Proteomic analysis of energy metabolism and signal transduction in irradiated melanoma cells

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Abstract
• AIM: To analyze proteomic and signal transduction alterations in irradiated melanoma cells.
• METHODS: We combined stable isotope labeling with amino acids in cell culture (SILAC) with highly sensitive shotgun tandem mass spectrometry (MS) to create an efficient approach for protein quantification. Protein–protein interaction was used to analyze relationships among proteins.
• RESULTS: Energy metabolism protein levels were significantly different in glycolysis and not significantly different in oxidative phosphorylation after irradiation. Conversely, tumor suppressor proteins related to cell growth and development were downregulated, and those related to cell death and cell cycle were upregulated in irradiated cells.
• CONCLUSION: Our results indicate that irradiation induces differential expression of the 29 identified proteins closely related to cell survival, cell cycle arrest, and growth inhibition. The data may provide new insights into the pathogenesis of uveal melanoma and guide appropriate radiotherapy.
• KEYWORDS: melanoma cell; 2D-LC-MS/MS; stable isotope labeling with amino acids; proteomic analysis; X-ray irradiation; protein-protein interaction
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INTRODUCTION
Uveal malignant melanoma is the most frequent primary intraocular neoplasm in adults [1]. Radiotherapy is a highly effective treatment option in cancer management; together with surgery and chemotherapy, it plays an important role in the treatment of 40% of patients who undergo radiotherapy are ultimately cured of their cancer [2]. Although radiotherapy has rapidly developed in recent years, the underlying cellular and molecular mechanisms of radioresistance are not fully understood. Furthermore, high doses of radiation can increase the risk of injury to normal tissues and lead to side effects. Thus, it is imperative to study the sensitivity of uveal melanoma cells to radiotherapy. Although a previous study reported that melanoma cells undergo cell cycle arrest after being exposed to high levels of irradiation, the molecular mechanism is not clear [3]. Furthermore, studying the complicated molecular changes in melanoma cells exposed to ionizing radiation is also challenging [4]. Quantitative proteomic provides a novel way to investigate large numbers of differentially expressed proteins in single experiments. In recent years, liquid chromatography coupled with tandem mass spectrometry (LC-MS/MS) has been increasingly used for proteomic studies because it has the advantages of high throughput and sensitivity, especially when combined with stable isotope labeling with amino acids in cell culture (SILAC) to create an efficient approach for large-scale protein quantification [5-7].

Aerobic glycolysis is generally accepted as a metabolic hallmark of cancer [8], but its relationship with cancer progression after irradiation is still unclear. New insights suggest that it would be useful to target tumor metabolism [9]. Recent research found that aerobic glycolysis would be closely associated with cancer cells survival and growth [10-13]. In addition, tumor progression is characterized by alterations in cell signaling [14]. With MS technologies and bioinformatics tools, it is possible to track and analyze signaling events in irradiated cancer cells. Recent research has shown that many signaling proteins regulate cancer cell growth and metabolism, including RAC-alpha serine/threonine-specific protein kinase (Akt), mammalian target of rapamycin (mTOR), cellular tumor antigen p53 (TP53), and others [15-17].
Furthermore, mapping signaling and metabolism protein interactions might help identify the molecular mechanisms at work in irradiated melanoma cells. Indeed, many studies have reported that energy metabolism is closely associated with radioresistance and radiosensitivity\cite{18,19}.

In this study, we applied SILAC technology coupled with 2D-LC-MS/MS and quantitative proteome and interaction network analysis to elucidate metabolism and signaling pathways in irradiated melanoma cells. The quantitative proteomics data described here provide new insights into novel targets for radiotherapy.

**MATERIALS AND METHODS**

**Materials**

**Cell culture** Human choroidal malignant melanoma cell line 92-1 was kindly provided by Modern Physics researchers at the Chinese Academy of Science. All cells were kept in a humidified atmosphere of 95% air and 5% CO\textsubscript{2} at 37\textdegree C, and 5\times10\textsuperscript{4} cells were seeded in 25cm\textsuperscript{2} culture flasks 2 days before irradiation, which resulted in <70% confluence at the time of irradiation. For SILAC experiments, the 15 hours post-irradiation group cells were cultured in heavy Dulbecco’s modified essential medium (DMEM) containing L-arginine-\textsuperscript{15}N, and control cells were maintained in light L-arginine-\textsuperscript{15}N, medium.

**Irradiation** After 48 hours normal incubation, cells were irradiated at room temperature using a Pantak-320S generator (Shimadzu, Tokyo, Japan) operated at 200kVp and 20mA with 0.5-mm Al and 0.5-mm Cu filters at a dose of 10Gy and were then cultured for 15 hours. The dose rate was 1Gy/min. After 15 hours further culture, SILAC cells were washed three times with ice-cold (0-4 \textdegree C) phosphate-buffered saline (PBS) separately for extraction.

**Sample preparation** Cells were scraped into 8mol/L urea and sonicated at 4\textdegree C to lyse the cells. After centrifugation for 30 minutes at 20 000xg, the supernatants were collected and kept at -80\textdegree C until analysis. Protein concentrations were measured using the Bradford method. Extracted protein samples from heavy and light serum were combined at a 1:1 ratio. The samples were dissolved in 8mol/L urea and 25mmol/L NH\textsubscript{4}HCO\textsubscript{3}, and reduced with 10mmol/L DTT for 1 hour at room temperature. Samples were alkylated by 40mmol/L iodoacetamide in the dark for 1 hour at room temperature, and then 40mmol/L DTT was added to quench the iodoacetamide for 1 hour at room temperature. After diluting 8mol/L urea with 25mmol/L NH\textsubscript{4}HCO\textsubscript{3} to 0.6mol/L, trypsin was added at a ratio of 1:40 and digested at 37\textdegree C overnight. In order to completely cleave the proteins, trypsin was added to the sample at a ratio of 1:40 again and digested at 37\textdegree C for 8 hours. Finally, trypsin digestion was stopped by adding formic acid (FA) for a 1% final concentration.

**Methods**

**2D–LC–MS/MS analysis** The samples were analyzed by two-dimensional liquid chromatography (2D-LC) on an LTQ mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) according to a previously published method \cite{20}. The tryptic peptide mixtures were analyzed by 2D-LC coupled to a linear ion trap mass spectrometer LTQ (ThermoElectron, Waltham, MA, USA). For each experiment, the peptide mixtures (from about 100µg proteins) were pressure-loaded onto a biphasic silica capillary column (250µm inner diameter [i.d.]) packed with 3cm reverse-phase C18 resin (SP-120-3-ODS-A, 3mm, the Great Eur-Asia Sci&Tech Development, Beijing, China) and 3cm strong cation exchange resin (Luna 5µm SCX 100A, Phenomenex, Torrance, CA, USA). The buffers used were 0.1% FA (buffer A), 80% acetonitrile (ACN)/0.1% FA (buffer B), and 600mmol/L ammonium acetate/5% ACN/0.1% FA (buffer C). After sample loading, the biphasic column was first desalted with buffer A and then eluted using a 10-step salt gradient ranging from 0 to 600mmol/L ammonium acetate. After each salt gradient, a gradient of buffer B ranging from 0 to 100% was applied. Step 1 consisted of a 100-minute gradient from 0 to 100% buffer B. For steps 2-9, after equilibrating with buffer A for the first 3 minutes, X% buffer C was applied for 5 minutes, and peptides were eluted using a linear gradient as follows: 0%-10% buffer B in 5 minutes, 10%-45% buffer B in 77 minutes, 45%-100% buffer B in 10 minutes, and 100% buffer B for 10 minutes, followed by re-equilibration with buffer A for 10 minutes. The 5-minute buffer C percentages (X) were 5%, 10%, 15%, 20%, 25%, 35%, 50%, and 75%. The gradient used in the final step contained 3 minutes of 100% buffer A, 20 minutes of 100% buffer C, a 5-minute gradient from 0% to 10% buffer B, a 72-minute gradient from 10% to 55% buffer B, and a 5-minute gradient from 55% to 100% buffer B. Then, 100% buffer B was applied for 5 minutes, followed by a 5-minute elution with buffer A and another 10-minute elution with buffer B. The effluent of the biphasic column in each case was directed into an in-house-packed 10-cm C18 analytical column (100µm i.d., SP-120-3-ODS-A, 3mm) with a 3- to 5-µm spray tip. The flow rate at the tip was maintained at about 500nL/min. Nano-electrospray ionization was performed at a spray voltage of 1.9kV and a heated capillary temperature of 170\textdegree C. The MS instrument was set to the data-dependent acquisition mode with dynamic exclusion turned on, and maximum ion injection time was set to 100ms. One MS survey scan, with mass range 400-2000m/z, was followed by five MS/MS scans. All tandem mass spectra were collected using a normalized collision energy (35% setting), an isolation window of 2Da, and 1 micro-scan. The XCalibur data system (ThermoElectron, Waltham, MA, USA) was used to control the high- performance liquid chromatographic (HPLC) solvent gradients and apply the MS scanning functions. MS data were searched using SEQUEST.
algorithm (Ver. 2.8) against the human database, which was released on May 27, 2008, and contains 37,869 protein sequences. The database was reversed and attached to estimate the false discovery rate (FDR).

**Western blot analysis** An antibody against lactate dehydrogenase B (LDHB) (sc-100775) was purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Cells were collected and lysed in appropriate amounts of lysis buffer (Biuyantian, Nanjing, China). The samples were mixed with sample buffer (250mmol/L Tris HCl, 5% β-mercaptoethanol, 50% glycerol, 10% sodium dodecyl sulfate [SDS], and 0.5% bromophenol blue) and boiled for 5 minutes, and equal amounts of protein (30 μg) were separated on 10% SDS-polyacrylamide gels (Bio-Rad, Hercules, CA, USA). Blotting was performed at 120 V for 1.5 hours in a wet transfer instrument (Bio-Rad). The membranes were then washed three times with PBS containing 0.1% Tween-20 and incubated with secondary antibody for 1 hour. Finally, the membrane was washed, protein bands were visualized using an enhanced chemiluminescence system (Amersham-Buchler, Braunschweig, Germany), and the membranes were exposed to X-ray medical film (Kodak, Rochester, NY, USA).

**Cell cycle assay** The cells were incubated for several different phases post-irradiation before they were collected and fixed for 24 hours at -20°C with 70% ethanol. The fixed cells were washed with PBS, treated with 100 μg/mL RNase A and 50 μg/mL propidium iodide for 30 minutes. Cell cycle distribution was analyzed with Modfit software (Verity Software, Topsham, ME, USA) from the histogram of DNA content measured with a flow cytometer (FACSan, Becton Dickinson, Franklin Lakes, NJ, USA).

**Cell proliferation assay** Exponentially growing cells were seeded on sterile cover slips at a density of 1 x 10⁶/well, incubated for 48 hours later, irradiated with X-rays, fixed in 4% paraformaldehyde for 10 minutes and in methanol at -20°C for 5 minutes, permeabilized with 0.5% Triton X-100 for 1 hour, blocked in buffer (5% skim milk with 0.5% Triton X-100) for 2 hours, and stained with a primary Ki67 antibody at 1:50 dilution (Cell Signaling Technology, Danvers, MA, USA) for 2 hours. The bound antibody was visualized using Alexa Fluor 594 anti-rabbit antibody (Molecular Probes, Eugene, OR, USA), and cell nuclei were counterstained with DAPI (4′,6-diamidino-2-phenylindole) solution (Invitrogen, Carlsbad, CA, USA). Photos were taken with a fluorescent microscope (Keyence, Tokyo, Japan). Ki67-positive cells were counted by using a fluorescent microscope, and at least 500 cells were scored for each sample.

**Glycolysis production measurements** Lactate production was measured using an enzymatic kit (Nanjing, Jiancheng, China) according to the manufacturer’s instruction. The results were normalized by cell counts. Briefly, NAD⁺ was added to media and stoichiometrically converted to NADH by lactate in the media. The levels of NADH were then colorimetrically quantified.

**Data analysis and bioinformation** Resulting data were collected using Xcalibur and interpreted by SEQUEST of Bioworks 3.3.1 (Thermo Fisher Scientific) using the NCBI database. Bioworks was used to analyze and filter peptide identifications, i.e., Xcorr ≥ 1.8 (z=1), 2.2 (z=2), 3.5 (z=3), Sp ≥ 500, Rsp ≥ 5, proteins with Number of peptide ≥ 2, and Consensus score ≥ 10. Subsequent analysis included assigning peptides to the spectra, validating the peptide assignments to remove incorrect results, determining relative quantitation ratios between heavy and light isotopic labeling, and inferring protein identifications from the assigned peptides. An open source tool called TPP (Version 4.4.1) could perform these steps: PeptideProphet validates peptides assigned to MS/MS spectra [21], XPRESS program quantitate peptides [22] and proteins in differentially labeled samples, and ProteinProphet [23] infers sample proteins. In addition, the minimum peptide length was considered as seven, and results below the PeptideProphet probability at 0.95 were filtered out. The annotations and biologic processes of proteins were obtained from the Human Protein Reference Database (HPRD Version 04-13-10, http://www.hprd.org). For proteins without descriptions, annotations were identified by searching the UniProt and National Center for Biotechnology Information (NCBI) protein databases. This study classified proteins according to go ontology (GO) biology processes from the HPRD database. Proteins were considered to be significantly dysregulated if the fold change in protein expression between sham and irradiated samples was ≥ ± 1.50 and P≤ 0.05 (t-test).

**Protein–protein interaction network analysis** The identified proteins were mapped to networks available in the STRING 9.0 database (http://string-db.org/) and displayed by Cytoscape 2.8.1 (http://www.cytoscape.org/). In this study, Ingenuity Pathway Analysis (IPA Ingenuity Systems, www.ingenuity.com) was applied to obtain information regarding the relationships, biological mechanisms, functions, and pathways of differentially regulated proteins. The fold change with log₂ ratio and IPI accession number of dysregulated proteins were submitted to IPA.

**Statistical Analysis** Statistical analysis was performed using Statistical Product and Service Solutions (SPSS13.0). The band densities were quantified as fold-change using Student's t-tests based on the mean of three replicates. P value of 0.05 or less was considered significant.

**RESULTS**

Identification and quantification of dysregulated proteins The simple LC-MS/MS workflow is depicted in Figure 1. In the filtered results, 189 proteins involved in cell communication and signal transduction (CC and ST) and
metabolism and energy pathways (M and EP) were identified, among which 19 proteins were upregulated, 29 were downregulated, and 141 were unchanged. These proteins were sorted by GO from the HPRD Database. LDHB was identified by 13 unique peptides, and the coverage ratio was 47.3% (Supplementary File). In Table 1, we listed the 10 proteins with the largest increase in expression levels and the 10 proteins with the largest decrease. Less dysregulated proteins are listed in Supplementary file 3 and were mainly involved in metabolism processes. For example, proteins such as SOD3, GSS, and ST13 have oxidative reduction functions, suggesting that the cells were under oxidative stress. Some slightly upregulated proteins, such as ATP5O, COX6A1, and HADHA, were mitochondrial and involved in ATP production. Other minor upregulated proteins such as NANS, CNP, and GNPDA1 were mainly involved in glycolysis and nucleotide metabolism, indicating that the cells maintained high metabolism levels to combat the stress of radiation.

**Western blotting analysis** LDHB and PFKM proteins were selected to verify MS data. The SILAC ratios of LDHB and PFKM showed dramatic up- and downregulation post-irradiation with 10-Gy X-rays, respectively (Figure 2). The expression changes of LDHB and PFKM were confirmed by Western blot analysis, which showed that changes were basically identical with SILAC (Table 2).

**Phenotype assessments** We performed cell cycle assays and found that irradiated cells were subject to cell cycle arrest (Figure 3A), similar to as reported in previous studies [1]. The lactate production level was used to investigate glycolysis.

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**Table 1 Proteins with largest increased and decreased expression in melanoma cells post-irradiation for 15 hours**

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Xpress ratio</th>
<th>Unique peptides</th>
<th>Coverage (%)</th>
<th>Description</th>
<th>Biological process</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDHB</td>
<td>11.11±0.04</td>
<td>13</td>
<td>47.3</td>
<td>lactate dehydrogenase B ferrochelatase isom b precursor guanine nucleotide binding protein (G protein) tumor necrosis factor receptor 1 precursor acetyl-Coenzyme A acetyltransferase 1 hexokinase 1</td>
<td>metabolism (GO:0008152); energy pathways (GO:0006091); signal transduction (GO:0007165)</td>
</tr>
<tr>
<td>PFKM</td>
<td>0.55±0.06</td>
<td>1</td>
<td>73.0</td>
<td>Macrophage migration inhibitory factor peroxisomal enoyl-coenzyme A hydratase-like protein adenylate kinase 2 isoform</td>
<td>metabolism (GO:0008152); energy pathways (GO:0006091); signal transduction (GO:0007165)</td>
</tr>
</tbody>
</table>

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**Table 2 Comparison of SILAC ratio and Western blot quantitation**

<table>
<thead>
<tr>
<th>Gene name</th>
<th>SILAC ratio</th>
<th>Normalized ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDHB</td>
<td>11.11±0.04</td>
<td>2.48±0.23</td>
</tr>
<tr>
<td>PFKM</td>
<td>0.55±0.06</td>
<td>0.74±0.05</td>
</tr>
</tbody>
</table>
alterations. The lactate generated at the end of the glycolysis pathway and the level of lactate production would reflect the glycolysis flux. Figure 3B shows that the level of lactate production was significantly higher post-irradiation, indicating that cells increased glycolysis in response to irradiation. Cell proliferation post-irradiation was another area of interest in this study. Ki67 is a widely accepted marker for determining cell proliferation \[24\]. As shown in Figure 3C, Ki67 immunoreactivity was significantly decreased post-irradiation, which indicated that radiation inhibited proliferation.

Changes in energy pathway protein levels induced by irradiation We assessed changes in energy pathway protein levels in melanoma cells 92-1 after irradiation. Differentially expressed proteins in response to 10-Gy X-Ray irradiation are depicted in Figure 4. Energy pathways are separated into four types by dashed lines: (glycolysis, TCA cycle, oxidative phosphorylation, and pantose phosphate pathway). Expression fold change $\geq 1.5$ or $\leq 0.67$ was considered significant \[25\]. Previous research demonstrated that cell cycle arrest in 92-1 melanoma cells is induced by irradiation with 10Gy for 15 hours \[3\]. HK1, LDHB, COX6A1, and ATP5O were significantly increased and PFKM, TKT, and NDUFB10 were significantly decreased.

Irradiation–induced changes in signal transduction protein levels ST proteins that regulate metabolism were
sorted by the PANTHER Classification system (http://www.pantherdb.org/). Their relative quantifications are shown in Figure 5. CUZD1, MAP2K4, and PDGFRB were upregulated. MAP2K4 is a tumor suppressor in many cancer cells and is a component of a stress and cytokine-induced signaling pathway involving MAPK proteins [26,27]. PDGFRB is considered as a cytoplasmic ATM activator that protects some cells from oxidative stress [28].

**Post-irradiation protein network analysis of metabolism and signaling transduction pathways** To further understand cell regulation post-irradiation, IPA was applied to analyze the protein networks of metabolic and signaling transduction processes. For metabolic processes, we found two networks with significant scores, and these are shown in Figure 6. One was "Molecular Transport, Small Molecular Biochemistry, and Cell Cycle," with a highly significant score of 40 (Figure 6A), and the second was "Free Radical Scavenging, Small Molecule Biochemistry, Protein Synthesis," with a score of 28 (Figure 6B).

For signaling transduction, the most significantly dysregulated network was "Cell Death and Survival, Cell Cycle, and Post-Translational Modification," with a score of 42 (Figure 7).

**DISCUSSION**

Our analysis employs targeted proteomics to study metabolism and signal transduction in melanoma cells. Glycolysis-related proteins are elevated in most cancer cells according to the "Warburg effect" [29]. We found that HK1

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**Figure 4** Relative quantification of the energy pathway proteome in response to 10-Gy X-ray treatment of 92–1 melanoma cells. The energy pathway was separated into four sections by dashed lines, to divide the glycolysis, TCA cycle, oxidative phosphorylation, and pentose phosphate pathways. Bars above the top horizontal line represent significantly increased proteins. Bars lower than bottom horizontal line represent significantly decreased proteins.

**Figure 5** Relative quantification of signal transduction proteins involved in metabolism in response to 10-Gy X-ray treatment of 92–1 melanoma cells. Bars above the top horizontal line represent significantly increased proteins. Bars lower than bottom horizontal line represent significantly decreased proteins.

**Figure 6** One was "Molecular Transport, Small Molecular Biochemistry, and Cell Cycle," with a highly significant score of 40 (Figure 6A), and the second was "Free Radical Scavenging, Small Molecule Biochemistry, Protein Synthesis," with a score of 28 (Figure 6B).

For signaling transduction, the most significantly dysregulated network was "Cell Death and Survival, Cell Cycle, and Post-Translational Modification," with a score of 42 (Figure 7).
Figure 6 Schematic representation of proteins associated with metabolism networks using IPA  
A: The network "Molecular Transport, Small Molecular Biochemistry, and Cell Cycle" had a highly significant score of 40;  
B: The network "Free Radical Scavenging, Small Molecule Biochemistry, Protein Synthesis" had a highly significant score of 28. Green and red colors indicate proteins found to be downregulated and upregulated in our study, respectively.

Figure 7 Schematic representation of proteins associated with signal transduction networks using IPA  
The network "Cell Death and Survival, Cell Cycle, and Post-Translational Modification" had a highly significant score of 42. Green and red colors indicate proteins found to be downregulated and upregulated in our study, respectively.
and LDHB were upregulated and PFKM was downregulated. Previous studies reported that HK2, LDHA, and PFKM were direct targets of Myc. However, the significance of LDHB in tumor development remains elusive, and the regulation of its expression is also less characterized. However, recent studies show that mTOR is a positive regulator of LDHB. HK1 is considered as the first step in glycolysis and is a likely candidate for the control of glucose metabolism. Generally, HK1 expression increase was indicative of previous glucose uptake. Recent evidence suggests that HK1 may also have an antiapoptotic role. Indeed, previous experiments showed that irradiated cells do not undergo apoptosis.

CSNK2A2, CSNK2B, DPYSL2, FKB1P1A, PHB, and ST13 were downregulated 15 hours after irradiation CSNK2A2 and CSNK2B are catalytic and regulatory subunits of the protein complex CKII (casein kinase II), respectively. CKII plays an important role in cell cycle progression, and CKII-inactivated cells may arrest in G1 due to upregulation of Sic1 (cyclin-dependent kinase inhibitor). Previous research indicated that p53 may lose the ability to maintain mitotic integrity in CKII absence or dysfunction via a loss of its ability to appropriately regulate cyclin B levels, as well as an inability to execute apoptosis. CSNK2B/CK2β is a regulatory subunit of CKII, which is considered as a vital protein that regulates cell cycle in both G1 and G2/M phases. Yde et al. found that downregulating CK2β by RNA interference resulted in delayed cell cycle progression at the onset of mitosis.

PHB and ST13 are considered to be tumor suppressor proteins. PHB binds to Rb proteins and represses E2F transcriptional activity. The mechanism by which DPYSL2, FKB1P1A, PHB, and ST13 are downregulated following irradiation remains unclear.

Based on the cell phenotypes observed post-irradiation (Figure 3), we inferred that the antiapoptosis pathways might be closely associated with glycolysis enhancement and other key proteins involved in signal transduction.

For metabolic processes, we found that the second network included many key proteins (Figure 6B). Recent research has shown that GLO1 is upregulated in various human malignant tumors, including melanoma. Moreover, GLO1 overexpression has been associated with cancer cell survival and resistance to chemotherapeutic agents. We found that GLO1 was significantly downregulated, and cell cycle arrest was observed at 15 hours. Moreover, antioxidant enzymes including PRDX1, PRDX2, and PRDX3 were also downregulated, although PRDX5 was not. Besides its role as an antioxidant, the effects of PRDX5 upregulation after radiation remain unknown. We inferred that attenuating antioxidant functions in irradiated melanoma cells might induce more serious DNA damage.

With regard to signaling transduction processes, the affected network was closely associated with cell survival and cell cycle. Some important regulators, such as NF-κ B complex, PI3K/Akt, and MAPK/Jnk might play vital roles in cell cycle arrest and cell survival. One limitation of this 2D-LC/MS/MS study is that it was difficult to detect regulator proteins due to their low levels.

In summary, we demonstrated changes in metabolism and signal transduction protein pathways in irradiated melanoma 92-1 cells with SILAC and 2D-LC-MS/MS. We set out to investigate mechanisms of cell cycle arrest, antiapoptosis, and growth inhibition following irradiation and identified several key regulatory proteins in cell cycle and cell survival. We hope that these findings will be useful in finding regulators that can be targeted to improve uveal melanoma radiotherapy.

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