Grade-Dependent Metabolic Reprogramming in Kidney Cancer Revealed by Combined Proteomics and Metabolomics Analysis

Hiromi I. Wettersten, A. Ari Hakimi, Dexter Morin, Cristina Bianchi, Megan E. Johnstone, Dallas R. Donohoe, Josephine F. Trott, Omran Abu Aboud, Steven Stirdivant, Bruce Neri, Robert Wolfert, Benjamin Stewart, Roberto Perego, James J. Hsieh, and Robert H. Weiss

Abstract

Kidney cancer [or renal cell carcinoma (RCC)] is known as “the internist’s tumor” because it has protean systemic manifestations, suggesting that it utilizes complex, nonphysiologic metabolic pathways. Given the increasing incidence of this cancer and its lack of effective therapeutic targets, we undertook an extensive analysis of human RCC tissue employing combined grade-dependent proteomics and metabolomics analysis to determine how metabolic reprogramming occurring in this disease allows it to escape available therapeutic approaches. After validation experiments in RCC cell lines that were wild-type or mutant for the Von Hippel-Lindau tumor suppressor, in characterizing higher-grade tumors, we found that the Warburg effect is relatively more prominent at the expense of the tricarboxylic acid cycle and oxidative metabolism in general. Further, we found that the glutamine metabolism pathway acts to inhibit reactive oxygen species, as evidenced by an upregulated glutathione pathway, whereas the β-oxidation pathway is inhibited, leading to increased fatty acylcarnitines. In support of findings from previous urine metabolomics analyses, we also documented tryptophan catabolism associated with immune suppression, which was highly represented in RCC compared with other metabolic pathways. Together, our results offer a rationale to evaluate novel antimitabolic treatment strategies being developed in other disease settings as therapeutic strategies in RCC.

Introduction

Examination of clinical materials using several distinct and complementary techniques has the potential to yield new perspectives in physiology and pathophysiology that might not be evident using each technique in isolation. With the acceptance of mainstream omics technologies (1), it has become possible to probe the molecular biology of disease with a high degree of resolution. However, due to the highly specialized nature of each omics technique and its associated analysis, each investigator tends to have a relatively narrow range of expertise directed toward but a single omics strategy; for this reason, it is rare to find a merging of several of these powerful techniques in a single study. Indeed, such single omics studies can be problematic for interpretation, for example, the quantity of proteins (e.g., enzymes) as the output of proteomics does not always correlate with their activities, and many gene transcripts undergo posttranscriptional processing. It is also common to see publications that focus on technical and analytical aspects of the omics process at the expense of the overarching biologic interpretation. Because the interpretation of omics data can be easily misconstrued due to covariant and systemic effects that are not biologically relevant, it is critical to validate the hypotheses built upon the omics data. For these reasons, it is a logical extension in this field to combine data from several complementary omics techniques to attempt to eliminate some of the bias inherent in a single method, and it is critical to validate the proposed hypotheses that come out of the omics data both in vitro and in vivo (2, 3).

The vast majority of published basic research in oncology is based on the behavior of commercially available cell lines. In these situations, experiments have been performed on cells representing predominantly a single (usually high) grade of cancer. In a parallel situation, authors of studies using in vivo xenografts, derived from immortalized cell lines, frequently ignore original
tumor grade information or that which could be inferred from cell line characteristics. It is becoming abundantly clear from work in our and other laboratories that the biology of tumors of different grades is markedly distinct, at least in renal cell carcinoma (RCC; refs. 4, 5). Indeed, work in our laboratory is focused on determining whether personalized therapy based on such grade classification may have clinical utility, a concept that is often foreign to clinicians. The utility of integrating multiple omics approaches is especially important when evaluating grade dependence with respect to therapy, because of the subtleties that can be difficult to recognize in comparing morphologically similar tissues.

In our continuing pursuit of physiologic alterations that occur in kidney cancer, which may enable us to discover and validate new biomarkers and targets for this highly morbid disease, we have previously utilized both metabolomics and proteomics separately in a variety of biofluids and tissues (4–8). In the present study, for the reasons discussed above, we have merged the proteomics and metabolomics datasets derived from human RCC tumors of distinct Fuhrman grade in order to expand the pathway data previously obtained (4, 5); for validation purposes, we utilize several commercial cell lines, both with and without Von Hippel-Lindau (VHL) mutations, because VHL status affects energy metabolism through hypoxia-inducible factor (HIF) regulation. Using this approach, we show here incontrovertible and validated evidence of grade-dependent metabolic reprogramming in human RCC, characterized by increasing utilization of the Warburg effect in higher grade, and we further show grade-dependent alterations in fatty acid, glutamine, and glutathione metabolism. In addition, we show that the tryptophan pathway feeds directly into known immunosuppressive metabolites, confirming a previous hypothesis formulated from our urinary metabolomics analysis (7).

From the data presented here, it appears that combining omics techniques leads to synergism in knowledge and yields concepts of grade-dependent tumor biology that would otherwise not be obvious using a single technique in isolation. In addition, our work addresses a major issue regarding therapy for RCC patients, namely that there is at present no grade or mutation-specific therapy, despite the fact that many approved targeted therapeutics have been recently developed. Thus, our work will lead to both a better understanding of RCC biology and important clinical advances especially related to potential grade-specific therapeutics.

Materials and Methods

Materials

Low-glucose DMEM, glucose-free DMEM, glutamine-free DMEM, and glucose were obtained from Life Technologies. DMSO, 2-deoxy-D-glucose (2-DG), etomoxir, and mouse monoclonal anti-β-actin antibody were obtained from Sigma. Stabilized glutamine was obtained from Gemini Bio-Products. Goat anti-mouse and goat anti-rabbit horseradish peroxidase–conjugated IgG were obtained from Bio-Rad. Human IFNγ was purchased from PeproTech. Methylthiohydantoin-DL-tryptophan (MTH-trp) was purchased from Enzo Life Sciences. Anti-human indoleamine 2,3-dioxygenase (IDO) antibody was obtained from Cell Signaling Technology. After appropriate Institutional Review Board (IRB) approval at UC Davis or Memorial Sloan Kettering Cancer Center (MSKCC), and after pathologic examination of partial or total nephrectomy samples, excess tissue was analyzed by shotgun proteomics and metabolomics, respectively (Supplementary Table S1).

Cell culture

The RCC cell lines Caki-1 (VHL-wt) and 786-O (VHL-mut) were obtained from the ATCC; the primary “normal human kidney” proximal tubular cells (NHK) were obtained from Lonza. All ATCC and Lonza cell lines undergo extensive authentication tests during the accessioning process as described on their website; in addition, all cells were frequently tested for mycoplasma in the author’s laboratory. The cells were maintained in DMEM supplemented with 10% FBS, 100 units/mL streptomycin, and 100 mg/mL penicillin (complete medium). The cells were maintained at 37°C with 5% CO₂.

Primary cell cultures

After appropriate IRB approval at the University of Milano-Bicocca, primary cell cultures were obtained from human normal cortex and RCC tissues, culture conditions and immunophenotypical characterization were performed as described (9).

Tryptophan and kynurenine analysis

After the indicated treatments, cell media were collected and polar metabolites were extracted using a modification of a previous method (10). Briefly, media aliquots were extracted with ice-cold chloroform, methanol, and 3 mmol/L PIPES-3, mmol/L EDTA, pH 7.4, and vortexed. Extraction mixtures were then stored at −80°C. Immediately before analysis, samples were thawed and centrifuged for 10 minutes at 10,000 rpm. The upper, aqueous phase was subjected to high-performance liquid chromatography (HPLC) analysis using a modification of a previous method (11). HPLC separations were performed on an Agilent 1100 instrument (Agilent Technologies) using a Thermo 50 × 2.1 Hypersil GOLD C18 column. HPLC conditions were as follows: mobile phase: 15 mmol/L potassium phosphate, pH 6.4, with 2.7% (v/v) acetonitrile; Isocratic flow rate: 0.8 mL/min; injection volume: 20 μL; run time: 10 minutes. Column temperature was maintained at 30°C. Detection was accomplished using an Agilent 1100 diode array detector, with detection of kynurenine at 360 nm and detection of tryptophan at 286 nm. Analyte concentrations were calculated using peak area under the curve and standard curves generated using authentic standards for each metabolite of interest.

MTT assay

Cell viability assays were performed as described previously (12). Briefly, cells were plated in 96-well plates, and after appropriate treatments, the cells were incubated in MTT solution/media mixture. Then, the MTT solution was removed, and the blue crystalline precipitate in each well was dissolved in DMSO. Visible absorbance of each well at 540 nm was quantified using a microplate reader.

Lactate assay

Cells were plated in 24-well plates, and after appropriate treatments, the media were processed to measure lactate levels using the Lactate Assay Kit from Biovision following the manufacturer’s instructions.

Oxidative metabolism measurements

Oxygen consumption rates (OCR) were measured using the XF24 Extracellular Flux Analyzer from Seahorse Bioscience. For
experiments utilizing etomoxir or 2-deoxyglucose, cells were pretreated 30 minutes before running the plate on the XF24 instrument. Media were changed to glucose-free or glutamine-free media 30 minutes before running the assay.

**Oil red O staining of RCC tissue**

Cryostat sections (8–10 μm) of RCC tissue samples of distinct Fuhrman grade were fixed in 10% neutral buffered formalin solution (Sigma-Aldrich) for 1.5 hours and rinsed with PBS. Staining was performed with freshly prepared Oil Red O (ORO) working solution, obtained by 3:2 dilution in H2O of ORO stock solution (0.35% in isopropanol), for 1.5 hours. The slides were then rinsed with H2O, and counter staining was performed with haematoxylin solution for 3 minutes. The stained slides were mounted with CC/Mount (Sigma) and analyzed by Nikon Eclipse E800 with 20× objectives. Three to four pictures for each slide were randomly captured and analyzed by LuciaG Image Analysis System (Nikon).

**Oil red O staining and quantitative assay in vitro**

Primary cell cultures and cell lines, plated in triplicate in 96-well plates, were treated with different etomoxir concentrations for 72 hours, well plates, were treated with different etomoxir concentrations quantified by extracting the dye from cells with 200 μL isopropanol and measuring the absorbance of each well at 510 nm using a microplate reader.

**Immunoblotting**

Immunoblotting was done as described previously (12). Briefly, after appropriate treatments, the cells were washed with PBS, lysed in lysis buffer, and cell lysates were immunoblotted. Membranes were blocked in 5% nonfat dry milk for 1 hour at room temperature and probed with appropriate antibodies. Membranes were then probed with horseradish peroxidase–tagged anti-mouse or anti-rabbit IgG antibodies. Signal was detected using enhanced chemiluminescence solutions.

**Oxidized glutathione and glutathione measurement and quantitation**

Cells were plated into 96-well plates (5,000/well) and the next day treated with or without stabilized glutamine (Gemini Bio-Products) for 24 hours. The glutathione (GSH) and oxidized glutathione (GSSG) concentrations were measured in each well of cells using the GSH/GSSG-Glo Assay Kit (Promega) following the manufacturer’s instructions.

Whole frozen tissues that were leftover from the metabolomics analysis were utilized to measure absolute GSH and GSSG levels using the GSH/GSSG-Glo Assay Kit (Promega) following the manufacturer’s instructions.

**Acylcarnitine quantitation**

Deuterium-labeled internal standards were added to the tissues, and the mixture was solubilized in methanol followed by a crash extraction. The extracted mixture was injected onto an Atlantis HIILIC Column connected to a Waters Xevo triple quadrupole mass spectrometer (Waters). The analytes were ionized via positive electrospray, and the mass spectrometer is operated in the tandem MS mode. The absolute concentration of each acylcarnitine was determined by comparing the peak with that of the relevant internal standard. The composition of 15 markers is included in this panel.

**NADP/NADPH assay**

After indicated treatments, the NADPH/NADP ratio was measured using the NADP/NADPH-Glo Assay Kit (Promega) following the manufacturer’s instructions.

**Nontargeted metabolomics analysis**

For metabolomics analysis, after appropriate IRR approval at MSKCC, clear cell RCC (ccRCC) tumors and normal kidney tissues from matched controls were obtained (Supplementary Table S1). Tumors used for the study were from archived frozen ccRCC specimens. The majority of the samples were snap frozen within 1 to 2 hours of resection (maximum up to 4 hours). Samples were inventoried and immediately stored at −80°C. At the time of analysis, samples were extracted and prepared for analysis using Metabolon’s standard solvent extraction method. The extracted samples were split into equal parts for analysis on the GC/MS and LC/MS/MS platforms. Samples were analyzed over 9 consecutive run days. Also included were several technical replicate samples created from a homogeneous pool containing a small amount of all study samples (“Client Matrix”).

Metabolite extraction and measurement were conducted at Metabolon Inc. as previously described (6, 13). All samples were maintained at −80°C until processed, and raw data are shown in Supplementary Table S2 with statistics in Supplementary Table S3. The mass spectrometer platforms, sample extraction and preparation, instrument settings and conditions, and data handling have previously been described in detail (14).

The major components of the process can be summarized as follows: the sample preparation process was carried out using the automated MicroLab STAR system (Hamilton). Samples were prepared using a proprietary series of organic and aqueous extractions to remove the protein fraction while allowing maximum recovery of small molecules. A cocktail of recovery standards was added before the first step in the extraction process for QC purposes. The resulting extract was divided into three fractions for untargeted metabolic profiling and randomized for analysis. Each sample was dried under vacuum to remove organic solvent. Samples were characterized using three independent platforms: ultra high-performance liquid chromatography/tandem mass spectrometry (UHPLC-MS/MS) in the negative ion mode, UHPLC-MS/MS in the positive ion mode, and gas chromatography-mass spectrometry (GC-MS) after silylation. The reproducibility of the extraction protocol was assessed by the recovery of the xenobiotic compounds spiked in every tissue sample before extraction. The LC/MS portion of the platform was based on a Waters ACQUITY UPLC and a Thermo-Finnigan LTQ mass spectrometer, which consisted of an electrospray ionization source and linear ion-trap mass analyzer. One aliquot was analyzed using acidic positive ion optimized conditions and the other using basic negative ion optimized conditions in two independent injections using separate dedicated columns. Samples destined for GC/MS analysis were redried under vacuum desiccation for a minimum of 24 hours before being derivatized under dried nitrogen using bistrimethyl-silyl-trifluoroacetamide. Samples were analyzed on a Thermo-Finnigan Trace DSQ fast-scanning
Identification of known chemical entities was based on comparison to metabolomic library entries of purified standards based on chromatographic properties and mass spectra. As of this writing, more than 4,000 commercially available purified standard compounds had been acquired and registered into the LIMS for distribution to both the LC and GC platforms for determination of their analytical characteristics. MS peaks were identified using Metabolon’s proprietary peak integration/identification software, by comparing MS peak data with that of a library of purified standard. The combination of chromatographic properties and mass spectra gave an indication of a match to the specific compound or an isobaric entity. The integrated ion peaks for each compound were normalized in MS run-day blocks by registering the medians to equal one (1.00) and normalizing each data point proportionately (termed the “block correction”). This integrated peak normalization generates a median scaled relative value for each metabolite measured in a given sample. Minimum median scaled metabolite values were imputed for specific samples in which the metabolite levels fell below the lower limit of quantitation. Following normalization procedures and imputation, statistical analysis was performed on log-transformed data to identify significant differences in metabolite levels between comparator groups.

Proteomics analysis
Tissue protein levels were obtained from a previously published proteomics study in RCC tissues that was performed under appropriate UC Davis IRB approval as described (4). The raw data and statistics for protein levels in tumors compared with normal kidney tissues are in Supplementary Table S4.

Creation of pathway maps
Data from a prior nontargeted proteomics study on grade-dependent formalin-fixed paraffin-embedded ccRCC tissues (4) and metabolomics data described above (Supplementary Table S1) were used to compare proteins/enzymes with metabolic pathways. Quantitative changes of individual metabolites as compared with matched control tissues, and proteins in relation to adjacent “normal” tissue, were calculated and graphed on a Kyoto Encyclopedia of Genes and Genomes (KEGG)-based pathway map as a function of grade. Although it is true that levels of many enzymes are not altered when their catalytic activity is increased, by combining proteomics and metabolomics data, we were able to identify activity of enzymes that had no change in protein concentration because enzymatic activity is quantitatively proportional to the appearance of the end metabolite.

Results and Discussion
The purpose of this study was to evaluate specific biochemical pathways associated with RCC, and moreover to do so at a high degree of resolution, in a continuing effort to attain new insights regarding metabolic reprogramming in these tumors, as well as to identify potentially new grade-dependent therapeutic targets for RCC. This work is unique from other metabolic studies in RCC due to its concurrent utilization of two distinct although complementary techniques to generate and validate reprogrammed energy metabolism and tryptophan pathways in RCC, thereby ameliorating some of the drawbacks and deficiencies of each omics technique as used in isolation.

Aerobic glycolysis is upregulated in high-grade RCC
It has long been known that most solid tumors utilize aerobic glycolysis/lactic acid fermentation (the Warburg effect) in order to provide sufficient energy for cell metabolism, as well as to synthesize key membrane components and other cellular building materials (4, 15–17). Given the prominence of the Warburg effect observed in RCC in many studies (including our proteomics studies; refs. 4, 5), we first asked whether there is evidence of aerobic glycolysis in RCC tissues using the combined metabolomics/proteomics approach. The glycolysis-relevant metabolites (glucose, glucose-6-phosphate, and fructose-6-phosphate) are increased significantly with higher grade, suggesting that glucose metabolism is more prominent with increasing tumor grade (Fig. 1; Supplementary Tables S3 and S4). Consistent with these data, most of the enzymes involved in this process, especially glyceraldehyde 3-phosphate dehydrogenase (GAPDH), enolase 2, and pyruvate kinase-muscle-2 (PKM2), are increased in tumors as compared with normal tissues, although their levels do not show the same unambiguous grade dependence as most of the metabolites, likely due to the fact as previously noted that enzymatic activity can change without an alteration of enzyme (i.e., protein) concentration. L-lactate was similarly increased in a grade-dependent manner (up to 2-fold in grade 4), suggesting that lactate fermentation was upregulated in RCC, particularly at higher grades. In support of these data, lactate dehydrogenase A (LDHA), involved in converting pyruvate to lactate, was increased in all grades of RCC as compared with control tissue possibly through increased HIF activity (18) due to VHL inactivation that is seen in more than 90% of RCC tumors (19). On the other hand, pyruvate carboxylase (PC), the enzyme that converts pyruvate to oxaloacetate in the tricarboxylic acid (TCA) cycle, was decreased in all grades, suggesting that glucose and sugars involved in glycolysis were utilized for lactate fermentation in
RCC rather than for metabolism within the TCA cycle or other downstream oxidative metabolism pathways.

Although the omics-derived pathway data are suggestive of biochemical and metabolic changes, we next asked whether such biology, which would occur as a result of such reprogramming, could be validated using available RCC cell lines. Both VHL-mut (786-O) and VHL-wt (Caki-1) human RCC cell lines showed decreased viability in decreased glucose concentrations or when

Figure 1.
Aerobic glycolysis is grade-dependently upregulated in RCC. Combined proteomics and metabolomics data of human RCC tissue were overlaid onto a stylized KEGG-based pathway diagram. Green, metabolite; orange, enzyme; black dotted arrow, metabolism; red arrow, upregulated pathway; blue arrow, downregulated pathway; G6P, glucose-6-phosphate; PEP, F6P, fructose 6-phosphate, phosphoenolpyruvate; OAA, oxaloacetate; GPI, glucose-6-phosphate isomerase; ALDOB, aldolase B; TPI1, triosephosphate isomerase 1; PGK1, phosphoglycerate kinase 1; ENO, enolase; PKM, pyruvate kinase; AKR1A1, aldo-keto reductase family 1, member A1; LDH, lactate dehydrogenase.
incubated with 2-DG (a glycolysis inhibitor) at various concentrations (Fig. 2A). To confirm that, as a signature of aerobic glycolysis, lactate production in these cells was highly active under normoxia, both cell lines were incubated in 5 mmol/L glucose media, glucose-depleted media, or with 2-DG and 5 mmol/L glucose, and lactate concentrations were measured in the media. Both procedures (2-DG and glucose depletion) markedly decreased lactate production (Fig. 2B), demonstrating that both RCC cell lines studied rely primarily on aerobic glycolysis (Warburg effect), consistent with the data obtained from the human RCC tissues.

Acylcarnitines are increased, yet fatty acid β-oxidation mechanism is attenuated, with higher grade

Many tumors and other tissues are able to oxidize fatty acids to satisfy their energy requirements (20). Free fatty acids (FFA) are esterified to fatty acyl-CoA and then transported into the mitochondria by carnitine palmitoyltransferase-1 (CPT1) and the carnitine system, at which point in normal tissue they are subjected to β-oxidation as fatty acyl-CoA to feed into the TCA cycle (21). To study this process in human RCC as a function of tumor grade, we evaluated the levels of the key components of this pathway as well as the enzymes that control their interconversions. Although the levels of shorter chain FFAs (6:0, 8:0, 9:0, 10:0, and 12:0) were generally decreased in a grade-dependent manner, the levels of carnitine and the acylcarnitines were increased in all grades (Fig. 3; Supplementary Tables S3 and S5) consistent with previous human urinary metabolomics studies in our laboratory (22). Absolute quantitation of acylcarnitines in a selection of tumor and matched normal control tissues revealed that most of these metabolites were higher in RCC tissues, and the increase was significant for the major metabolite L-carnitine (Supplementary Table S5). However, surprisingly, most of the enzymes of β-oxidation were decreased in high-grade tissues, suggesting a lack of oxidation of the acyl-CoAs in RCC tissue, which is a possible mechanism for the accumulation of acylcarnitines.

Adipogenesis with clear cytosol is a known signature of RCC (23), and increased activity of adipogenesis-related proteins, such as acyl-CoA:cholesterol acyltransferase, has been reported to be an indicator of aggressiveness (24). Thus, we first hypothesized that the grade-dependent decrease of FFAs is due to increased adipogenesis: accumulation of lipids (triglycerides and esterified fatty

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Acylcarnitines are increased and β-oxidation is downregulated in RCC. Combined proteomics and metabolomics data of human RCC tissue were overlaid onto a stylized KEGG-based pathway diagram. Green, metabolite; orange, enzyme; black dotted arrow, metabolism; red, upregulated pathway; blue arrow, inhibition. Acylcarnitines are increased and β-oxidation enzymes were downregulated in RCC. Combined proteomics and metabolomics data of human RCC tissue were overlaid onto a stylized KEGG-based pathway diagram. Green, metabolite; orange, enzyme; black dotted arrow, metabolism; red, upregulated pathway; blue arrow, inhibition.

The glutamine pathway feeds the GSH/GSSG antioxidant system

There is increasing evidence that the glutamine pathway is essential in a variety of cancers, including RCC, to support efficient carbon metabolism of both anabolism and cell growth (25, 26). Indeed, VHL-mut RCC has been shown to be sensitive to glutaminase inhibition, as in some cases these tumors use glutamate to regulate the TCA cycle and subsequent lipogenesis (27). Moreover, the prominence of glutaminase may be a vestige of nontransformed renal epithelial cell biology in which it is utilized for systemic acid secretion through the renal tubules (28).

Glutamine feeds into various downstream metabolic pathways, including the TCA and urea cycles as well as the GSH pathway. Although those enzymes detected by the metabolomics analysis which catabolize glutamine or glutamate to feed into the TCA and urea cycles were decreased in our analysis, the metabolite levels in the pathway from glutamine to GSH and GSSG were grade-dependently increased and associated with generally decreased levels of the enzymes that catalyze GSH to glutamate, suggesting that glutamine is reprogrammed to the GSH pathway at higher grades (Fig. 4; Supplementary Fig. S2A). Indeed, the marked increase in GSH in the global metabolomics study (Fig. 4; Supplementary Table S2) was one of the highest magnitude changes observed in our entire metabolomics analysis. These data suggest that glutamine is being utilized by the cancer cell in large part to attenuate oxidative stress, rather than for energy and cell component production through the TCA cycle, thereby yielding a survival advantage to these cells. In fact, this hypothesis is validated by the finding that the GSH/GSSG ratio in grade 3 and 4 tumors, after direct quantitative measurement of these metabolites, was significantly higher than that of normal kidneys (Supplementary Fig. S2A and Supplementary Table S5), indicative of decreased oxidative stress in the high-grade tumors.

To validate this pathway, we examined the GSH pathway as a function of glutamine in vitro; cell viability assay results showed that RCC cell viability was decreased in both cell lines dose-dependently with glutamine depletion (Supplementary Fig. S2B). To further analyze the function of glutamine as a source for the GSH pathway, absolute quantitative GSH and GSSG levels and GSH/GSSG ratios were measured in cells grown in with and without glutamine. Levels of both GSH and GSSG and GSH/GSSG ratios decreased significantly in both cell lines in glutamine-depleted media (Supplementary Fig. S2C), consistent with our postulated role of glutamine in the tumoral antioxidant response system. To confirm that the effect of glutamine depletion correlated with a change in redox state, we evaluated the NADPH/NADP⁺ ratio in the glutamine-depleted media and found that the ratio was significantly decreased in the glutamine-depleted state in both cell lines (Supplementary Fig. S2D), further supporting an increased signature of oxidative stress in the absence of glutamine.
Taken together, these findings confirm the necessity of glutamine for attenuating oxidative stress by kidney tumors.

The TCA cycle is downregulated as a function of tumor grade

PC, one of the main routes by which pyruvate enters the TCA cycle, was decreased in RCC (Figs. 1 and 5A; Supplementary Table S4), and several enzymes regulating the “input” pathways to the TCA cycle, such as enzymes involved in β oxidation and gamma-aminobutyric acid (GABA) transaminase, were downregulated as a function of grade (Figs. 3, 4, and 5A). Furthermore, the increased acetylcarnitine level suggested that acetyl-CoA was not being used to feed the TCA cycle (Fig. 3). Taken together, these findings suggest that three major inputs to the TCA cycle (glycolysis, fatty acid metabolism, and the glutamine pathway) are reprogrammed in RCC. To confirm this finding, we measured OCR, as a proxy for oxidative metabolism, in RCC cell lines treated with glucose deprivation, etomoxir (inhibitor of CPT1), or glutamine depletion. OCR was not affected by inhibition of glycolysis, fatty acid metabolism, or glutamine metabolism in RCC cells (786-O and Caki-1), while NHK cells had a significantly decreased OCR with etomoxir treatment (Fig. 5B), indicating that these major energy pathways, all converging on the TCA cycle, were not significantly contributing to oxidative metabolism in two RCC cell lines while normal cells did rely more heavily on this pathway. These findings, in light of the above data indicating an increased transit through the aerobic glycolysis pathway, suggest that RCC cells do...
The tryptophan pathway results in IDO-dependent production of anti-inflammatory metabolites

Tryptophan is an essential amino acid that undergoes multiple pathways of metabolism in normal and cancerous tissue, resulting in immunosuppressive and serotoninergic properties (29). Based on previously published work from our laboratory demonstrating high levels of immunosuppressive tryptophan metabolites in human RCC urine (7) and in human RCC xenograft mice urine and serum (6), we next asked whether, and in what manner, the tryptophan metabolic pathway is upregulated as a function of RCC grade. When tryptophan pathways were examined by combined metabolic and proteomic analyses, the immunosuppressive metabolite kynurenine increased in tumors compared with normal tissues, the immunosuppressive metabolite quinolinate increased in grade 2 tumors compared with normal tissues, and the activity of enzymes that catalyze tryptophan to the other known arms of tryptophan metabolism (including serotonin and indoleacetic) was decreased inversely proportional to tumor grade (Fig. 6A), consistent with our previously published data.

The cytokines IFNγ, IFNα, and IFNβ stimulate the transcription of IDO, the primary enzyme in the tryptophan pathways that produce immunomodulatory metabolites, and this finding may explain the relative lack of success when interferons are used clinically to treat advanced RCC. To validate the tryptophan pathway data, we showed that IFNγ increases IDO levels (Fig. 6B), and inhibition of IDO by MTH-trp results in increased tryptophan and decreased kynurenine levels in the culture media (Fig. 6C). The fact that an IDO inhibitor, 1-D-MT, is available and currently in clinical trials for other cancers (Clinicaltrials.gov NCT01021608) suggests that targeting IDO may provide a new therapeutic avenue for advanced RCC.
Indole-3-acetaldehyde

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viability assay kit (EMD Millipore) on a MUSE (EMD Millipore). Data shown are mean (conditioned media were harvested and tryptophan (TRP) and kynurenine (KN) were measured by HPLC and normalized for cell number counted using the cell

Strengths and weaknesses of current study

A strength of our study is that it demonstrates that each Fuhrman grade of tissue possesses distinct biochemical characteristics with a specific magnitude of metabolic reprogramming, a phenomenon that is logical given the markedly disparate clinical behavior of tumors of different grades. In addition, despite tumor heterogeneity observed in RCC (30), it would appear that in this era of personalized medicine, and based on the differences that we have observed, it may now be more appropriate to evaluate patient responses to drug therapies with regard to the differences that we have observed, it may now be more appropriate to evaluate patient responses to drug therapies with the use of experienced pathologists for determination of such, we have attempted to minimize potential inconsistencies from this issue. While there exists variability in any omics experiment, this possibility was minimized in this work by utilizing laboratories with abundant experience in nontargeted metabolomics and proteomics, both of which were based on mass spectrometry. To further minimize potential errors from this issue, validation of findings from each pathway was performed in vitro.

Although some of our findings, such as enhancement of the Warburg effect as a function of increasing tumor grade, were not surprising given the known activation of hypoxia pathways in the majority of RCCs, other pathway alterations were quite unexpected and represent novel potential therapeutic targets for further research (Fig. 7). The overarching theme of our findings regarding energy metabolism is that (i) the Warburg effect seemed to be of prime importance for these tissues, especially of high grades, at the expense of oxidative metabolism, including fatty acid oxidation and transit through the TCA cycle; (ii) the glutamine pathway functions to attenuate the high levels of reactive oxygen species that have long been known to be present in RCC (31); and (iii) tryptophan metabolism was upregulated, leading to the

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Figure 6.

Tryptophan metabolism favors a grade-dependent increase in immune-suppressive metabolites. A, combined proteomics and metabolomics data of human RCC tissue were overlaid onto a stylized KEGG-based pathway diagram. Green, metabolite; orange, enzyme; black dotted arrow, metabolism; red arrow, upregulated pathway; blue arrow, downregulated pathway; MAO, monoamine oxidase; DDC, dopa decarboxylase; ALDH, aldehyde dehydrogenase; IDO, indoleamine 2,3-dioxygenase; TDO, tryptophan 2,3-dioxygenase. B, 786-O (VHL-mut) and Caki-1 (VHL-wt) cells were plated in 6-well plates the day before treating with either human IFN-g (50 ng/mL) and/or MTH-trp (100 μmol/L) for 4 days and immunoblotted with the antibodies indicated. C, 786-O (VHL-mut) and Caki-1 (VHL-wt) cells were plated in 6-well plates the day before treating with either human IFN-g (50 ng/mL) and/or MTH-trp (100 μmol/L) for 3 days. The conditioned media were harvested and tryptophan (TRP) and kynurenine (KN) were measured by HPLC and normalized for cell number counted using the cell viability assay kit (EMD Millipore) on a MUSE (EMD Millipore). Data shown are mean (n = 3) and SD. *P < 0.05. μM/1M cell, μM/1 million cells.

NCT00567931) presents an incentive to evaluate IDO as a novel RCC therapeutic target, which work is under way in our laboratory.

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Figure 7. Summary of grade-dependent metabolic pathway alterations in RCC. With higher grade, glycolysis was directed toward lactate metabolism at the expense of TCA cycle intermediaries. Fatty acid β-oxidation was decreased, whereas the glutamine pathway served to attenuate oxidative stress, thereby increasing cancer cell survival. Tryptophan was metabolized preferentially to immunosuppressive compounds. Red, increased; blue, decreased.

References


Disclosure of Potential Conflicts of Interest

B. Neri is Sr. Director Diagnostic R&D at Metabolon Inc. No potential conflicts of interest were disclosed by the other authors.

Authors’ Contributions

Conception and design: H.I. Wettersten, O. Abu Aboud, B. Neri, R. Wollert, J.J. Hsieh, R.H. Weiss
Development of methodology: H.I. Wettersten, D. Morin, M.E. Johnstone, O. Abu Aboud, R. Wollert, J.J. Hsieh
Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): H.I. Wettersten, A.A. Hakimi, C. Bianchi, M.E. Johnstone, D.R. Donohoe, J.F. Trott, R. Wollert, B. Stewart, R. Perego, J.J. Hsieh
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): H.I. Wettersten, D. Morin, M.E. Johnstone, D.R. Donohoe, O. Abu Aboud, S. Stiindivant, B. Neri, R. Wollert, J.J. Hsieh, R.H. Weiss
Writing, review, and/or revision of the manuscript: H.I. Wettersten, A.A. Hakimi, C. Bianchi, J.F. Trott, S. Stiindivant, B. Neri, B. Stewart, R. Perego, J.J. Hsieh, R.H. Weiss
Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): H.I. Wettersten, D. Morin
Study supervision: R.H. Weiss

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Hiromi I. Wettersten, A. Ari Hakimi, Dexter Morin, et al.

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