Proteomic and functional analyses demonstrate the involvement of oxidative stress in the anticancer activities of oridonin in HepG2 cells

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Proteomic and functional analyses demonstrate the involvement of oxidative stress in the anticancer activities of oridonin in HepG2 cells

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Abstract. Oridonin exhibits a curative effect on liver carcinoma in patients and experimental animals. In the present study, we performed proteomic and functional analyses to explore the mechanism involved in the anticancer activity of oridonin. Oridonin treatment for 24 h resulted in a dose-dependent decrease in cell viability with an IC50 value of 37.90 µM. Treatment with 40 µM oridonin for 24 h induced apoptosis as typical apoptotic nuclear alterations were observed following DAPI staining. Using a 2-DE-based proteomic approach, 3 upregulated oxidative stress markers, Hsp70-1, Hop and Prdx2, were identified in the HepG2 cells treated with 40 µM oridonin for 24 h. A pattern of alteration in Hsp70-1 was verified by western blotting. The mRNA expression patterns of Hsp70-1 and Hop as determined by qPCR were comparable to their protein expression patterns. Further investigations showed that oridonin treatment for 24 h resulted in reactive oxygen species (ROS) generation, and ROS scavenger N-acetylcysteine (NAC) completely inhibited ROS production and restored cell viability, suggesting that oxidative stress contributed to oridonin-induced HepG2 cell death. Western blot analysis of oxidative pathway-related proteins demonstrated that oridonin treatment increased p-JNK, p-p38 and p-p53, and decreased Bcl-2 protein expression levels, promoted cytochrome c release, decreased mitochondrial membrane potential, and activated caspase-9 and caspase-3. Furthermore, knockdown of Hsp70-1 expression with specific shRNA significantly decreased the viability of the cells treated with oridonin, suggesting a protective role of Hsp70-1 in oridonin-mediated oxidative stress. The results of the present study provide evidence for a link between oxidative stress and oridonin-induced apoptosis in HepG2 cells.

Introduction

Rabdosia rubescens (HemsI.) Harra (Dong Ling Cao in Chinese) is a medicinal herb and has long been used for the treatment of hepatoma in China (1). Oridonin (Fig. 1) is the main active constituent of R. Rubescens. Although laboratory and clinical data have confirmed the inhibitory activities of oridonin in hepatocellular carcinoma (2-5), the molecular mechanisms of action of oridonin have not been fully elucidated. In the present study, we found that expression levels of a number of oxidative stress markers were upregulated in oridonin-treated HepG2 cells using a proteomic approach.

Oxidative stress is a disturbance in the oxidant-antioxidant balance leading to potential cellular damage. The imbalance can result from a lack of antioxidant capacity caused by disturbances in production and distribution, or by an overabundance of reactive oxygen species (ROS) from other factors. ROS are potential carcinogens due to their roles in mutagenesis, tumor promotion and progression (6). It is commonly accepted that cancer cells have an increased ROS steady state level and are likely to be more vulnerable to damage by further ROS insults induced by exogenous agents. Thus, manipulating ROS levels by redox modulation could be a way to selectively kill cancer cells without causing significant toxicity to normal cells (7). A unique anticancer strategy termed ‘oxidation therapy’ has been developed by inducing cytotoxic oyxstess for cancer treatment. Many antitumor agents, such as vinblastine, cisplatin, mitomycin C, doxorubicin, camptothecin, mitomycin, neocarzinostatin exhibit antitumor activity via ROS-dependent activation of apoptotic cell death (8, 9). To explore the role of oxidative stress in the anticancer activities of oridonin in HepG2 cells, we further performed functional analysis. Results showed that ROS production was responsible for oridonin-induced HepG2 cell death.

Materials and methods

Reagents. Oridonin was purchased from Shanghai Shamrock Imp. & Exp. Trading Co., Ltd. (Shanghai, China). Its purity was determined to be 97% by HPLC. Immobilized pH gradient (IPG) strips, dithiothreitol (DTT), the 2D Clean-Up
Kit and the 2D Quant Kit were obtained from GE Healthcare. MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] was from USB Corporation. Trypsin was from Promega Corporation (Madison, WI, USA). Silver nitrate, Rhodamine 123 and 4,6-diamidino-2-phenylindole (DAPI) were from Sigma-Aldrich Biotechnology. Power SYBR-Green PCR Master Mixture was obtained from Applied Biosystems. Protease inhibitor cocktail, phosphatase inhibitor cocktail and FuGENE HD transfection reagent were from Roche Bioscience. The mitochondria isolation kit was from Beijing Applygen Technologies Inc. Bio-Rad protein assay was from Bio-Rad Laboratories, Inc. Iodoacetamide (IAA), nitrucellose membranes and ECL detection reagents were purchased from Amersham Biosciences. p-JNK (Thr183/Tyr185), p-p38 (Thr180/Tyr182), p-p53 (Ser15), cytochrome c and cleaved caspase-3 antibodies were from Cell Signaling Technology. Hsp70, Bcl-2, caspase-3, caspase-9, β-actin and anti-rabbit IgG antibodies (horseradish peroxidase-conjugated) as well as the Hsp70 shRNA and control shRNA were from Santa Cruz Biotechnology (Santa Cruz, CA, USA). The SuperScript II reverse transcription kit, TRizol reagent, CM-H2DCFDA and the anti-mouse IgG antibody (horseradish peroxidase conjugated) were obtained from Invitrogen Biotechnology.

**Cell culture.** HepG2 cells (ATCC, Manassas, VA, USA), grown in Dulbecco's modified Eagle's medium (DMEM; Gibco, Carlsbad CA, USA) supplemented with 10% fetal bovine serum (FBS; Gibco) and 1% penicillin/streptomycin (P/S, Gibco) were cultured at 37˚C in an atmosphere containing 5% CO₂.

**Determination of cell viability.** Cell viability was assessed by the MTT assay. HepG2 cells (5.0x10⁵/well) were seeded and grown in 96-well plates for 24 h and then treated with various concentrations of oridonin. Control cells were treated with 0.1% DMSO. After a 24-h incubation, 10 µl of 5 mg/ml MTT solution was added to each well, and the plates were incubated at 37°C for 4 h. Following medium removal, 100 µl of DMSO was added to each well and the plates were gently shaken for 5 min. Optical absorbance was determined at 570 nm with a microplate spectrophotometer (BD Biosciences, Franklin Lakes, NJ, USA). The absorbance obtained by vehicle-treated cells was considered to represent 100% cell survival. Each treatment was performed in triplicate and each experiment was repeated six times.

**Fluorescence microscopic analysis of apoptosis.** Apoptotic morphological alterations were monitored in DAPI-stained cells. Cells (40x10⁴) were grown for 24 h on coverslips in 35-mm dishes in the presence or absence of 40 µM oridonin. Coverslips were carefully washed with PBS, fixed with 4% paraformaldehyde for 10 min and incubated with 10 µg/ml DAPI for 10 min. Cells were washed with PBS and observed under a fluorescence microscope (Nikon, Tokyo, Japan).

**Proteomic sample preparation and 2-DE.** Cells were seeded in 100-mm culture dishes at 1.5x10⁶ cells/dish, incubated overnight and then treated with 40 µM oridonin for 24 h. Cells were harvested by trypsinization, washed three times with isotonic buffer (10 mM Tris, 250 mM sucrose, pH 7.2) and lysed in a lysis buffer (8 M urea, 4% CHAPS, 40 mM DTT and 0.5% IPG buffer pH 3.0-10.0 NL) by gentle shaking for 60 min on ice. Extracts were centrifuged at 25,000 x g for 60 min at 4°C. Supernatants were purified with the 2D Clean-Up Kit following the manufacturer's instructions. The protein concentration of each purified sample was determined using the 2D Quant Kit, and protein samples were stored in aliquots at -80°C until further analysis.

First-dimension separation was performed using 13-cm IPG strips (pH 3.0-10.0 NL). Samples were diluted in rehydration solution containing 8 M urea, 2% CHAPS, 0.5% IPG buffer, 0.002% bromophenol blue and 0.28% DTT to reach a final protein load of 100 µg (in 250 µl) per strip. IPG strips were actively rehydrated at 30 V for 12 h, focused at 500 V for 2.5 h, 1,000 V for 0.5 h, and then the voltage was increased to 8,000 V gradually over the next 3 h and maintained at 8,000 V for 40,000 Vh. Prior to the second-dimension separation, IPG strips were equilibrated in equilibration buffer (6 M urea, 75 mM Tris-HCl pH 8.8, 29.3% glycerol, 2% SDS) containing 1% (w/v) DTT for 15 min and then in the same equilibration buffer containing 2.5% IAA for a further 15 min. The second-dimension separation was carried out on 10% SDS-PAGE (20 mA/gel, 10 min; 25 mA/gel, 260 min).

**Silver staining and image analysis.** Protein spots were visualized by silver staining. Gels were fixed overnight in fixing solution (40% v/v methanol and 10% v/v acetic acid) and then treated for 30 min with sensitizing solution (30% ethanol, 4.05% w/v sodium acetate and 0.2% sodium thiosulfate). Each gel was washed three times with Milli-Q water for 5 min each, stained for 40 min in 0.1% silver nitrate solution and then developed by incubation with a developing solution (2.5% w/v sodium carbonate and 0.02% formaldehyde) until protein spots appeared. The developing reaction was terminated by putting gels in the stopping solution (1.46% w/v EDTA, disodium salt) for 10 min, and finally gels were washed three times with Milli-Q water for 5 min each.

The stained gels were scanned with a LabScan 6.0 software installed image scanner (GE Healthcare, Pittsburgh, PA, USA), and data were analyzed using the ImageMaster 2D Platinum 6.0 software (GE Healthcare). The intensity volume of each spot was processed by background subtraction and total spot volume normalization, and the resulting spot volume percentage was used for comparison. Only those significantly (Student's t-test; P<0.05) and consistently upregulated or downregulated spots (>2 fold) or spots which appeared or disappeared after treatment in three independent experiments were selected for in-gel digestion and MALDI-TOF-MS/MS analysis.
In-gel digestion and MALDI-TOF-MS/MS analysis. Protein spots were excised from gels, and each sample was transferred to a 1.5-ml Eppendorf tube and de-stained in a freshly prepared de-staining solution (15 mM potassium ferricyanate and 50 mM sodium thiosulfate) until the brownish color disappeared. Each de-stained sample was washed in 10 mM ammonium bicarbonate (NH₄HCO₃) for 5 min, 10 mM NH₄HCO₃ containing 10 mM DTT for 15 min at 56°C, 10 mM NH₄HCO₃ containing 55 mM IAA for 20 min at room temperature in the dark, and 10 mM NH₄HCO₃ containing 50% ACN for 15 min. After drying in a Vacuum concentrator (Eppendorf, Hamburg, Germany), each sample was incubated at 37°C overnight in 5 µl of 5 µg/ml Trypsin Gold solution. The supernatant was collected and the gel was further extracted with 1% formic acid. The extracts were combined and dried in a Vacuum concentrator and resuspended in formic acid for MS analysis. The mass spectra were obtained using the Bruker Autoflex III MALDI-TOF/TOF mass spectrometer (Bruker Daltonics, Billerica, MA, USA). Protein identification was performed automatically by searching the Swiss-Prot 55.3 database using the Mascot 2.04 search engine (Matrix Science, Ltd., London, UK). Database searches were carried out using the following parameters: type of search, MS/MS ion search; enzyme, trypsin; and allowance of one missed cleavage. Carbamidomethylation was selected as a fixed modification and oxidation of methionine was allowed to be variable. The peptide and fragment mass tolerance were set at 50 ppm and 0.5 Da, respectively. The instrument was selected as MALDI-TOF-TOF. Proteins with probability-based MOWSE scores (P<0.05) were considered to be positively identified.

Western blot analysis. Cells were collected and proteins were extracted with RIPA lysis buffer [50 mM Tris-Cl, 1% v/v NP-40, 0.35% w/v sodium deoxycholate, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, pH 7.4, 1 mM phenylmethylsulfonyl fluoride (PMSF), 1 mM NaF, 1 mM Na₂VO₄] containing a protease inhibitor cocktail and phosphatase inhibitor cocktail. Mitochondrial and cytoplasmic extracts were prepared with a mitochondria isolation kit following the manufacturer's instructions. The protein concentration was determined using the Bio-Rad protein assay. Individual protein samples (20-50 µg) were separated by SDS-PAGE and then electro-transferred onto nitrocellulose membranes. The membranes were blocked for 30 min with 3% skim milk in TBST buffer composed of 50 mM Tris (pH 7.6), 150 mM NaCl and 0.1% Tween-20 and incubated with primary antibodies overnight at 4°C followed by incubation with horseradish peroxidase-conjugated secondary antibodies. Protein signals were visualized by ECL detection reagents according to the manufacturer's instructions.

qPCR. Cells were treated with 40 µM oridonin for the indicated durations, and total RNA was isolated using TRIzol reagent according to the manufacturer's protocol. Five micrograms of RNA was used for reverse transcription by oligo-dT using the SuperScript II reverse transcription kit. The qPCR primers were designed as follows: Hsp70-1 (sense 5’-caga acaagggcgctggagg-3’ and antisense 5’-tgagctggtggtggctg tcc-3’), Hop (sense 5-tgaaggagccgcagccgagc-3’ and antisense 5’-taagge gcatggtgggtgca-3’). To normalize the amounts of RNA in samples, a PCR reaction was also performed with primers of β-actin (sense 5’-gactatcctggaagtc-3’ and antisense 5’-gtctccagggcctggaagc-3’). qPCR was performed in a total volume of 20 µl, with 1X Power SYBR-Green PCR Master Mix in the 7500 Fast Real-time PCR system (Applied Biosystems, Foster City, CA, USA). All samples were run in triplicate. The cycling parameters for qPCR reaction included 40 cycles of 95°C for 15 sec and 60°C for 60 sec. The amplification of specific transcripts was confirmed by melting curve profiles generated at the end of the PCR program. Cycle threshold (Ct) values were normalized to β-actin, and comparative quantification of the target gene was carried out using the ΔΔCT method.

Transient transfection of Hsp70 shRNA. Transient HSPA1 gene silencing was attained by transfection of Hsp70 shRNA into HepG2 cells using FuGENE HD transfection reagent according to the manufacturer's instruction. HepG2 cells were grown in 6-well plates overnight to achieve >80% confluency at the time of transfection and then were transfected with 4 µg of Hsp70 shRNA or control shRNA and 6 µl FuGENE HD transfection reagent. The transfection reagents/shRNA complex was diluted with DMEM (FBS and antibiotic-free) to a total transfection volume of 2 ml and incubation was carried out for 6 h at 37°C until the medium was discarded and replaced with growth medium (DMEM with 10% FBS). Cells were incubated at 37°C for a further 30 h and then subjected to 40 µM oridonin treatment for 12 h. The cells were then collected for verification of protein expression by western blotting. Cell viability was determined by MTT assay.

Measurement of intracellular ROS production. The dye, CM-H₂DCFDA, which becomes fluorescent after cellular oxidation, was used for ROS detection. HepG2 cells were treated with 40 µM oridonin in the absence or presence of 2.5 mM NAC, an ROS scavenger. After 24 h, cells were harvested and washed with PBS. The washed cells were resuspended in PBS containing 10 µM CM-H₂DCFDA and incubated at 37°C in the dark for 30 min. The levels of ROS were then determined by flow cytometry.

Observation of mitochondrial membrane potential (MMP). MMP was determined by the fluorescent dye Rhodamine 123. After incubation with 40 µM oridonin for 6 h, cells were collected, resuspended in 1 µM Rhodamine 123 and stained at 37°C for 15 min. The fluorescence intensity of the cells was analyzed by flow cytometry.

Data analysis. All results are expressed as the mean ± SEM, and statistical analyses were performed using the Student's t-test.

Results

Oridonin inhibits cell proliferation and induces apoptosis in HepG2 cells. MTT assay showed that treatment with oridonin for 24 h resulted in a marked decrease in cell viability in a dose-dependent manner (Fig. 2A). The IC₅₀ value was determined as 37.90 µM for the 24-h treatment from a dose-response curve with GraphPad Prism 5.0 software (GraphPad Software, San Francisco, CA), which indicated a statistically significant difference compared to the control group (p<0.05). Additionally, the cell cycle was analyzed by flow cytometry. The results showed that oridonin treatment led to a significant increase in the percentage of cells in the S phase (Fig. 2B). These results suggest that oridonin may have a potential therapeutic effect on HepG2 cells.
Table I. Summary of differentially expressed proteins in the oridonin-treated HepG2 cells.

<table>
<thead>
<tr>
<th>Protein names</th>
<th>MW</th>
<th>pI</th>
<th>MS/MS scores</th>
<th>Fold-change*</th>
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</thead>
<tbody>
<tr>
<td>Heat shock 70-kDa protein 1</td>
<td>70,294</td>
<td>5.48</td>
<td>212</td>
<td>+2.08</td>
</tr>
<tr>
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<td>63,227</td>
<td>6.40</td>
<td>88</td>
<td>4*</td>
</tr>
<tr>
<td>Peroxiredoxin-2</td>
<td>22,049</td>
<td>5.66</td>
<td>192</td>
<td>+2.45</td>
</tr>
</tbody>
</table>

MW, molecular weight; pI, isoelectric point; MS, mass spectrometry; *Fold-change in the oridonin-treated HepG2 cells at the time-point of 24 h compared with the control (+, increase; -, decrease). *Protein expression was not detected in the control samples.

Differentially expressed proteins in the oridonin-treated HepG2 cells. In an attempt to identify molecular changes following oridonin treatment, we monitored differential protein expression in the HepG2 cells treated with 40 µM oridonin or vehicle for 24 h using 2-DE-based proteomics. To ascertain reproducibility of results, 2-DE was performed three times for each protein sample, and each treatment was repeated three times. Fig. 3A shows the representative gel images. Proteins within the range of 15-225 kDa and with isoelectric point (pI) between 3 to 10 were well separated. Over 1,000 spots were detected on each gel. Three protein spots of interest with fold-changes ≥2 in terms of volume intensity in both triplicate gels and three independent experiments were cut from the gels and analyzed by MALDI-TOF-MS/MS after trypsin digestion. As listed in Table I, these upregulated spots were identified as heat shock 70-kDa protein 1 (Hsp70-1, gene name HSPA1), Hsc70/Hsp90-organizing protein (Hop/Sti1, gene name STIPI) and peroxiredoxin-2 (Prdx2, gene name PRDX2).

The protein expression level of Hsp70-1 in response to 40 µM oridonin treatment was verified by western blotting (Fig. 3B). Expression levels of mRNA as determined by qPCR
Oridonin-induced ROS production is responsible for cell death in HepG2 cells.

Hsp70-1 and Hop were 2.64- and 1.68-fold higher than these levels in the control cells (Fig. 3C), which are comparable to their protein expression patterns.

**Oridonin-induced ROS production is responsible for cell death in HepG2 cells.** Hsp70 acts as a stress sensor and its expression is induced by a variety of stimuli that give rise to cellular oxidation (10). Hop is an extensively studied co-chaperone that not only directly binds to Hsp70 and Hsp90, but also modulates the activities of these chaperones. Overexpression of the Hop gene at the earliest time during stress may be required to mediate the stress response of the inducible Hsp70 (11). Prdx2 belongs to the 2-Cys subgroup of peroxiredoxins (Prdxs).

Prdxs are a family of antioxidative proteins and in mammals the family consists of six members (12,13). They have been reported to be increased in stressed cells induced by oxidants such as H$_2$O$_2$ (14-16) or ionizing radiation (17). Taken together, the overexpression of Hsp70-1, Hop and Prdx2 indicates that oridonin induces oxidative stress.

To investigate whether oridonin induces oxidative stress in HepG2 cells, the intracellular ROS level was measured by detecting the fluorescent intensity of ROS-specific probe CM-H$_2$DCFDA after a 40 µM oridonin treatment. As shown in Fig. 4A, the ROS level increased at 24 h, and 2.5 mM of the ROS scavenger NAC markedly decreased ROS production in the oridonin-treated HepG2 cells. To explore whether ROS act

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**Figure 4. Role of ROS in oridonin-induced cell death.** (A) Flow cytometric analysis of ROS production. HepG2 cells were stained with ROS-specific probe CM-H$_2$DCFDA after the indicated treatments for 24 h. The intracellular fluorescence was determined by flow cytometry. Results are representative of three independent experiments. (B) Representative micrographs of HepG2 cells after the indicated treatments for 24 h. Micrographs were examined by phase-contrast microscopy (original magnification, x100). (C) Effects of NAC on the viability of HepG2 cells treated with 40 µM oridonin for 24 h. Cell viability was determined by MTT assay. Data are presented as the mean ± SEM of three independent experiments. *P<0.01 vs. the control; **P<0.01 vs. 40 µM oridonin treatment.
as a protective factor in the oridonin-treated HepG2 cells, cells were co-incubated with 2.5 mM NAC and 40 μM oridonin for 24 h. NAC completely inhibited oridonin-induced cell death (Fig. 4B). Cell viability was 104.50±4.31% in the NAC and oridonin-treated cells vs. 60.49±2.89% in the oridonin-treated cells (Fig. 4C). These results suggest that ROS generation plays a role in the antiproliferative activity of oridonin in HepG2 cells.

Oridonin-induced ROS generation results in apoptosis through the activation of JNK and p38 pathways. Oxidative stress resulting from excessive production of ROS triggers apoptotic cell death through p38 and JNK pathways (18). JNK and p38 function as upstream kinases of p53 phosphorylation and subsequently stabilize and activate p53 transcriptional activity, leading to diverse cellular responses, such as apoptosis (19). Bcl-2 and caspase families are required to mediate p53-dependent apoptosis (20).

To determine whether p38 and JNK pathways are involved in the anticancer effects of oridonin in HepG2 cells, immunoblotting was conducted. As shown in Fig. 5A and B, oridonin treatment significantly increased phosphorylated p38 (p-p38), p-JNK, and p38, and decreased the expression of Bcl-2, mitochondrial cytochrome c, caspase-9 (46 kDa) and caspase-3, increased the expression of cytoplasmic cytochrome c, as well as activated the cleavage of caspase-9 (35 kDa) and caspase-3. The integrity of the mitochondrial membranes was examined by flow cytometry in Rhodamine 123-stained cells. Results showed that treatment with 40 μM oridonin for 6 h significantly decreased Rhodamine 123 fluorescence intensity (Fig. 5C). This observation suggests that oridonin treatment decreased MMP in HepG2 cells.

Hsp70-1 knockdown increases the antiproliferative effect of oridonin. To investigate the involvement of Hsp70-1 in the apoptotic effect of oridonin, HepG2 cells were transfected with Hsp70 shRNA. Control cells were transfected with the control shRNA plasmid. Oridonin treatment caused Hsp70-1 induction and transient transfection of Hsp70-specific shRNA suppressed Hsp70-1 expression with or without 12-h oridonin treatment (Fig. 6A). Moreover, Hsp70 shRNA transfection, oridonin treatment or the combination of the two treatments significantly reduced cell viability (Fig. 6B and C) when compared with the control cells. These data confirm that Hsp70-1 protects HepG2 cells against cell death. Furthermore, Hsp70-1 knockdown led to the increased sensitivity of HepG2 cells to oridonin cytotoxicity, which suggests that Hsp70-1 acts as a protective factor against oridonin-induced oxidative stress in HepG2 cells.

Discussion

Oridonin, one of the main bioactive ent-kaurane diterpenoids of R. rubescens, possesses anticancer activities. Oridonin can induce G2/M arrest and apoptosis in human hepatocellular carcinoma cells (21,22). Multiple mechanisms such as inhibition of telomerase, regulation of MAPK signaling pathways and induction of endoplasmic reticulum stress have been implicated in the anticancer activity of oridonin (4,21,23,24). In the present study, using a 2-DE-based proteomic approach, we found that the expression levels of oxidative stress markers Prdx2, Hsp70-1 and Hop were altered following oridonin treatment.

Free radicals and reactive molecules containing oxygen such as singlet oxygen, superoxides, peroxides are known as ROS and can cause oxidative stress in cells (18). It has been suggested that oxidative stress in response to a low concentration of ROS may be essential to cellular signaling and proliferation. However, excessive production of ROS may damage various intracellular molecules, leading to severe oxidative stress accompanied by loss of cell function and possibly apoptosis (25). Many studies have shown that apoptosis is closely associated with excessive production of ROS (26-28). Since our proteomic analysis identified several
proteins related to oxidative stress, we investigated whether oxidative stress occurs in oridonin-treated HepG2 cells and explored the role of oxidative stress in the anticaner activities of oridonin in HepG2 cells. We detected the level of reactive oxygen species (ROS) following oridonin treatment for 24 h and found that oridonin promoted ROS generation, and ROS scavenger N-acetylcysteine (NAC) completely inhibited ROS production and restored cell viability, suggesting that ROS generation contributed to oridonin-induced HepG2 cell death. Although a low dose of oridonin (1.4 µM) is reported to decrease ROS formation and improve cell survival after arsenic treatment (29), therapeutic doses of oridonin are shown to selectively induce apoptosis in various cells through sustained oxidative stress (30-33).

Among many signaling pathways known to be redox sensitive, the MAPK pathway has been well studied. A relative high concentration of ROS may induce activation of JNK and p38 MAPK pathways and contribute to apoptosis (25,34). MAPKs have been shown to phosphorylate p53 at various residues in response to different stressful stimuli, and such phosphorylation can initiate p53 activation and lead to p53-dependent apoptosis (19). Activation of p53 suppresses the expression of anti-apoptotic Bcl-2 protein which prevents the release of mitochondrial cytochrome c. Through inhibition of Bcl-2, p53 promotes the release of cytochrome c from the mitochondrial intermembrane space into the cytoplasm (20). Cytochrome c release is frequently coincident with a disruption of the mitochondrial membrane potential (MMP). Released cytochrome c forms an apoptosome complex with caspase-9 and adaptor protein Apaf-1. Caspase-9 is activated following recruitment into the apoptosome and then cleaves and activates effector caspases, such as caspase-3 (35). The cleavage of substrates of caspase-3 significantly alters the cell physiology toward apoptosis (20). We next investigated the involvement of these molecules in oridonin-induced apoptosis by immunoblotting. In oridonin-treated cells, increased expression of p-p38 and p-JNK was accompanied by the upregulation of phosphorylated p53 at serine 5. The activation of p53 decreased the expression of Bcl-2, which triggered mitochondrial cytochrome c release, thus suppressing MMP and activating caspase cascades in HepG2 cells. In agreement with our observations, activation of p38 was reported to be dependent on ROS generation, and p38 and p53 inhibitors have been reported to significantly reduce oridonin-induced apoptosis in HepG2 cells (32).

Abnormal changes in the cellular redox status cause chaperone induction. Chaperones may act as protective factors against cytoplasmic oxidative stress (10). Since upregulation of oxidative stress marker Hsp70-1 was associated with oridonin
treatment in HepG2 cells, we investigated the role of Hsp70-1 in the antiproliferative effect of oridonin by Hsp70-1 shRNA plasmid transfection. Hsp70-1 knockdown enhanced the sensitivity of HepG2 cells to oridonin-induced cell death. These findings suggest that Hsp70-1 acts as a protective factor against oridonin-induced oxidative stress in HepG2 cells, and also demonstrates the induction of oxidative stress by oridonin.

Collectively, proteome analysis revealed that oxidative stress markers Prdx2, Hsp70-1 and Sirt1 are involved in the anticancer effects of oridonin. Oxidative stress pathways are activated in oridonin-treated HepG2 cells. Hsp70-1 acts as a protective factor against oxidative stress in HepG2 cells. These results shed new light on the anti-hepatoma mechanisms of oridonin. Further research is needed to explore the mechanisms by which ROS mediate the apoptotic activity of oridonin.

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