Comparative membrane proteomics: a technical advancement in the search of renal cell carcinoma biomarkers†

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Renal Cell Carcinoma (RCC) is the most common kidney cancer, accounting for 3% of adult malignancies, with high metastatic potential and radio-/chemo-resistance. To investigate the protein profile of membrane microdomains (MD), plasma membrane supramolecular structures involved in cell signaling, transport, and neoplastic transformation, we set up a proteomic bottom-up approach as a starting point for the identification of potential RCC biomarkers. We purified MD from RCC and adjacent normal kidney (ANK) tissues, through their resistance to non-ionic detergents followed by ultracentrifugation in sucrose density gradient. MD from 5 RCC/ANK tissues were then pooled and analysed by LC-ESI-MS/MS. In order to identify the highest number of proteins and increase the amount of membrane and hydrophobic ones, we first optimized an enzymatic digestion protocol based on Filter Aided Sample Preparation (FASP), coupled to MD delipidation. The MS analysis led to the identification of 742 ANK MD and 721 RCC MD proteins, of which, respectively, 53.1% and 52.6% were membrane-bound. Additionally, we evaluated RCC MD differential proteome by label-free quantification; 170 and 126 proteins were found to be, respectively, up-regulated and down-regulated in RCC MD. Some differential proteins, namely CA2, CD13, and ANXA2, were subjected to validation by immunodecorration. These results show the importance of setting up different protocols for the proteomic analysis of membrane proteins, specific to the different molecular features of the samples. Furthermore, the subcellular proteomic approach provided a list of differentially expressed proteins among which RCC biomarkers may be looked for.

Introduction

Renal Cell Carcinoma (RCC) is the deadliest tumor of the genitourinary tract. Clear cell renal cell carcinoma represents the most common histotype, accounting for 70–80% of all kidney cancers. Due to its asymptomatic development, approximately 80% of RCCs are discovered incidentally during unrelated diagnostic abdominal imaging, when the disease is already in the advanced phase.1,2 Furthermore, RCC is chemotherapy and immunotherapy resistant, and nephrectomy is the only beneficial therapy.

Once metastases develop, the prognosis for long-term survival is poor; the 5 year survival rate for non-treated patients is usually less than 14%. Nevertheless, there are currently no circulating validated biomarkers able to confirm the identity of renal masses, whether benign or malignant.3,4

Plasma membrane proteomics seems to be a promising tool to address this issue, as membrane proteins perform endocytosis and signaling, and modifications of plasma membrane composition are encountered in several cancers.5,6 Moreover, about two-thirds of membrane proteins are drug targets, hence crucial for therapy advancement.

In particular, within the membrane, there are specialized microdomains, also known as lipid rafts. Membrane microdomains (MD) are small (10–200 nm), heterogeneous, highly dynamic, sterol- and sphingolipid-enriched lateral assemblies that compartmentalize cellular processes. Caveolae, a subclass of such MD, are flask-like invaginations of the plasma membrane that characteristically display caveolin-1 expression.7

MD are highly dynamic structures and act as selective signal transduction mediators, enabling interactions between the
intra- and extra-cellular compartments. Furthermore, they play a key role in drug resistance, cell migration, cell adhesion, and cell survival, as well as in metastasis and tumor progression.\textsuperscript{8–11} Since membrane microdomains represent a minor and highly selected subset of the cellular proteome, they are particularly well suited to investigation using MS techniques. However, their peculiar lipid composition and the enrichment in highly hydrophobic membrane proteins make the assessment of the microdomain proteome a hard issue to address, therefore requiring the optimization of specific protocols. In fact, membrane proteins are usually under-represented in classical proteomic approaches relying on 2D gel electrophoresis (2DE) and MS analysis. This is due to them being generally not very abundant, their isolectric points (pI's) being generally alkaline and their solubility in the aqueous buffers used for isoelectrofocusing (IEF) being poor.\textsuperscript{11–13} Recent data on the quantitative proteomics of MD – lipid rafts or caveolae – have underlined the puzzling role of various signaling proteins in cancer development, but these studies were mainly performed on cell cultures, such as metabolic labelling, are not suitable for the study of human tissue MD, making label-free quantification a valid choice to achieve differential expression profiles. Moreover, the protocol needs to be adapted to small amount of samples, as feasible when working with human surgical specimens.

For these reasons, starting from human surgical samples, we focused on the pre-analytical phase of sample preparation, in order to improve the identification of MD membrane proteins, using a gel-free approach through liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS). We evaluated three different digestion protocols based on the Filter Aided Sample Preparation (FASP) technique, and, after optimization, we assessed differential RCC MD proteomes by label-free quantification, as a starting point for the identification of potential RCC biomarkers.

Materials and methods

Chemicals

Solutions were prepared with Milli-Q water. Water of HPLC grade, trifluoroacetic acid (TFA), isopropyl alcohol, tributyl phosphate (TBP), ammonium bicarbonate (ABC), dithiothreitol (DTT), iodoacetamide (IAA), urea, Triton X-100, Tween-20, Trizma-base, MES, sucrose, BCA protein assay reagent and BSA were from SIGMA Chemical Co. (St. Louis, MO, USA). Acetonitrile (ACN), methanol, and glycerol were from Merck (Darmstadt, Germany). Porcine trypsin was from Promega (Madison, WI, USA). Bradford protein-binding colorimetric assay was purchased from Bio-Rad (Hercules, CA, USA). Anti-protease inhibitor cocktail (Complete) was from Roche (Mannheim, Germany). NuPAGE\textsuperscript{20} SDS-PAGE gel electrophoresis system components (mini gels, running and loading buffers, molecular weight markers and Coomassie brilliant blue staining–CBB) were supplied by Life Technologies (Paisley, Renfrewshire, UK). Hybond-ECL nitrocellulose membrane was from GE (Little Chalfont, Buckinghamshire, UK). Anti-aminopeptidase N (CD13) monoclonal antibody, anti-annexin A2 (ANXA2) and anti-carbonic anhydrase (CA2) polyclonal antibodies were from Abcam (Cambridge, UK).

Species-specific secondary peroxidase conjugated antibodies and ECL reagents were from Pierce (Rockford, IL, USA).

Patients and specimens

Patients with suspected RCC, not receiving any previous chemotherapy, were subjected to radical nephrectomy, after their informed consent and the local research ethics committee approval (U.O. Comitato di Etica e Sperimentazione Farmaci Direzione Scientifica Fondazione IRCCS Ca’Granda Ospedale Maggiore Policlinico, Milano). RCC was classified according to the WHO recommendations\textsuperscript{18} and the 2009 TNM (tumor-node-metastasis) system classification also using immunohistochemical techniques; only samples diagnosed as conventional clear cell RCC were included in the study. Tumor grading was performed according to the Fuhrman grading system.\textsuperscript{19}

Immediately after removal, the pathologist collected samples of primary RCC, selected inside homogeneous areas and avoiding grossly necrotic or fibrotic parts, and of homologous normal cortical tissue (adjacent normal kidney, ANK) contiguous to the tumoral mass. A minimum of one cm\textsuperscript{3} of tumor and normal cortex were placed in sucrose buffered solution (250 mM sucrose, 10 mM Tris-HCl, pH 7.4, 1 mM EDTA), kept on ice and immediately transferred to the laboratory.

The clinical and personal features of the studied patients are summarized in Table 1. Their neoplasms had quite homogeneous characteristics, being in the early phase, and neither metastasis nor positive lymph nodes were present at diagnosis.

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ptT, tumor stage; G, nuclear grade.\textsuperscript{a} Samples used for the preparation of the ANK MD pool in order to optimize the FASP protocol.\textsuperscript{b} Samples used for the label-free quantitation.\textsuperscript{c} Samples analyzed by 1DE.\textsuperscript{20}

Subcellular fractionation and microdomain isolation

After nephrectomy, fresh RCC and ANK tissue samples were submitted to subcellular fractionation through differential
centrifugation; all steps were performed in a cold room (4 °C) or on ice. Subcellular and microdomain-enriched fractions were isolated and characterized as described.20

Electrophoresis and western blotting

Protein separation was performed by the NuPAGE® electrophoresis system (Life Technologies). Equal amounts (10 μg) of subcellular fraction proteins were separated by 4–12% NuPAGE and blotted onto nitrocellulose membranes using a mini tank electrophoretic transfer apparatus (Hoefer). Membranes were developed with the respective primary antibodies (anti-CD13 1 : 2000, anti-ANXA2 1 : 2000, and anti-CA2 1 : 1000), followed by peroxidase-conjugated secondary antibodies (Pierce) and enhanced chemiluminescence detection (SuperSignal West-Pico ECL, Pierce). Images were captured on a CCD camera, LAS4000 (GE Healthcare).

Delipidation and protein precipitation by tri- n-butylphosphate/acetone/methanol

Aliquots (100 μL) of the MD pools were mixed with 14 volumes of ice-cold TBP/acetone/methanol mixture (1 : 12 : 1) and incubated at 4 °C for 90 minutes. Proteins were precipitated by centrifugation at 2800 × g for 15 minutes (4 °C), washed sequentially with 1 mL of TBP, acetone and methanol, and then air-dried.21 During the set-up of delipidation experiments, we assessed a loss of MD proteins of about 20%. Accordingly, we performed delipidation on 150 μg of ANK and RCC MD, since 100 μg of proteins were required for FASP protocols. In order to evaluate the reproducibility of protein extraction, we performed quantitative protein assay using the BCA reagent on delipidated samples and verified the recovery after NuPAGE/CBB staining, by loading equal aliquots of ANK and RCC MD delipidated samples.

Filter aided sample preparation: protocol optimization

Due to the peculiar MD lipid composition, we optimized an enzymatic digestion protocol based on Filter Aided Sample Preparation (FASP),22 comparing three different protocols (Table 2), with and without previous delipidation. For this purpose, equal amounts of MD isolated from ANK tissues of 11 patients (Table 1) were pooled in order to eliminate inter-individual differences.

Protocol 1 was an adapted version of the FASP digestion by Wiśniewski et al.,22 already reported for the MS analysis of urinary exosomes.23 The lysis of MD, about 100 μg of proteins, was performed by a 30 min incubation in RIPA buffer (50 mM Tris-HCl pH 7.4, 1% NP40, 0.25% sodium deoxycholate, 150 mM NaCl, and 1 mM EDTA). MD were then submitted to disulfide bond reduction with 50 mM DTT in 50 mM ABC (95 °C for 5 min). After cooling the sample, the lysates were transferred into the ultrafiltration units (Amicon Ultra-0.5 mL 30 kDa, Millipore) and centrifuged at 14 000 × g for 15 min in order to eliminate lysis buffer and DTT. The filters were washed with 300 μL of 50 mM ABC (14 000 × g for 10 min) and the samples were incubated with 80 μL of 100 mM IAA in darkness for 30 min. IAA was discharged by centrifugation at 14 000 × g for 5 min, and the filters were washed with 200 μL of 50 mM ABC five times (14 000 × g for 10 min). Proteins were digested overnight at 37 °C by 4 μg of trypsin for each sample. After digestion, the filtered tryptic peptides were collected by centrifugation with two washes of water (40 μL and subsequently 100 μL).

In protocol 2, lysis and reduction steps were the same as protocol 1; alkylation was performed by incubating the samples with 100 μL of 50 mM IAA in darkness for 20 min. After IAA elimination, the filters were washed with 100 μL of 50 mM ABC four times (14 000 × g for 15 min). Protein digestion was performed overnight at 37 °C adding 1 μg of trypsin. After centrifugation at 14 000 × g for 10 min, the filtered tryptic peptides were collected, and the filters washed with 50 mM ABC and 500 mM NaCl, two times each. The eluted peptides were acidified by 0.1% TFA.

In protocol 3, the sample was solubilized by a denaturation buffer containing SDS and DTT (2% SDS, 50 mM DTT, 50 mM Tris-HCl pH 7.6), incubated at 95 °C for 5 minutes. Dissolved MD membranes (70 μL) were mixed with 430 μL of 8 M urea in 100 mM Tris-HCl, pH 8.5 (UA 8.5 solution), transferred into the ultrafiltration device and centrifuged at 14 000 × g for 15 minutes. Alkylation was performed as in protocol 2, followed by four washes, two with 8 M urea in 100 mM Tris-HCl, pH 7.9 (UA 7.9 solution) and two with 50 mM ABC (14 000 × g for 15 min). Protein digestion and peptide elution were the same as protocol 2.

Digestion and peptide extraction

MD isolated from RCC and ANK tissues of 5 patients were pooled (Table 1), and the resulting samples were lysed and subjected to protein extraction and digestion by the FASP protocol 3, after delipidation.

MD proteome analysis by nLC-ESI-MS/MS

All the MS analysis and protein identification were performed under the same conditions. Before LC-MS analysis, tryptic digests for each sample were quantitated using a NanoDrop spectrophotometer and desalted using ZipTip® μ-C18 (Millipore) following manufacturer's instructions.24 About 1 μg of digested proteins for each sample run were injected at least 3 times into a Dionex UltiMate 3000 rapid separation (RS) LC nanosystem (Thermo Scientific, Germany) coupled online with a nano ESI ion trap mass spectrometer, amaZon ETD (Bruker Daltonics GmbH, Germany). After the μ-trapping column desalting (Dionex, Acclaim PepMap 100 C18, cartridge, 300 μm ID × 5 mm, 5 μm) and concentration steps, peptides were then separated.
by a 360 min-multistep gradient on the analytical 50 cm column (Dionex, 0.075 mm ID, Acclaim PepMap100, C18, 2 μm) with a flow rate of 300 nl min$^{-1}$ as already described. The ion-trap mass spectrometer equipped with an on-line nanospray source was operated in data-dependent-acquisition mode. For MS generation, enhanced resolution and a trap ICC value of 400 000 were used; for MS/MS acquisition, the ICC target was increased to 1 000 000; a narrow range for Smart Fragmentation from 50 to 150% was adopted. CID MS/MS fragmentation was set to fragment the ten most abundant MS peaks with strictly active exclusion after one spectrum and released after 9 seconds. The obtained chromatograms were elaborated using Compass DataAnalysis™, v.4.0 Sp4 (Bruker Daltonics, Germany), and the resulting mass lists were processed using an in-house Mascot search engine (v.2.4.0), through Mascot Daemon. Database searching was restricted to the human Swissprot database (accessed Apr 2014; 544 996 sequences; 193 815 432 residues). Trypsin as an enzyme and carbamidomethyl (C) as a fixed modification were set in search parameters. Mass tolerance for all modifications was generally fixed at 2 Da for the precursor ions and 0.8 Da for the product ions. Automatic decay database search and a built-in Percolator algorithm were applied to calculate the posterior error probabilities for each peptide-spectrum match and rescoring search results with a unique significance threshold. Data were filtered using a global false discovery rate < 1% and only proteins with at least one unique identical peptide sequence (p-value < 0.05) were considered identified.

Label-free protein abundance evaluation

Protein abundance in MD of RCC and ANK tissues was determined using Progenesis LC-MS software v 4.1 (Non-linear Dynamics, Newcastle, England). The raw LC-MS/MS data were imported and the ion intensity maps of all runs (3 runs for each class) were visually examined for defects and used for the alignment process, performed using the “Automatic Alignment” function. One sample is thus set as the reference run and the retention times of all other samples within the experiment were automatically aligned to create a maximal overlay across the data. The default sensitivity and a peak width of 0.2 min were selected as parameters for the feature detection algorithm. Samples were then allocated to their experimental class (RCC vs. ANK). The identification of peptides was performed using an in-house Mascot search engine as described above. The filtered MASCOT search results were imported back into Progenesis LC-MS and the identified peptides with a score lower than 13 were discarded (p-value > 0.05). Experimental variation affecting the protein expression was minimized by analyzing the same amount of sample (1 μg) and by the normalization process performed using Progenesis software (http://www.nonlinear.com/progenesis/qi-for-proteomics/v2.0). Protein quantification results were exported and used for further analysis.

Bioinformatic analysis

Gene ontology (GO) analysis was performed using the LocDB database (www.rostlab.org/services/locDB/) for investigating the subcellular localization of the identified proteins and the Protein ANalysis THrough Evolutionary Relationship (PANTHER) database (http://www.pantherdb.org/) for the analysis of their molecular functions. In addition, membrane proteins were deeply studied and transmembrane helix prediction was performed using the online TMHMM 2.0 prediction server (http://www.cbs.dtu.dk/services/TMHMM/). Palmitoylation was predicted using CSS-Palm 2.0, GPI-modification site prediction was conducted through the online big-PI predictor server (http://mendel.imp.ac.at/sat/gpi/gpi_server.html) and myristoylation was predicted using the online Expasy Myristoylator prediction server (http://web.expasy.org/myristoylator/). Protein–protein interactions were predicted using the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) database v9.0 (http://www.string-db.org/) and proteins were linked based on the following six criteria: neighbourhood, gene fusion, co-occurrence, co-expression, experimental evidence and existing databases.

Tissue specificity of the identified proteins was evaluated using different databases available in the Human Protein Atlas (HPA) (http://www.proteinatlas.org/). In particular, MD proteins were compared with: (i) the human specific proteome of 32 tissues, summing up “tissue enriched”, “group enriched” and “tissue enhanced” gene categories; (ii) the housekeeping proteome (proteins detected in all tissues); and (iii) the renal cancer proteome, considering only the proteins detected in at least 80% of the HPA analysed patient tissues.

Results and discussion

Protocol setup

For the optimization of the digestion protocol, we pooled MD isolated from ANK tissues of 11 patients in order to eliminate inter-individual differences. Then we subjected aliquots of 100 μg of proteins to the three protocols, and analyzed them by LC-MS/MS.

In order to select the best protocol, we took into consideration: (i) the number of total proteins identified by the Mascot search engine; (ii) the amount of hydrophobic proteins on the basis of the hydrophobicity score, GRAVY index (assigned by ProtParam http://web.expasy.org/protparam/); and (iii) the percentage of membrane proteins [classification assigned by LocDB and confirmed by TMHMM 2.0].

We first tested an adapted version of FASP digestion, already used for the MS analysis of membrane proteins (hereafter named “Protocol 1”) (Table 2). FASP is a method that combines strong detergents for universal solubilization with efficient pre-digestion ‘clean up’ of the proteome in order to obtain purified peptides, avoiding the disadvantages of the gel format. The main feature of Protocol 1 is the lysis of MD, performed by a 30 min incubation in lysis RIPA buffer. The resulting analysis with LC-MS/MS led to the identification of only 32 proteins with a few signals of low intensity shown by the base peak chromatogram (ESI, Fig. S1).

Due to the peculiar lipid composition of MD, we introduced a step of protein delipidation and precipitation with tri-n-butyl-phosphate, acetone, and methanol. This pretreatment ensured the highest protein recovery, according to Shevchenko and
Using delipidation followed by Protocol 1, 103 proteins were identified (Fig. 1), a number still lower than in our previous study (286 proteins). We then tested another protocol (Protocol 2), by adding NaCl washing, in order to increase the recovery of hydrophobic peptides, and one acidification step with 0.1% TFA, on both delipidated and non-delipidated samples (Table 2). These settings determined an increased, yet not fully satisfactory, recovery of proteins (Fig. 1).

The conditions were then further modified in Protocol 3, which, unlike the other protocols, involves the use of SDS buffer and urea (Table 2). In particular, we performed all washings with 8 M urea, which should fully wash out detergents from membrane proteins. When applied without delipidation, this protocol led to the identification of 170 proteins, of which only 23% were transmembrane proteins (Fig. 1). This could be due to the lipids-SDS interaction: as a consequence, the SDS cannot be efficiently removed by the filter, interfering with the MS analysis (ESI, † Fig. S1). Moreover, the filter may retain some lipid components, together with hydrophobic lipid-bound proteins.

Conversely, Protocol 3 coupled to delipidation was the most efficient way to identify proteins in MD. In fact, firstly, the introduction of the delipidation step resulted in a significant increase in the yield of total protein identification (N = 342). Secondly, the total number of membrane proteins greatly increased (N = 222), boosting the percentage of transmembrane proteins to 50%.

These results may depend on the depletion of the lipid components, which likely enables the “release” of transmembrane proteins. Once freed, they become more suitable for trypic digestion, thus generating peptides that can be eluted through a 30-kDa cut-off FASP filter.

**LC-MS protein identification and label-free quantification**

The optimized protocol was implemented to perform the comparison between RCC and ANK MD. In fact, we speculated that the delipidation step could eliminate some differences in protein identification due to the different lipid composition (data not shown), making the comparative proteomic analysis more reliable. We first evaluated the recovery after MD delipidation as shown in the ESI, † Fig. S2. The extraction of ANK and RCC MD proteins after delipidation is reproducible, and the protein profiles were comparable with the non-delipidated ones (ESI, † Fig. S2). Proteins of microdomain-enriched fractions prepared from 5 paired samples of RCC and ANK tissues were pooled and analyzed. Identification was accepted with at least one unique peptide exceeding the Mascot score of identity cut-off, in order to ensure a better understanding of MD proteomes. Exploiting the powerful peak detection of the Progenesis algorithm (used during the label-free protein expression evaluation), and applying the above detailed criteria, MS/MS analysis after FASP digestion led to the identification of 742 proteins in ANK and 721 in ccRCC MD. Overall, more than 800 protein species were identified; their full details, including scores and the number of matched peptides, are given in the ESI, † Tables S1 and S2.

The molecular characteristics of the identified proteins were assessed by bioinformatics and prediction tools. The results showed that the majority of the identified proteins (53.1% and 52.6% in ANK and RCC, respectively) were membrane-associated, with about 58% of them being transmembrane proteins (ESI, † Tables S1 and S2). Many of them were typical raft proteins, such as caveolins (1 and 2), flotillins, aquaporin-1, prohibitins, VDAC1 and Thy-1. Comparing our list of total MD identified proteins to RaftProt, a recently published Mammalian Lipid raft Proteome Database (http://lipid-raft-database.d.uq.edu.au/index.html), 580 (72%) were classified as human lipid raft associated proteins and 394 (49%) as High Confidence lipid raft proteins. Moreover, we investigated whether protein sequences contained sites for post-translational modifications such as glypiation, palmitoylation, and myristoylation, using prediction algorithms. They assigned putative myristoylation sites to about 15% of total identified membrane proteins and GPI anchoring to 5% of them. Much more frequent, about 44%, was the prediction of palmitoylation sites. Additionally, STRING analysis classified...
67% of proteins as constituents of protein complexes. These findings are intriguing, because both such post-translational modification and protein–protein interactions are likely to determine a possibly reversible association with the membrane MD of otherwise soluble proteins, allowing for their dynamic binding to microdomains. After applying these tools for the prediction of direct or indirect membrane interactions, only 79 proteins (9.8%) were left out.

These results are in good agreement with a recently published work, concerning chicken inner ear membrane MD proteomics. Mass spectrometry of MD fractions identified over 600 putative raft proteins, and most of them were predicted as membrane-associated and involved in trafficking and metabolism.

Very few other papers broached tissue MD proteomics. In particular, a study by Yu et al. adopted a “tube-gel” protein digestion label-free shotgun proteomic strategy to quantify raft proteins in the neonatal mouse brain – about 200 raft proteins were identified from a single sample, thus defining a core proteome qualitatively quite similar to ours and previous ones.40 In fact, membrane MD, on the one hand, are highly dynamic, with protein (and lipid) components shuttling rapidly between raft and non-raft membranes, and on the other hand, present a fixed and common proteomic core, accounting for their structural and specific characteristics.

In our previous work, we analyzed the same RCC and ANK MD samples by a gel based method (Table 1). The present optimized gel-free approach seems to be more suitable for the study of membrane microdomains, providing a higher number of total identified proteins (742 vs. 98 in ANK MD and 721 vs. 93 in RCC MD). It is noteworthy that more than 88% of the proteins identified with the gel-based approach were also found using the shotgun proteomic strategy (Fig. 2), confirming their localization in the MD.

We assessed the tissue specificity of the MD proteome, comparing the new list of MD proteins with the Human Protein Atlas database (HPA) – a comparison with the human tissue-specific proteomes of 32 different tissues showed a peak matching with kidney-specific ones (Fig. 3). Moreover, 30.5% of proteins identified in RCC MD were already detected in HPA renal cancer tissue microarrays. It has to be stressed that subcellular fractionation, and in particular MD purification, greatly enhances the possibility to identify low-abundance proteins, undetectable in the whole tissue. Several protein classes (i.e. G proteins, RAS-related proteins, components of the vacuolar ATPase–synthase complex, etc.) cannot indeed be tissue specific because they are typical for the MD proteome, regardless of their tissue localization.

We then evaluated the differential proteome by label-free quantification, considering proteins to be “up” or “down-regulated” when the ratio was, respectively, higher than 1.50 and lower than 0.67. Applying these criteria to 657 quantified protein species, 170 (25.9%) were classified as “up” and 126 (19.2%) as “down” in RCC MD (ESI,† Table S1).

The differential proteins (“up” and “down”) were correlated with subcellular localization (by LocDB) and molecular function (by Panther) (Fig. 4). The majority of “down” proteins were membrane-associated (70.6%) (Fig. 4A); this result could be
dependent on the process of de-differentiation typical for cancer cells, which lose their tubular epithelial specialization. In fact, among “down” proteins, we found aquaporin-1 (AQP1), a water-channel that is considered a kidney differentiation marker, Na+/glucose co-transporters 2 and 5, involved in glucose reabsorption by proximal tubule cells, and renal dipeptidase (DPEP-1), a zinc-dependent membrane metalloprotease. In contrast, the “up” proteins are found to be partially membrane (42.3%) and partially cytoplasmic proteins (38.2%) (Fig. 4A). It has to be underlined that some cytosolic proteins may associate with the plasma membrane and MD, through cytoskeletal components. Several cytoskeleton associated-proteins, such as ezrin, radixin and moesin, members of the ezrin–radixin–moesin (ERM) family, involved in cancer progression, are in fact increased in RCC MD. Interestingly, focusing on post-translational modifications, the predicted palmitoylation sites appeared nearly doubled (from 42.5 to 71%) among “up”, compared to “down” proteins. It is tempting to speculate that this behavior is driven by a modification of the RCC MD lipid composition, thus allowing cancer cells to deliver new proteins, perhaps involved in aberrant signaling events, towards MD. The elucidation of this hypothesis deserves further investigation.

By analyzing this list of proteins using Panther, an on-line tool for the classification of genes according to the Gene ontology classes, we obtained the picture shown in Fig. 4B. From a functional point of view, the presence of neoplasm seems to increase some peculiar protein classes involved in signaling and adhesion, typical MD-associated processes. In fact, G proteins, structural, and binding proteins (e.g., flotillins, tubulins, annexins, and caveolins), were more abundant in RCC MD, compared to ANK.

It has to be underlined that most of differential proteins in our previous gel-based study were confirmed here, after label free quantification, for example, Basigin, DPEP1, AQP1, and caveolin-1 (ESI, Table S1). This result may indeed represent a preliminary validation.

Validation of differential RCC MD protein content by immunoblotting analysis

Relative quantification results obtained by the label-free approach were verified by immunoblotting of three differential proteins, carbonic anhydrase 2 (CA2), annexin A2 (ANXA2), and aminopeptidase N (CD13), comparing the signal intensity in different subcellular fractions prepared from ANK and RCC (Fig. 5).

The choice of these three proteins was firstly based on their significantly high or low RCC/ANK ratio (CA2, 2.54; ANXA2, 1.97; CD13, 0.37) obtained after label-free quantification. Secondly, ANXA2 and CD13 were already reported as MD associated, and listed as High Confidence Raft Proteins in the RaftProt database. On the other hand, CA2 is a typical cytosolic enzyme, responsible for more than 95% of the total kidney carbonic anhydrase activity and it is not usually described as a MD bound protein.

WB results regarding ANXA2 confirmed label-free quantification, being up-regulated in RCC samples, compared to ANK. This difference is higher at the MD level than in whole lysates, suggesting the occurrence of an altered subcellular localization.
in the RCC tissue. ANXA2 is actually considered a marker of several cancer cells (reviewed in ref. 47), and its overexpression in RCC has already been reported.\(^{48}\) Its key role in the organization of lipid raft signaling domains by binding to membrane phospholipids, under physiological conditions, was also described.\(^{49}\)

A very recent paper\(^{50}\) outlines the regulatory role of ANXA2 in RCC cell motility and the change in ANXA2 localization from cytoplasm to membranes in the RCC tissue (by immunohistochemistry), supporting our findings.

Aminopeptidase N is a transmembrane zinc-dependent ectoenzyme, expressed by various tissues, including the kidney.\(^{51}\) Besides its peptidase activity, it is physiologically involved in endocytosis and signaling.\(^{52}\) CD13 is down-regulated in RCC MD, as determined by label-free analysis; accordingly, its signal was undetectable in RCC subfractions, while highly enriched in RCC MD level, while undetectable in the corresponding ANK MD. These data suggest a possible tumor-dependent mis-localization of CA2 in RCC MD. Interestingly, an increased expression of CA2 has been reported in the neovessel endothelium of several tumors, such as melanoma and esophageal, renal, and lung cancer, but not in the corresponding healthy vascular cells.\(^{57}\) Since we prepared fractions from whole RCC tissue, the signal intensity detected in MD may indeed be derived from tumoral neovessels.

Overall, the validation of these three proteins further supports the potency of label-free MS approach.

Conclusions

In conclusion, our results show the importance of setting up a specific protocol for the proteomic analysis of membrane proteins, according to the different molecular features of the sample. Moreover, our data confirm that the sensitivity of proteomic profiling can be significantly enhanced by focusing on highly enriched subcellular fractions, such as MD. The optimization of all analytical steps, from sample pre-fractionation and preparation to the MS analysis and label-free quantification, provided us with a large panel of differential RCC MD proteins, among which tumor biomarkers may be looked for.

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References