CASP6       | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDC2;CDK1   | 20.344   | 10.175   | YES (Upregulated) | 1.386  | 1.118 | NO          |
| CDK4        | 1.664    | 0.095    | NO            | 4.016  | 2.809 | YES (Upregulated) |
| CDKN2AP3    | 8.448    | 0.470    | YES (Upregulated) | 0.947  | 0.021 | NO          |
| CDK5RAP3    | 6.325    | 0.988    | YES (Upregulated) | NaN    | NaN   | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
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| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.837    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
| CDK5RAP3    | 5.212    | 1.577    | YES (Upregulated) | 1.232  | 0.428 | NO          |
be differentially expressed after treatment with DHT, compared with control cells. Treatment with EST resulted in 18 differentially expressed proteins compared with control cells. As shown in the Venn diagram, 2 proteins were found to be differentially regulated by both male and female sex hormones. Among these 76 proteins, cyclin-dependent kinases 1 (CDK1) and 4 (CDK4) were regulated by DHT and EST, respectively (supplemental Table S4A, S4B). Supporting our findings, CDK1 has been reported to phosphorylate AR preventing its degradation and thereby increasing AR stability and protein expression (49–51), whereas CDK4 was up-regulated after 17β-estradiol treatment in MCF-7 breast cancer cells (52) and rat vascular smooth muscle cells (53).

Log2 transformed normalized ratios are shown as heat maps for all DHT/CONT and EST/CONT differentially regulated proteins (Fig. 4). Interestingly, all DHT-regulated proteins were up-regulated relative to control (Fig. 4A, supplemental Table S4). Similarly, only RING-finger type E3 ubiquitin ligase (UHRF1), an important nuclear protein that plays critical roles in regulating various processes such as DNA methylation maintenance and G1/S transition (54), was found to be downregulated after EST treatment (Fig. 4B, supplemental Table S4). The intrinsic variability across passages is also reflected in the values of the ratios between conditions, as replicates 1 and 2 represent passage 4, and replicates 3 and 4 represent passage 5.

**Gene Ontology and Molecular Pathway Analysis**—We next evaluated Gene Ontologies of our proteins differentially regulated by sex hormones. Some of the most relevant and significant biological processes enriched among all DHT-regulated proteins were fatty acid beta-oxidation (FAO), glutamine metabolic process, N-acetylglucosamine metabolic process, glucose catabolic process, tricarboxylic acid (TCA) cycle, and response to hydrogen peroxide, indicating DHT-induced alterations in energy metabolism (Fig. 5, supplemental Table S5). In this sense, gene ontology revealed mitochondrion as the most enriched cellular organelle (supplemental Table S5). Steroid hormone receptor signaling biological process was also significantly enriched by DHT treatment (Fig. 5). In addition, positive regulators of androgen signaling pathway such as protein deglycase DJ-1 ( PARK7) (55, 56) and calreticulin (CALR) (57) were up-regulated, providing further confirmation of the PTEC biological response to DHT treatment. Biological processes linked to apoptosis were also significantly enriched among EST-regulated proteins (supplemental Table S5).

Dysregulated metabolism is a key factor in many kidney diseases (58). In DKD, increased intracellular glucose metabolism leads to excessive activation of hexosamine and protein kinase C pathways, as well as accentuated non-enzymatic glucose oxidation to advanced glycation end products (59). As a consequence, mitochondrial oxidative phosphorylation and reactive oxygen species (ROS) release are promoted. Increased FAO and acetyl-CoA production in TCA cycle accentuate these processes in the mitochondria (60). DHT alone was able to dysregulate processes that are also altered by diabetes such as glucose metabolism, N-acetylglucosamine metabolism (a relevant step in hexosamine biosynthetic pathway), HBP, and FAO. Thus, we selected GPI, GPNPNT1 and HADHA as candidates for validation that represented each one of these three processes. It is remarkable that GPNPNT1 protein expression was increased in both DHT- and EST-treated PTEC, compared with control cells. Fig. 6 illustrates the role of the most representative up-regulated proteins in cellular energy metabolism pathways.

**In Vitro and In Vivo Validation**—We next aimed to confirm SILAC ratios by Western blot analysis in the same protein extracts used for the proteomic study. In consistence with our findings from SILAC, Western blot analysis in treated PTEC confirmed a significant up-regulation of HADHA, GPI and GPNPNT1 by DHT. To a lesser extent, these proteins were also up-regulated by EST (Fig. 7A–7D). We next examined whether these observations were consistent in new PTEC stimulation experiments, at both passages studied in SILAC experiments. Cells at passage 4 and 5 were thus serum-starved for 18 h and then treated with control, DHT or EST for 8h (n = 5–6 in each passage). Expression of proteins HADHA, GPI and GPNPNT1 was numerically increased in DHT-treated
cells at passage 4 (supplemental Table S6). At passage 5, both DHT- and EST-treated PTEC exhibited a significant increase in GNPNAT1 protein expression, a result concordant with SILAC protein expression data. However, HADHA and GPI at passage 5 were not up-regulated by DHT, simulating our SILAC experiments (supplemental Table S4).

**Fig. 4.** Heat map representation of sex-regulated proteins in PTEC. Heat maps were constructed using Perseus software and reflect the value of Log2 transformed normalized DHT/CONT (A) and EST/CONT (B) ratios for each individual protein (rows) in each experiment (columns). The color intensity of red and green boxes correlates with the degree of protein over- and under expression, respectively. A gray box indicates that the ratio was not assessed for that particular protein in that particular experiment.

**Fig. 5.** Enriched biological processes among the 60 proteins differentially regulated by DHT. Bar graphs illustrating the Log2 transformed DHT/CONT and EST/CONT ratios of several DHT-regulated proteins involved in the most representative, informative and significantly enriched biological processes according to gene ontology analysis using BinGO plugin in Cytoscape. An asterisk indicate that these protein was significant by significance A with $p < 0.01$ in at least 2 experiments.
We next aimed to examine whether our \textit{in vitro} observations in PTEC have relevance \textit{in vivo}. With this purpose, we determined expression of HADHA, GPI, and GNPNAT1 proteins in the kidney of healthy female and male mice by immunohistochemistry. As shown in supplemental Fig. S2, both sexes showed strong positive staining for all three candidates in cells from the proximal tubule and other tubular segments. These findings suggest that changes in renal HADHA, GPI and

**Fig. 6. Role of DHT- and EST-regulated proteins in energy metabolism.** The figure illustrates a simplification of several processes related to energy metabolism and mitochondrial compartment. Proteins that appeared to be up-regulated by DHT (orange), or both DHT and EST (blue) are also represented. GAD-3P, glyceraldehyde-3-phosphate; FFA, free fatty acids; HBP, hexosamine biosynthetic pathway; Gin, glutamine; Glu, glutamate; ROS, reactive oxygen species; $\alpha$-KG, alpha-ketoglutarate; OAA, oxaloacetate.

**Fig. 7. \textit{In vitro} and \textit{in vivo} validation studies for HADHA, GPI, and GNPNAT1.** A, shows the immunoblots representing HADHA, GPI, and GNPNAT1 protein expression in the same treated PTEC used for the spike in SILAC study. Intensities for HADHA (B), GPI (C), and GNPNAT1 (D) were calculated and normalized to $\beta$-actin (B, C, D). E, shows the immunoblots representing HADHA, GPI, and GNPNAT1 protein expression in the kidney of healthy female and male mice. Densitometry analysis of each band was performed using Image J software. Intensities for HADHA (F), GPI (G) and GNPNAT1 (H) were calculated and normalized to $\beta$-actin (F, G, H). Data are expressed as means $\pm$ S.E. *p < 0.05 compared with control cells.
GNPNAT1 across sexes will be mostly ascribed to sex-specific regulation of these proteins in the tubular compartment, strengthening the significance of our *in vitro* observations. In concordance with our results from SILAC proteomics, Western blot analyses revealed that, under physiological conditions, renal HADHA and GPI were significantly increased in males, compared with females. In contrast, GNPNAT1 was not significantly different between female and male mice (Fig. 7E–7H). Expression of HADHA and GPI renal proteins strongly and significantly correlated with the animal kidney weights (r = 0.675, p = 0.023; r = 0.772, p = 0.005; respectively) which were significantly increased in males, compared with females (supplemental Table S7A).

**Biological Significance: Validation of Top Candidates in the Diabetic Kidney**—Because our top candidate proteins belong to biological processes critical for DKD (59), namely glycolysis, HBP and FAO, we evaluated the effect of sex on renal HADHA, GPI, and GNPNAT1 protein expression in female and male mice of two different models of diabetes: the STZ-induced and the Akita (Ins2WT/C96Y) models of type 1 diabetes. We specifically chose diabetes as an experimental model because of the significant role of metabolism and the poorly understood impact of sex on the natural history of diabetic nephropathy. Of importance, both STZ-induced and Ins2WT/C96Y diabetic males showed accentuated hyperglycemia compared with females, as well as a more severe renal hypertrophy in terms of kidney weight and kidney to body weight ratio (supplemental Table S7B–S7C). In the setting of type 1 diabetes, male sex was associated to increased HADHA, GPI and GNPNAT1 protein expression, compared with females (Fig. 8). Specifically, renal HADHA was significantly increased, whereas GPI and GNPNAT1 were modestly higher, in STZ males than in females (Fig. 8A–8D). We demonstrated that the expression of these three proteins was predominantly in renal tubules, similar to the control animals (Fig. 8E). These findings were further validated in diabetic Akita mice, a genetic model of type 1 diabetes. Akita males showed significantly higher protein levels of HADHA and GNPNAT1, as well as a modest increase in GPI protein expression (Fig. 8F–8I). Correlation analyses in all the studied diabetic mice revealed a moderate but significant association between HADHA protein expression and kidney weight (r = 0.471, p = 0.013).

**Enriched Functional Category Analysis**—We explored enriched biological processes in our SILAC data that were reproducible in prior studies of diabetes to assess the associ-
Fig. 9. **Functional enrichment of oxidative stress among our DHT-regulated proteins.** The enrichment map (EM) was constructed imposing our list of DHT-regulated proteins to a renal transcriptomic data set from human diabetic kidneys in Cytoscape (A). EM is depicted in the left bottom, in the panel. Each circle represents an enriched GO term, with red color indicating significance. Green edges indicate that up-regulated proteins are shared by the two GO terms they connect. The analysis revealed that functional groups associated with oxidative stress (A) were shared with our data set. The pink edge represents overlap between the target gene set and the enriched biological processes from our experiment. To validate the link between male sex and renal oxidative stress, nitrotyrosine immunostaining was used to evaluate superoxide and peroxynitrite levels in kidney sections from female and male, control and STZ-diabetic mice (B). Scale Bar = 20 μm. Original magnification ×40. Brown-stained areas were quantified with ImageJ software (C). Data are expressed as means ± S.E. of positive area fraction (%). *p < 0.05 compared with females.

**DISCUSSION**

This study was designed to shed new light on the biological processes and cellular compartments involved in the deleterious effects of androgens in CKD by studying proteins regulated by male and female sex hormones in the kidney. Our specific goal was to capture a DTH and EST "protein signature" in vitro that reflected an early and consistent response of the PTEC to sex hormones.

To define the proteome of sex hormone-stimulated PTEC, we utilized SILAC methodology, which is the current standard for relative quantification of cellular proteomes by MS (25). We identified and quantified over 5000 proteins, thus providing a unique depth of insight into PTEC responses to sex hormones. The strengths of our approach include good proteome coverage, use of instruments with high sensitivity and accuracy, consideration of the effect of cell passage on protein expression, and control experiments to minimize false-positive hits. To the best of our knowledge, this is the first effort to date to characterize proteomic responses of human kidney cells to DHT and EST stimulation.

We first noticed that among our DHT-differentially regulated proteins, many were enzymes that have been shown to play a role in CDK and DKD. As an example, in cultured fibroblasts from type 1 diabetic patients with and without nephropathy, mRNA levels of GPI and other enzymes of the glycolytic pathway were found to be increased in the context of DKD (62). In turn, HADHA and HADHB, as subunits of the mitochondrial trifunctional protein, are involved not only in FAO but also in molecular events relevant to kidney disease, such as lysine acetylation (63), and modulation of renin expression (64), respectively. In addition, GNPNAT1 (which catalyzes a
reaction critical for HBP activation) participates in proliferative and hypertrophic processes associated with diabetes (65).

Our next aim was to use systems biology to demonstrate biological processes and molecular pathways fundamental in PTEC response to DHT and EST. Three of the dominant biological processes identified among our 60 DHT/CONT differentially regulated proteins were glucose catabolic process, N-acetylglucosamine metabolic process, and FAO, and one of the most enriched cellular compartments was the mitochondrion. In concordance with our results from SILAC proteomics and MS, validation experiments by Western blot confirmed that GPI and HADHA protein levels were augmented in DHT-treated renal cells and in kidneys from C57BL/6 males, supporting the idea that androgens impair carbohydrate and fatty acid metabolism in the kidney through the action of DHT.

It is known that alterations in the processing of carbohydrates and fatty acids may ultimately compromise the mitochondrial function and the oxidative stress status within the cell (59). Thus, the increase in glycolytic enzymes and proteins responsible for FAO and glutamine metabolism after DHT treatment may explain the up-regulation of several TCA cycle enzymes such as malate dehydrogenase, as well as the increased expression of many proteins related to the mitochondrial compartment (Fig. 6). Furthermore, in this work, we demonstrated that perturbed energy metabolism and up-regulation of mitochondrial proteins by male sex hormones is associated with increased oxidative stress levels in male kidneys under non-pathological conditions. Therefore, our findings suggest that these processes may play a critical role in mediating the deleterious effects of androgens in the physiology of the renal cells. In this sense, DHT has been shown to alter glycolytic metabolism and impair mitochondrial function in other cell types (66, 67). As recently reported by Wang et al., pancreatic islets from DHT-treated rats showed significant changes in the levels of several key genes involved in mitochondrial biogenesis, mitochondrial oxygen consumption rate, and ATP production. The authors also demonstrated that androgens can directly impair beta cell function by inducing mitochondrial dysfunction in vitro in an AR-dependent manner (66). In a different context, DHT enhanced glucose consumption and lactate export in prostate cancer cells (67).

We speculate that increased metabolism of sugar, lipid and amino acids in DHT-treated PTEC induced an imbalance on the levels of several metabolites such as pyruvate, α-ketoglutarate and acetyl-CoA. Thus, the increase in the protein expression of pyruvate dehydrogenase and other enzymes involved in N-acetylglucosamine metabolism and TCA cycle may be explained, at least in part, as an attempt of the tubular cell to counterbalance the abnormal accumulation of these metabolites. For example, GNPNAT1, one of the key enzymes of the HBP that was found to be significantly increased by DHT in both SILAC and Western blot experiments, uses acetyl-CoA as the acetyl group donor for the conversion of glucose-6-phosphate to N-acetylglucosamine-6-phosphate.

The enrichment in energy metabolism processes after DHT stimulation, together with the association between male sex and renal oxidative stress levels, pointed out a possible activation of the classic and well defined mitochondrial-dependent apoptotic pathways (68–70). It has been reported that long exposure to testosterone (24 h to 48 h) promoted apoptosis in HK-2 cells and PTEC (8, 13). We decided to treat PTEC for only 8 h, a time that we considered long enough to observe both genomic and nongenomic actions of sex hormones, but still prevented us from the interference of androgen-induced apoptosis and cell death in our cell culture. Thus, the lack of a more significant number of apoptotic players among our DHT proteomic signature despite the increase in proteins related to energy metabolism may be indicative of an early, adaptive response of the cell to DHT signaling, preceding the gradual development of pathological consequences because of a continuous impaired metabolism and subsequent accumulation of ROS. In this sense, protein expression of the antioxidant enzyme SOD2 was also found to be up-regulated by DHT, together with CASP6 and PARK7 as part of the response to hydrogen peroxide. SOD2 is responsible for the conversion of superoxide anion to hydrogen peroxide (71). Decreases in SOD2 expression and activity have been associated with exacerbated levels of oxidative stress and mitochondrial dependent apoptosis in HK-2 cells and murine kidneys after long exposures to high glucose, angiotensin II (72), or nephrotoxic reagents such as cisplatin (73). From this point of view, evidence for the decreased but increased SOD2 after 8 h of DHT treatment in PTEC suggests that oxidative stress and apoptosis are partially blunted by a physiological early response of the tubular cell. Furthermore, several proteins related to mitochondrial p53-mediated apoptosis, namely BAX, BCLAF1, BCL2L13, TP53BP1, and TP53BP2, were detected and quantified in all four experiments but were not affected by DHT treatment (data available in Proteome Xchange).

Cumulative evidence has linked altered energy metabolism and mitochondrial modification to the pathogenesis of diabetes and its complications (74). In this sense, proteomic efforts in different tissues from mouse models of type 1 diabetes (T1DM) detected increased expression of several mitochondrial proteins related to FAO, TCA cycle and oxidative stress (75, 76). Within the diabetic kidney, label-free quantification of the mitochondrial proteome revealed increased FAO protein content and induction of TCA cycle enzymes (76). In this sense, renal mitochondrial complex III of the respiratory chain, one of the major sites for ROS generation (71), was found to be altered in the early stage of STZ-induced T1DM in rats (77). We provide strong evidence supporting that these metabolic processes relevant to diabetes are impaired by male sex hormones and increased in diabetic males, compared with females. In addition, the increase in renal oxidative stress
levels in diabetic males indicates that these alterations may ultimately lead to a more rapid kidney disease progression in males. In accordance with our results, male sex has been associated with higher urinary and kidney levels of oxidative stress in animal models of hypertension (4) and renal ischemia (78), respectively. To our knowledge, we are the first to report a sexual dimorphism in renal nitrotyrosine levels in the diabetic kidney.

As defended by Forbes and coauthors, it is important for the cells to maintain glucose homeostasis when exposed to hyperglycemic conditions by reducing the transport of glucose inside the cells. However, certain cell groups are unable to decrease glucose concentration, and are thus susceptible to damage (79). From this perspective, it is conceivable that several molecular pathways related to energy metabolism are activated in the tubular cell as a mechanism to compensate the tremendous increase in glucose influx under diabetic conditions. Among these pathways, the role of HBP in the pathophysiology of diabetic cardiorenal disease has been extensively studied (80–82). UDP-GlcNAc, the donor sugar for O-GlcNAcylation of proteins, is synthesized from glucose, glutamine, and UTP via the HBP. In turn, GNPNAT1 requires acetyl-CoA, which can be generated through FAO or transformation of citrate from TCA cycle by ATP-citrate lyase activity. Thus, it is generally accepted that HBP sits at the nexus of glucose, nitrogen, fatty acid and nucleic acid metabolism, which are altered in the context of diabetes (83). According to our SILAC data, these metabolic pathways were enhanced in DHT- but not EST-treated PTEC. This may explain the fact that renal GNPNAT1 appeared to be higher in our diabetic but not control male mice as compared with females.

It is of importance that diabetic male mice developed renal hypertrophy and accentuated hyperglycemia in comparison to diabetic females. In addition, significant correlations between renal HADHA and GPI protein expression and kidney weight were found in the present study, even under non-diabetic conditions. In this sense, metabolic remodeling has been extensively associated to maladaptive hypertrophic processes in other organs such as the heart (84), the pancreas (85) and the adipose tissue (86). In particular, remodeling of glucose (87) and fatty acid (88) metabolism promoted ventricular hypertrophy in experimental models of heart dysfunction. When accompanied by increased oxidative stress levels, these hypertrophic changes are more likely to become maladaptive and lead to organ failure (89). In the endocrine pancreas of mice exposed to a high-fat diet, metabolic changes associated to hypertrophy were found to be more accentuated in males (85). Alterations in processes related to mitochondrial biogenesis also play a role in hypertrophy (90). Interestingly, these processes can be differentially regulated by female and male sex hormones (91). In the kidney, increased amino acid delivery to the proximal tubule cells has been associated with renal growth and hypertrophy; regulation of molecular mechanisms such as AKT-pathways, that are also regulated by sex hormones, play a role in these renal alterations (92). It has been proposed that HBP activation leading to O-GlcNAcylation is also involved in hypertrophy, especially under hyperglycemic conditions (65). Therefore, we presume that glucose influx into HBP was augmented in the kidneys of diabetic males, probably contributing to the increase in protein expression of renal GNPNAT1 and the accentuated renal hypertrophy. Overall, strong experimental evidence demonstrating a link between these hypertrophic processes and perturbations in bioenergetics in the kidney is still lacking. With our data, we reinforce the idea that the maladaptive hypertrophy as a consequence of metabolic remodeling observed in other organs under certain pathological conditions also occurs in the kidney, and in a sex-specific manner.

To date, little is known about the effect of sex on energy metabolism within the kidney. In this work, we are the first to provide strong evidence that, under physiologic conditions, DHT and male sex promote higher activation of energy metabolism in the tubular cell and in the renal cortex. We also demonstrate that renal expression of the three candidates representing glycolysis, N-acetylgluosamine metabolism and FAO was increased in males in the context of T1DM. Further bioinformatics analyses and validation experiments allowed us to link this effect of male sex hormones in renal energy metabolism with accentuated renal hypertrophy and oxidative stress levels. For the first time, we suggest that altered metabolic activity in male kidneys under physiologic conditions confer a major susceptibility to develop renal complications, especially in the context of diabetes.

The strengths of our work include (1) the use of a solid, robust and accurate quantitative proteomic approach that was not previously used in renal cells; (2) good proteome coverage by employing a mass spectrometer with high sensitivity and accuracy, (3) control experiments to minimize false-positive hits, (4) the use of two different models of DKD for validation; (5) the inclusion of publically available data on renal transcriptomics in our systems biology analyses, and (6) the discovery of a new, biologically relevant link between male sex and perturbed renal energy metabolism.

Despite the novelty of our findings, our study has several limitations. Most significantly, we studied PTEC responses to sex hormones at a single time point, and the expression levels of proteins are likely to be dynamic. Although we studied a single time point, the variety of identified processes has been implicated to CKD and DKD processes. In turn, the intrinsic variability of PTEC proteome across passages may have probably contributed to the relatively low number of differentially expressed proteins between conditions. The MS analysis of EST-treated PTEC in passage 4 was limited to a single experiment and thus could contribute only marginally to the overall findings.

In conclusion, DHT alone led to dysregulated metabolic processes that are also altered in the diabetic kidney. These
processes, including glucose metabolism, HBP and FAO, are associated with diabetes and may represent the link to understand the more rapid progression of CKD in males. Top candidate proteins representative of each of these processes were verified and validated in sex hormone-treated PTEC and in male and female control and diabetic mice. Specifically, sex-specific regulation of GPI, HADHA, and GNPNAT1 was demonstrated in vitro and in vivo. To the best of our knowledge, we are the first to demonstrate that GPI, HADHA, and GNPNAT1 are responsive to androgens. Our results suggest that detrimental effects of androgens in diabetic nephropathy and other kidney diseases are mediated, at least in part, by altered energy metabolism within the tubular cell (Fig. 10).

Future studies focused on evaluating the influence of male sex hormones on the connection between these metabolic processes, together with other “omics” efforts in DKD approached from a “sex-specific perspective,” may potentially shed new light on the identification of new serum and urine biomarkers for sex-directed therapies in renal disease.

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