Kidney Tumor Biomarkers Revealed by Simultaneous Multiple Matrix Metabolomics Analysis

Sheila Ganti1, Sandra L. Taylor2, Omran Abu Aboud1, Joy Yang3, Christopher Evans3,4, Michael V. Osier6, Danny C. Alexander7, Kyoungmi Kim2, and Robert H. Weiss1,4,5

Abstract

Metabolomics is increasingly being used in cancer biology for biomarker discovery and identification of potential novel therapeutic targets. However, a systematic metabolomics study of multiple biofluids to determine their interrelationships and to describe their use as tumor proxies is lacking. Using a mouse xenograft model of kidney cancer, characterized by subcapsular implantation of Caki-1 clear cell human kidney cancer cells, we examined tissue, serum, and urine all obtained simultaneously at baseline (urine) and at, or close to, animal sacrifice (urine, tissue, and plasma). Uniform metabolomics analysis of all three “matrices” was accomplished using gas chromatography– and liquid chromatography–mass spectrometry. Of all the metabolites identified (267 in tissue, 246 in serum, and 267 in urine), 89 were detected in all 3 matrices, and the majority was altered in the same direction. Heat maps of individual metabolites showed that alterations in serum were more closely related to tissue than was urine. Two metabolites, cinnamoylglycine and nicotinamide, were concordantly and significantly (when corrected for multiple testing) altered in tissue and serum, and cysteine–glutathione disulfide showed the highest change (232.4-fold in tissue) of any metabolite. On the basis of these and other considerations, three pathways were chosen for biologic validation of the metabolomic data, resulting in potential therapeutic target identification. These data show that serum metabolomics analysis is a more accurate proxy for tissue changes than urine and that tryptophan degradation (yielding anti-inflammatory metabolites) is highly represented in renal cell carcinoma, and support the concept that PPAR-α antagonism may be a potential therapeutic approach for this disease. Cancer Res; 72(14); 1–9. ©2012 AACR.

Introduction

The use of metabolomics to identify tumor biomarkers as well as potential targets for therapy has entered mainstream clinical medicine and is beginning to lead to payoffs in kidney (1, 2), prostate (3), and pancreatic (4) cancer, as well as in kidney disease in general (5). Although most of the existing published studies in the field have focused on either tumor tissue or a specific biofluid for metabolomics analysis, there are limited available data examining the use of a biofluid to serve as a proxy for tumor metabolomic changes through simultaneous examination of tissue in addition to several biofluids.

Because the ultimate purpose of a metabolite biomarker is to reflect biochemistry of the tumor of interest, it is essential to determine how metabolomic profile changes occurring in a tumor are reflected in blood and urine.

To begin to address this question, as well as to extend our ongoing work in kidney cancer metabolomics and biomarker discovery, we used a xenograft mouse model of highly metastatic renal cell carcinoma (RCC) represented by subcapsular implantation of human Caki-1 cells. This disease model has the advantage of closely recapitulating human RCC in that the cancer cells are implanted under the renal capsule, using sham surgery animals to control for any metabolic changes that might occur after surgical stress. From this model, we obtained terminal tissue, serum, and urine simultaneously, and we carried out global metabolomics analysis on each matrix using identical analytical platforms and run at the same time.

We now show that most of the identified metabolites that were detected in all 3 matrices were concordant in their direction of change and that blood serves as a more accurate proxy of tissue change than does urine, as the magnitude of metabolite changes in each matrix show a gradation from urine (most) to serum to tissue (least). Consistent with other published studies in RCC (1, 6, 7) and to further confirm the veracity and consistency of our data, we biologically validated several relevant pathways whose signatures were significantly altered in one or several matrices, specifically the tryptophan metabolism pathway, which is linked to immunosuppressive
metabolites. In addition, from the finding that the 2 metabolites, which were altered in all 3 matrices, are present in the peroxisome proliferator pathway, the target receptor PPAR-α was identified and validated in vitro. Thus, global metabolomics of a mouse xenograft model indicates that serum, and to a lesser extent urine, show utility as proxies for tumoral metabolism changes. Furthermore, validation of several identified altered pathways suggests that these could be further evaluated as potential markers and therapeutic targets for this disease.

Materials and Methods

Materials

Four human proximal tubule epithelial cancer cell lines, ACHN, A498, 786-O, and Caki-1, and one "normal" derived kidney epithelial cell line, HK-2, were obtained from the American Type Culture Collection (ATCC). Primary proximal renal tubular epithelial cells (NIK) were from Lonza. All ATCC and Lonza cell lines undergo extensive authentication tests during the accessioning process as described in vendor websites. ACHN, A498, and HK-2 cells were maintained in Dulbecco's Modified Eagle's Medium, and Caki-1 and 786-O cells were maintained in RPMI, all supplemented with 10% FBS and 100 units/ml streptomycin and 100 µg/ml penicillin. Cells were maintained at 5% CO2 at 37°C.

Mouse subcapsular xenograft model

Human Caki-1 cells (10⁶) were mixed with 30% of nongrowth factor reduced Matrigel (BD) and injected into the right flanks of 2 donor nude mice. After the tumors reached 500 mm³ of size, mice were sacrificed and tumors excised to prepare tumor cell suspension. Tumors were minced and passed through 70 µm Nylon cell strainer (BD Falcon) and washed with PBS to collect cell pellets. Tumor cells were resuspended in 30% Matrigel (10⁶ cells/20µL/mouse) for renal subcapsular implantation. Nude mice were anesthetized with isoflurane and the kidneys were exteriorized for subcapsular injection. The sham control group of mice received no injection of tumors. After injection, the kidney was returned to the abdominal cavity and the peritoneum and skin were closed by suture and metal wound clips, respectively. All of the mice were sacrificed when the xenografted animals became moribund, 34 days after surgery. Kidney plus tumor size averaged 1.62 grams at sacrifice. Terminal serum was collected, and tumor (from xenografted animals) and normal kidneys (from sham surgery animals) were removed and split for snap freezing and 10% buffered formalin fixation at sacrifice. For urine, we collected at 2 time points, one day before surgery and 32 days after surgery (2 days before sacrifice). From the frozen tissue, tumors were dissected out from adjacent noncancerous tissue, and this as well as sham control kidneys were subjected to nontargeted metabolomic analysis.

MTT assay

A total of 5 × 10⁴ cells were plated in 96-well plates and incubated for 16 hours at 5% CO₂ at 37°C. After appropriate treatments, the cells were incubated in 20 µL of thiazolyl blue tetrazolium bromide (MTT) solution (5 mg/mL in PBS) with 180 µL of the growth medium for 3 hours. Then, the MTT solution was removed and the blue crystalline precipitate in each well was dissolved in DMSO (dimethyl sulfoxide; 200 µL). Visible absorbance of each well at 540 nm was quantified using a microplate reader.

Immunoblotting

Cells were treated with either vehicle, LPS, or 1-MT for 72 hours. The cells were then lysed, protein was collected, and immunoblotting was carried out as previously described (8).

Nontargeted metabolomic analysis

The metabolomic platforms, including sample extraction process, instrumentation configurations and conditions, and software approaches for data handling, were previously described in detail (1, 9). Urinary metabolite values were creatinine normalized to account for urine concentration differences among samples, and tissue samples were normalized to equal mass before chromatographic analysis. Due to space restrictions, these techniques are further described in Supplementary Data.

Statistical methods

Processing of the raw data yielded 299 known metabolites from tissue, 251 from serum and 274 from urine samples from xenograft and sham surgery mice. For each matrix, metabolites observed in fewer than 3 of the samples from each of the experimental groups (xenograft and sham surgery) were excluded. This screening resulted in 267, 246, and 267 metabolites for tissue, serum, and urine, respectively, that were subsequently statistically analyzed. Missing metabolite values were imputed using the minimum observed metabolite value. Before statistical analyses, metabolite intensities were log transformed to meet underlying assumptions of normality with a constant variance and to reduce the dominant effect of extreme values. Before log transformation, urinary metabolite values were creatinine normalized to account for urine concentration differences among samples.

The primary objective of the statistical analysis was to identify metabolites in each matrix whose concentration differentiates xenograft and sham surgery mice that potentially could serve as diagnostic biomarkers for kidney cancer as well as to elucidate alterations of metabolite signals in pathways associated with the presence of kidney cancer. To identify metabolites as potential diagnostic biomarkers for mice with kidney cancer, we aimed to (i) identify metabolites that distinguish xenograft and sham surgery mice using differential analysis, and (ii) identify sets of relevant metabolites that act synergistically within functionally defined pathways using functional score analysis.

For each tissue and serum metabolite, we identified differentially expressed metabolites between xenograft mice and sham surgery mice using t tests. For urine, because we
measured twice 1 day before surgery and 32 days after surgery from the same animals, we used an analysis of covariance to identify differentially regulated metabolites, modeling postsurgery metabolite values as a function of cancer status, with presurgery metabolite values included as a covariate. Significance was determined based on a permutation null distribution consisting of 6,433 permutations for tissue and serum and 3,432 permutations for urine. The number of permutations was lower for urine because we had one less sample (i.e., urine samples from 8 xenograft mice and 7 sham surgery mice) available for the statistical analysis. False discovery rates (FDR) were also calculated to account for multiple testing. We also conducted a partial least squares (PLS) regression analysis for each matrix separately using all metabolites. For urine, we used the postsurgery values only for consistency with the other matrices. Leave-one-out cross-validation was conducted using up to 10 latent components. \( R^2 \) was calculated as a measure of the amount of variability explained by the PLS regression, whereas the \( Q^2 \) values were a measure of the predictive error of the regression. For each metabolite, the variable importance in the projection (VIP) score was calculated to determine a metabolite’s influence on predicting the outcome as well as its weight in the predictor matrix, while considering the presence of multicollinearity among metabolites.

Results

Metabolomic analysis of all 3 matrices shows similarities among matrices

Eight nude mice were implanted with Caki-1 cells under the renal capsule, and 7 control mice of identical genetic background were subjected to sham surgery. Urines were collected one day before surgery and 32 days after surgery (i.e., 2 days before sacrifice). Blood (as serum) and kidney tissue were obtained simultaneously from all animals at sacrifice (34 days after surgery); the tumor + kidney weights at sacrifice averaged are given in Supplementary Table S1. All samples were subjected to simultaneous global metabolomics analysis by liquid chromatography (LC)- and gas chromatography–mass spectrometry (GC–MS) as described in Materials and Methods and Supplementary Data. The 267 metabolites identified from tissue, 246 from serum, and 267 from urine were used for further downstream analyses. As expected, many of the metabolites identified in one matrix were also found in the other(s). Of the suite of identified metabolites, 89 were detected in all 3 matrices (Fig. 1). Tissue and serum had the most identified metabolites in common, with 174 metabolites identified in both of these matrices. Interestingly, urine had fewer metabolites in common with the other matrices and had the largest number (122) of unique metabolites. Thus, there exists a closer link between tissue and serum metabolic profiles than between tissue and urine.

One goal of a biofluid study for biomarker discovery, especially in kidney cancer, is to determine whether easily accessible fluids (blood and urine in this case) can serve as proxies for what is occurring in the tumor tissue. About two-thirds of the 89 metabolites found in all 3 matrices were altered in the same direction (Table 1) supporting the hypothesis that metabolic changes in tissue are reflected in serum and urine. Notably, about one-third of the metabolites were altered in the opposite direction in biofluids as compared with tissue. Thus, although consistent changes in some metabolites can potentially be traced through all 3 matrices, other metabolites show differential regulation patterns, which may be consistent with the existence of nodes or convergence points in metabolic pathways.

Although most metabolites were altered in the same direction across the 3 matrices, metabolic differences between xenograft and sham surgery mice were most pronounced in tissue followed by serum and urine (Fig. 2). When metabolite changes in cancer versus control animals were compared using heat map visualization, it became obvious that differences in tissue metabolites between cancer and control are more pronounced than these differences in serum or in urine (Fig. 3A–C). This observation was also supported by \( R^2 \) and \( Q^2 \) values adapted to evaluate the prediction performance of PLS (Supplementary Table S2). \( R^2 \) values were high for all the 3 matrices when we used the first 2 or more components, suggesting the model fitness is good. \( Q^2 \) values for serum were considerably smaller than for tissue but higher than urine. The finding that serum changes between cancer and control animals are more marked than urine is expected, because blood is in intimate contact with the tumor because of its abundant angiogenesis (especially in RCC), and urine is subjected to additional processing of the plasma by the kidneys.

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<th>Table 1. Number of metabolites altered in the same direction (concordant) and opposite direction (discordant) between tissue, serum, and urine</th>
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that are regulated by modulators of PPAR-α (nicotinamide and cinnamoylglycine; Fig. 4B). These findings suggested modulation of specific pathways by the presence of the tumor and thus identification of potential therapeutic targets (see below).

Validation of tissue pathways supports veracity of the metabolomics data

In any high-throughput metabolomics study, key metabolic pathways, as suggested by statistical significance, need to be validated in benchtop experiments to ensure the veracity of the metabolomics data and to identify biologic correlates. Rationales for selection of metabolites in the discovery stage for further validation can include statistical significance and/or biologic evidence, such as the relative changes of metabolite expression in cancer compared with control or functional relevance of the metabolites as evidenced by their involvement in the same pathway. For such validation, we have chosen 2 separate pathways or metabolites based on the findings as discussed above: (i) the PPAR-α pathway that is associated with the metabolites that were significantly altered in tissue and serum at FDR < 0.05 and in urine using less stringent FDR < 0.1 and raw P value less than 0.01 (cinnamoylglycine and nicotinamide); and (ii) the tryptophan degradation pathway in which we have seen significant changes in metabolites in human urine (2) as well as in the mouse tissue described here.

PPAR-α modulation results in alteration of cell growth

As mentioned above, nicotinamide and cinnamoylglycine were 2 of the 5 significantly and concordantly changed (attenuated) in association with the cancer state in all 3 matrices (by FDR < 0.05 in tissue and blood and by FDR < 0.1 in urine; Fig. 4), and these compounds are altered by peroxisome proliferators. The PPAR-α pathway is known to regulate key enzymes of the tryptophan pathway (10) that leads, through quinolinate (found elevated in a previous study; ref. 1), to production of NAD⁺. NAD⁺ is a strong reducing agent and is involved to a large degree with cellular energetics and other electron donor processes. Animals fed the PPAR-α agonist Wy-14,643 showed altered urinary levels of nicotinamide and cinnamoylglycine (11). Relevant to RCC, other investigators used siRNA to attenuate PPAR-α in Caki-1 cells to show that this receptor regulates genes involved in fatty acid metabolism (12). These findings suggested that PPAR-α may be involved in tumor promotion or attenuation and prompted us to explore this nuclear receptor as a possible target in RCC. When incubated with several RCC cell lines, as well as a “normal” renal tubular epithelial cell line (HK-2) and primary RTE cells (NIH), the PPAR-α agonist Wy-14,643 showed increased cell proliferation whereas the PPAR-α antagonist GW/6471 had the opposite effect and was considerably more pronounced (Fig. 5). To recapitulate these data in cells, we used the Caki-1 cell line and showed the expected effects of the PPAR-α agonist Wy-14,643 upon tryptophan and nicotinamide levels (refs. 11, 13; Supplementary Fig. S1). Thus, PPAR-α may be a viable target for RCC therapy with PPAR-α antagonism resulting in significant inhibition of cell proliferation.

**Intermatrix comparison reveals common biochemical function among matrices**

Metabolites with large changes in tissue and showing concordant and detectable changes in serum and/or urine are the most logical candidates for potential biomarkers. Consequently, to evaluate specific metabolites as potential biomarkers, we next identified metabolites that significantly differed between xenograft and sham surgery control mice but with particular attention to shared metabolites altered in the same direction (i.e., concordant) among all matrices. Tissue had the largest number of differentially regulated metabolites with 186 of the 267 metabolites in this matrix differing significantly at FDR < 0.05 between xenograft and sham surgery mice (Supplementary Table S3A). In serum, 76 of 246 differed significantly at FDR < 0.05 (Supplementary Table S3B) and in urine, no metabolites were statistically significantly different (Supplementary Table S3C). The VIP scores obtained by the PLS regression for selecting most relevant metabolites, which have a significant effect on separation between sham surgery and control samples, were in agreement with the (FDR) P values from the differential analysis in terms of significance (Supplementary Table S4). Interestingly, urine has higher VIP scores compared with tissue and serum, which implies a lower proportion of relevant metabolites or a higher magnitude of correlation among metabolites with similar effects on the separation.

When examined across all 3 matrices, 5 metabolites (cinnamoylglycine, glucose, nicotinamide, phenylpropionoylglycine, and valine) were significantly downregulated in tissue and serum at FDR < 0.05 and close to significance in urine, all with FDR < 0.1 and raw P value less than 0.01 (Fig. 4A). Looking at tissue and plasma together, there were a plethora of metabolites that were significantly and concordantly altered in these matrices, including 2 metabolites

![Figure 2. Variation in metabolites in each matrix. Distribution of fold change (log2) of metabolites between xenograft and sham surgery mice in the 3 matrices. Dotted line shows a log2 fold change of 1 indicating no difference between xenograft and sham surgery.](image-url)
Attenuation of the tryptophan degradation pathway results in higher levels of inflammatory markers and may contribute to immune escape

By metabolomics analysis of tumor tissue, the tryptophan level was significantly lower (0.69-fold, FDR = 0.0024), and the downstream metabolite kynurenine was significantly elevated (2.78-fold, FDR = 0.0008) compared with controls. Our previous study using human urine showed that quinolinate, another downstream metabolite in addition to kynurenine, was elevated in human RCC (1). Taken together, these data indicated that tryptophan metabolism is increased in RCC, resulting in the decrease of tryptophan and the accumulation of downstream metabolites in xenograft tumor tissue and serum and the accumulation of such metabolites in human urine.

In our earlier metabolomics and proteomics studies, we have frequently observed altered tryptophan metabolism (1, 7). It has been shown that increased tryptophan metabolism is associated with both decreased proliferation of T cells and a reduced immune response, mediated by the enzyme indoleamine-2,3-dioxygenase (IDO; refs. 14, 15); patients with ovarian, endometrial, hepatocellular, and colorectal carcinomas have all been shown to have chronically activated IDO (14). Thus, to further validate the metabolomics data (in addition to our previously published work cited above), we reduced the
accumulation of the downstream metabolites by inhibiting IDO and assessed the inflammatory response in the tumor cells. RCC cell lines were treated with the specific IDO inhibitor 1-MT and COX2 induction, as a measure of inflammation (16), was detected by immunoblot (Fig. 6). A dose of 200 μmol/L of 1-MT significantly induced COX2 expression in A498 and ACHN cancer cells, thus decreased tryptophan metabolism mediated by increasing tryptophan metabolism, a means by which RCC tumors prevent an inflammatory response in the tumor itself (as well as in any systemic effects of the malignancy), but before this study, this assumption has not been validated. For these reasons, we have undertaken a comprehensive metabolomics analysis of tissue, blood, and urine taken simultaneously from an anatomically accurate RCC mouse xenograft model. In light of the data obtained from this study, it is evident that blood is a decent proxy of tumoral metabolic changes, whereas urine is less reflective of such changes.

Some of the suite of metabolomic changes seen here can be correlated to changes in metabolic pathways. In particular, glycolysis and the citric acid cycle are involved in the Warburg effect, and the examination of individual metabolites in these pathways shows that citrate, which is 21-fold increased in cancer tissue (FDR value = 0.005), is also increased in urine (although nonsignificantly in this study) as a possible tumor proxy. It is highly likely that, although tissue glycolysis is

Discussion

The clinical use of omics technologies in cancer research has evolved from the direct evaluation of tumor tissue searching for prognostic markers to the use of biofluids as proxies for changes in the cancer of interest. Although it is clear that the latter approach is more readily translatable to the clinical setting, there are no currently available studies showing the use or accuracy of biofluids as a reflection of metabolic changes within tumor tissue. That a specific biofluid can reflect metabolic changes in a cancer is a reasonable assumption because the concept behind the use of the clinical laboratory for blood chemistries assumes that tests for liver and kidney function, for example, accurately reflect changes that are occurring in these organs. Consistent with this concept, recent metabolomics studies, including some from our laboratory, have assumed that changes in the blood and urine metabolomes accurately reflect alterations in metabolic processes that are occurring in the tumor itself (as well as in any systemic effects of the malignancy), but before this study, this assumption has not been validated. For these reasons, we have undertaken a comprehensive metabolomics analysis of tissue, blood, and urine taken simultaneously from an anatomically accurate RCC mouse xenograft model. In light of the data obtained from this study, it is evident that blood is a decent proxy of tumoral metabolic changes, whereas urine is less reflective of such changes.

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![Figure 4. Comparison of metabolites across matrices. The differences in identified metabolites shown as cancer: control is indicated for each matrix [red, upregulated; blue, downregulated; solid, significant (P < 0.05); unfilled, not significant (P > 0.05)]. A, using raw P values; B, using FDRs. Dotted lines, concordant change across matrices.](image-url)
Multiple Matrix Metabolomics Analysis of Kidney Cancer

Figure 5. PPAR-α modulation affects RCC and other renal epithelial cell lines. The indicated cell lines (NHK, primary human renal tubular epithelial cells) were grown to confluence and incubated continuously with the indicated concentrations (10–100 μmol/L) of the PPAR-α agonist Wy-14,643 or antagonist GW6471. An MTT assay was carried out as described in Materials and Methods. The experiment shown is representative of 3 independent experiments. Asterisks indicate significance at a P value <0.05 when compared with the DMSO-treated control for that cell line.

Figure 6. 1-Methyltryptophan increases an inflammatory marker in cancer cells. The RCC cell lines A498 and ACHN were grown to confluence and treated for 72 hours with 1-methyltryptophan as indicated. Whole-cell protein extracts were then run using 10% SDS-PAGE and incubated with the indicated antibodies. Actin is a loading control and LPS is a positive control. The experiments shown are representative of 3 independent experiments. Densitometry measurements are shown for this blot: incubation of both cell lines with 1-MT at 200 μmol/L was significant (P = 0.015 for A498 and P = 0.034 for ACHN).

Among all measured metabolites, the compound that had the highest magnitude of change was tissue cysteine–glutathione disulfide (CSSG), which was 232.4 times higher in cancer (FDR = 0.0005). Although also identified in serum, this compound was not significantly changed in that matrix and was below the detection limit in urine. Elevated levels of CSSG and GSSG are a signature of oxidative stress and, in light of the finding of significant changes in CSSG, GSSG, and GSH in the xenograft tissue, implicate the oxidative stress pathway as being important in kidney cancer, as has been reported by others in RCC cell lines in general (18). However, in urine and serum, these signature metabolites were below the level of detection and thus, although an important pathway in cancer, glutathione and the oxidative stress pathways are unlikely capable of being biofluid proxies of tumor tissue events or biomarkers or RCC.

The PPARs belong to the nuclear hormone receptor superfamilly and are involved in the transcriptional activation of many target genes regulating energy metabolism, adipogenesis, angiogenesis, cell proliferation, and inflammation (19). The PPAR family consists of 3 subunits (α, β/δ, and γ) that heterodimerize with the retinoic acid receptor (RXR) and bind to DNA to either activate or repress expression of a variety of inflammation-related genes (19). PPAR-α is expressed in many tissues, but it is found at particularly high levels in tissues that require fatty acid oxidation as a source of energy such as liver, kidney, and heart, consistent with its known physiologic role (20); its signature was detected in all 3 matrices in our study. Consistent with our findings that a PPAR-α agonist increases elevated in our analysis consistent with the Warburg effect, it is being shunted into biosynthetic pathways rather than through the citric acid cycle as is evidenced by significantly lower tumor levels of fumarate (0.6525-fold, FDR = 0.0029; see Supplementary Table S2A). In addition, the acylcarnitines, which have been shown to be increased in the urine of RCC patients (17), were found to be increased consistently in tissue (e.g., acetylcarnitine increased 3.2016-fold, FDR = 0.0005; butyrylcarnitine increased 19.2149-fold, FDR = 0.0005) but not in serum and urine in the xenograft model, highlighting a difference between mouse and human. Nonetheless, the data from this study support our earlier data (17) and indicate that human urinary acylcarnitines are excellent proxies of tissue changes and, for this reason, could be developed further as potential biomarkers.

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RCC cell growth, administration of the agonist WY-14,643 has been shown to significantly induce proliferation of the breast cancer cell lines MCF-7 and MDA-MB-231 (21). It has also been proven that long-term feeding of WY-14,643 caused a 100% incidence of liver tumors in wild-type mice, whereas PPAR-α-null mice was refractory to the same treatment (22). Other PPAR-α agonists such as chlorinated paraffin (C12), cinnamyl anthranilate, perchorloethylene, and trichloroethylene also caused kidney tumors in rats (23). In accordance with the previous studies, our results parallel effects with a PPAR-α agonist and antagonist. A previous study with humans showed that administration of Wy-14,643 significantly attenuated urinary cinnamoylglycine (9-fold; ref. 11), a finding that is in agreement with our observations about this metabolite. Attenuation of the PPAR-α pathway may therefore present a novel therapy in treating not only cancer, as evidenced by our data, but also other solid tumors.

The metabolites in the tryptophan and nicotinamide pathways were well represented in the tumor tissue and intersect at the metabolite quinolinolate. We have regularly observed tryptophan metabolites and an altered tryptophan metabolism pathway in our RCC metabolomics and proteomics studies (1, 7), and other investigators have observed a decrease in serum tryptophan in RCCs of all grades (24), similar to what we have observed in this study. In addition to being elevated in mouse tissue and human urine, one of these metabolites, quinolinolate, stimulates proliferation of some cRCC cell lines (1). Perhaps more importantly, IDO, which catalyzes in early step in tryptophan metabolism, regulates conversion of tryptophan metabolites that we have observed in urine and tissue could be seen in the contralateral tissue. For the metabolomics analyses, we have used identical platforms as was used in our previous studies in human urine (1, 2), allowing relationships to be accurately made between species.

In summary, this study represents for the first time a comprehensive metabolomics evaluation of cancer, including tumor tissue as well as the 2 most significant and commonly accessed biofluids, serum and urine. Data presented here show that, not unexpectedly, in kidney cancer blood is a better proxy for tumoral changes than is urine. Furthermore, we have used pathway and network analysis to discover and validate several important metabolic processes that may result in the discovery of novel diagnostic targets and therapeutic approaches for cRCC.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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References


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