Identification of over-expressed proteins in oral squamous cell carcinoma (OSCC) patients by clinical proteomic analysis

Wan-Yu Lo a, Ming-Hsui Tsai b, Yuhsin Tsai c, Chun-Hung Hua b, Fuu-Jen Tsai a, Shiuan-Yi Huang a, Chang-Hai Tsai d, Chien-Chen Lai a,c,⁎

a Department of Medical Research, China Medical University Hospital, Taichung, Taiwan
b Department of Otorhinolaryngology, China Medical University Hospital, Taichung, Taiwan
c Graduate Institute of Chinese Medical Science, China Medical University, Taichung, Taiwan
d Department of Biotechnology and Bioinformatics, Asia University, Taichung, Taiwan
e Institute of Molecular Biology, National Chung Hsing University, Taichung, Taiwan

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Abstract

Background: Oral cancer is a worldwide problem. It is a universal aggressive disease in the population of smoking and drinking. The oral cancer mortality has been ranked 5th place in Taiwan in male cancer patients. A number of protein markers for oral cancer are still not applicable in large populations. Proteomic technologies provide excellent tools for rapid screening of a large number of potential biomarkers in malignant cells.

Method: Proteomics and real-time quantitative RT-PCR were used to analyze over-expressed proteins in 10 OSCC patients.

Result: Forty-one proteins were identified as commonly over-expressed in OSCC tissues. In OSCC tissues, αB-crystallin, tropomyosin 2, myosin light chain 1, heat shock protein 27 (HSP27), stratifin, thioredoxin-dependent peroxide reductase, flavin reductase, vimentin, rho GDP-dissociation inhibitor 2 (rho GDI-2), glutathione S-transferase Pi (GST-pi) and superoxide dismutase [Mn] (MnSOD) were significantly over-expressed (an average of 7.2, 6.0, 5.7, 4.3, 3.6, 3.4, 3.0, 3.0, 2.6, 2.5, 2.1-fold, respectively). In real-time quantitative RT-PCR analysis, the gene expressions of αB-crystallin, HSP27 and MnSOD were also increased in the cancer tissues, consistent with proteomic results.

Conclusion: The identified proteins in this experiment may be used in future studies of carcinogenesis or as diagnostic markers and therapeutic targets for OSCC.

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Keywords: Oral squamous cell carcinoma (OSCC); Oral cancer; Proteomics; LC-MS/MS; Real-time quantitative RT-PCR

1. Introduction

Oral cancer is one of the most common types of human cancer in the world [1]. Among them, oral squamous cell carcinoma (OSCC) occurs with the highest frequency. Nevertheless, factors deciding on its clinical characteristics including carcinogenesis, development, progression, invasion, and metastasis have not been elucidated yet. This cancer is a highly variable disease with multiple heterogeneous genetic and epigenetic changes associated with various types of clinical behavior. Globally about 500,000 new oral and pharyngeal cancers are diagnosed annually, and three quarters of these are from the developing world [2]. In Taiwan, the incidence of OSCC is one of the highest in the world, 20 per 1,000,000 among men, and comprising approximately 4–5% of all malignancies [3].

The mortality rate of oral cancer increased about 5.83-fold from 1981 to 2003 in Taiwan, making its prevention and early detection and treatment an important public health issue [4]. Generally, both smoking and drinking are the important risk factors for cancer of the oral cavity, oropharynx, hypopharynx, and larynx. Other risk factors for oral cancer include: human papillomavirus (HPV) infection, Epstein-Barr virus infection,
occupational exposure to wood dust and consumption of certain preservatives or salted foods [5,6]. It is different from other regions; chewing areca quid is the most important risk factor to induce oral cancer in Taiwan. Approximately 85% of all patients with OSCC habitually use betel quid [7]. Following previous reports, the 5-year survival rates for mouth, tongue, oropharynx, and laryngopharyngeal caners seldom exceed 40%. Therefore, the existence of a reliable, accurate, cost-effective, and noninvasive test for OSCC is desirable [8].

The alteration of genes has been traditionally revealed by the use of cytogenetics, immunohistochemistry, or molecular approaches based on one or a few genes that changed the expression of many genes such as oncogenes and tumor suppressor genes which have been associated with oral carcinogenesis [9–11]. But the molecular signals triggering the onset of oral carcinogenesis are still unknown. Genomics has been incorporated in oncology research and now, in the postgenomic era, there is a strong drive to incorporate proteomic technologies as well [12,13]. Proteomics is a promising approach in the identification of proteins with changed levels which may be useful as diagnostic markers for the early detection of cancer [14,8]. Proteomics have been successfully employed in studies of lung [15], breast [16], prostate [17], gastrointestinal cancer [18], tongue squamous cell carcinoma (SCC) [19] and head and neck squamous cell carcinoma (HNSCC) [20].

In this study, we combined the proteomics and real-time quantitative RT-PCR to approach the characterization of significantly over-expressed proteins from OSCC patients. We designed a rapid and convenient system to hunt all possible diagnostic markers from OSCC patients in a short term. We globally surveyed differential gene expression using matched normal and tumor tissues from patients with OSCC to identify genes that may predispose to the cancer.

2. Materials and methods

2.1. Patients and tissues

The pairs of surgical archived specimens of primary OSCC and their matched adjacent normal surrounding mucosa specimens from 10 patients were obtained from the China Medical University Hospital (Taichung, Taiwan). The patients were between 40 and 60 years old. Patient information about sex, age, and histories of cigarette smoking, betel quid chewing, and medical illness was obtained from medical charts or by telephone inquiry. The American Joint Committee on Cancer pathologic stage of ten patients was T3 (9 N0, 1 N2). The stage of the cancer was defined by the extent of the lesion and was pathologically defined for all ten cases. Consistently and significantly 2-fold) were averaged in each protein and normalized in tumor samples for the advance analysis. Comparisons were made between gel images of individual tumor and normal samples from 10 patients, we screened the over-expressed proteins (>2-fold) in the same equilibrium solution containing 50 mmol/l Tris–HCl (pH 8.8), 6 mol/l urea, 30% glycerol, 2% SDS, a trace of bromophenol blue, and DTE (1% w/v) for 20 min, and followed with second equilibration for 20 min in the same equilibrium solution containing iodoacetamide (2.5% w/v) instead of DTE. Finally, the strip was transferred to the top of 12% polyacrylamide gels and held in position with molten 0.5% agarose in running buffer containing 25 mmol/l Tris, 0.192 mol/l glycine and 0.1% SDS. Gels were run at 16 mA/gel for 30 min followed by 50 mA/gel for 4–5 h.

2.2. Protein extractions

For two-dimensional gel electrophoresis, to remove blood by PBS, all tissues were minced into pieces of about 1–2 mm³ and then homogenized with 0.5 ml lysis buffer (8 mol/l urea, 4% CHAPS), containing protease inhibitor cocktail (Sigma, St. Louis, MO), by a homogenizer (MagNA Lyser; Roche) for 60 s at 6500 rpm. The lysates were centrifuged at 8000 rpm for 20 min at 4 °C and supernatants were collected for acetone precipitation for protein purification. The protein pellet was denatured with sample buffer (8 mol/l urea, 4% CHAPS, 65 mmol/l DTE, 0.5% ampholytes). The denatured proteins were incubated at 4 °C 2–3 h before centrifugation at 13,000 rpm for 15 min. The protein concentrations of the resulting supernatants were measured using the Bio-Rad Protein Assay (Bio-Rad, Hercules, CA) according to the NenoDrop instruction.

2.3. Two-dimensional gel electrophoresis

We took 200 µg protein extraction per sample for 2D gel separation. The extracted sample was diluted to 350 µl with rehydration buffer consisted of an 8 mol/l urea, 4% CHAPS, 65 mmol/l DTE, 0.5% ampholytes, and a trace of bromophenol blue. Then, the rehydration solution (containing the sample) was applied into 17 cm immobilized pH gradient pH 3–10 IPG strip (ReadyStrip IPG strip, Bio-Rad) overnight. Electrophoresis in the first dimension was focused for a total of 60 kV h (Protein IEF cell, Bio-Rad) at 20 °C and then stored at −20 °C until SDS-PAGE. Before SDS-PAGE, IPG strips were equilibrated with 3 ml of an equilibrium solution containing 50 mmol/l Tris–HCl (pH 8.8), 6 mol/l urea, 30% glycerol, 2% SDS, a trace of bromophenol blue, and DTE (1% w/v) for 20 min, and followed with second equilibration for 20 min in the same equilibrium solution containing iodoacetamide (2.5% w/v) instead of DTE. Finally, the strip was transferred to the top of 12% polyacrylamide gels and held in position with molten 0.5% agarose in running buffer containing 25 mmol/l Tris, 0.192 mol/l glycine and 0.1% SDS. Gels were run at 16 mA/gel for 30 min followed by 50 mA/gel for 4–5 h.

2.4. Detection of protein spots and data analysis

Gels were routinely stained with silver nitrate and then scanned by GS-800 imaging Densitometer with PDQuest software ver. 7.1.1 (Bio-Rad). Protein spots were quantified using the PDQuest software. To compare the tumor and normal samples from 10 patients, we screened the over-expressed proteins (>2-fold) in tumor samples for the advance analysis. Comparisons were made between gel images of individual tumor and normal samples by PDQuest software. Normalized volume differences were statistically calculated for all ten cases. Consistently and significantly over-expressed spots (>2-fold) were averaged in each protein and selected for analysis with nanoLC-MS/MS.

2.5. Enzyme digestion

The in-gel digestion method was a modification of Gharahdaghi et al.’s [21] and Terry et al.’s [22]. Each protein spot (1–2 mm
diameter) of interest was cut out with a pipette tip and transferred into a microcentrifuge tube (0.6 ml). The gel pieces were washed twice with 50 μl of 50% acetonitrile (ACN):50% 200 mmol/l ammonium bicarbonate for 5 min, shrank with 100% acetonitrile until the gels turned white, then dried for 5 min in a speed vacuum at room temperature. The gel pieces were rehydrated in 15 μl of 50 mmol/l ammonium bicarbonate (37 °C, 4 min), and an equivalent volume (15 μl) of trypsin solution (20 ng/μl in 50 mmol/l ammonium bicarbonate; Promega, Madison, WI) was added, then it was incubated at 37 °C for 4 h or 30 °C for at least 16 h. After digestion, vortex and spin down of the gel pieces, the supernatant was the peptide solution and was stored at −20 °C until mass analysis.

2.6. Nanoelectrospray mass spectrometry

Nanoscale capillary LC-MS/MS was used to analyze the meaningful proteins involved in the reaction. LC-MS/MS analysis was performed using an Ultimate capillary LC system (LC Packings, Amsterdam, The Netherlands) coupled to a QSTARXL quadrupole-time of flight (Q-TOF) mass spectrometer (Applied Biosystem/MDS Scienx, Foster City, CA). The nanoscale capillary LC separation was performed on an RP C18 column (15 cm × 75 m i.d.) with a flow rate of 200 nl/min and a 60 min linear gradient of 5–50% buffer B. Buffer A contained 0.1% formic acid in 5% aqueous ACN; buffer B contained 0.1% formic acid in 95% aqueous ACN. The nanoLC tip used for online LC-MS was a PicoTip (FS360-20-10-D-20; New Objective, Cambridge, MA). Data acquisitions were performed by Automatic Information Dependent Acquisition (IDA; Applied Biosystem/MDS Scienx). The IDA automatically finds the most intense ions in a TOF MS spectrum, and then performs an optimized MS/MS analysis on the selected ions. The product ion spectra generated by nanoLC-MS/MS were searched against NCBI databases for exact matches using the ProID program (Applied Biosystem/MDS Scienx) and the MASCOT search program (http://www.matrixscience.com) [23]. A homo sapiens taxonomy restriction was used and the mass tolerance of both precursor ion and fragment ions was set to ±0.3 Da. Carbamidomethyl cysteine was set as a fixed modification, while serine, threonine, tyrosine phosphorylation and other modifications were set as variable modifications. All phosphopeptides identified were confirmed by manual interpretation of the spectra.

2.7. Validation of proteomic data by real-time quantitative RT-PCR

To define the relative expression of the genes, the PCR product from each tumor sample was compared with the grossly normal tissue from the same patient. A comparison real-time quantitative RT-PCR method was used to assay the relative expression of the genes in the normal and tumor tissues as described previously [24]. Total RNA was extracted using the TRIzol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. All PCR reactions were performed using the real-time fluorescence detection method using the LightCycler System (Roche Diagnostics, Mannheim, Germany) with a FirstStart DNA Master SYBR Green I kit (Roche Molecular Biochemicals, Indianapolis, IN). The LightCycler System was used for quantitative assessment of gene expression. The primers used for real-time quantitative RT-PCR are listed in Table 1. A reaction mixture containing the following components at the indicated end-concentration was prepared according to the manufacturer’s instructions: 9.4 μl water, 1.6 μl MgCl2 (2.5 mmol/l), 1 μl of forward primer (20 pmol), 1 μl of reverse primer (20 pmol), and 2 μl FastStart DNA Master SYBR Green I mix. Fifteen microliters of the reaction mixture was distributed into precooled capillaries and 20 ng of reverse-transcribed total RNA in a volume of 5 μl was added as PCR template. The GAPDH gene was chosen for normalization of the data. A negative control without cDNA template was performed to assess the overall specificity. To prepare the standard curve, 1 μg of RNA from oral normal tissue was reverse-transcribed and dilutions of 200, 20, 2, and 0.2 ng of cDNA were made. PCR cycle conditions were the following: initial denaturation was for 10 min at 95 °C followed by 45 cycles of amplification at 95 °C (10 s) for denaturation, 56–68 °C (10 s) for annealing, and 72 °C for extension, with a temperature slope of 20 °C/s. After amplification, the temperature was slowly elevated above the melting temperature of the PCR product to measure the fluorescence and thereby to determine the melting curve. A relative value for the initial target concentration in each reaction was determined on the basis of the kinetic approach using the LightCycler software, ver. 3.5. These real-time quantitative RT-PCR data were calculated by the 2−ΔΔCt method for RNA quantification [25]. A gene expression in tumor tissue less than 2-fold of the normal counterpart was defined as under-expression and one with greater than 2-fold the normal counterpart was defined as over-expression.

3. Results

3.1. Analysis of the over-expressed proteins from OSCC tissue extracts

In this study, we investigated the reliability of protein detections by coupling an improved extraction method and a 2D-PAGE method previously optimized for soluble proteins, using the OSCC as a model system. A flow chart of the methods used in this work is depicted in Fig. 1. To identify the over-expressed proteins in OSCC tissues, we used the 2D-PAGE technology to monitor changes in the abundance of proteins. Total protein extraction rates do not differ significantly between OSCC and matching surrounding normal tissues. The proteins were quite evenly distributed in gel with P/S in the range of 3–10 and molecular masses of 10–250 kDa. Fig. 2 shows representative 2D master gel images for both OSCC (Fig. 2A) and matching normal samples (Fig. 2B). Some proteins were present in train spots, indicating that modified proteins or isoforms were well separated by 2-DE.

Averages of 800 spots were detected across all gels. Fully automated spot detection and quantification were also performed by PDQuest software, image pairs from each patient, followed by automated image to image matching and statistical analysis. The ratio values were compared and abundance ratios
between different samples from different gels were calculated. The ratio values are expressed as fold changes, i.e., a 2-fold increase in OSCC is expressed as 2.0. In the OSCC tissues, we found 41 commonly over-expressed protein spots (p<0.001) compared to the normal tissue pairs (data not shown). Highlighted in squares are the 11 protein spots that consistently showed significant over-expression in OSCC tissues in all cases analyzed. Statistic data for these proteins (or the sum of the isoforms) are summarized in Table 2 with fold differences in OSCC tissues relative to the normal tissue pairs (for an average of 10 pairs indicated). The table shows the alterations in the over-expression levels of the identified protein spots in OSCC tissues. 

αB-crystallin was increased 7.2-fold in OSCC that was the highest level over-expression among the 11 proteins. GST-pi was the lowest level (2.5-fold).

3.2. Protein identification by nanoLC-MS/MS analysis

Each protein spot was excised and subjected to in-gel tryptic digestion, nanoLC-MS/MS and database matching. Table 3 is a summary of the 11 over-expressed proteins: most of the matched proteins had high sequence coverage, accession number, theoretical Mr (kDa)/pl and MOWSE scores.

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### Table 2

<table>
<thead>
<tr>
<th>Spot no.</th>
<th>Protein ID</th>
<th>Experimental Mr (kDa)</th>
<th>Experimental pl</th>
<th>Over-expression (fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>αB-crystallin</td>
<td>14.4–16</td>
<td>6–7</td>
<td>+7.2</td>
</tr>
<tr>
<td>2</td>
<td>Tropomyosin 2</td>
<td>31</td>
<td>5–5.5</td>
<td>+6.0</td>
</tr>
<tr>
<td>3</td>
<td>Myosin light chain 1</td>
<td>14.4–18</td>
<td>4–5</td>
<td>+5.7</td>
</tr>
<tr>
<td>4</td>
<td>HSP27</td>
<td>21.5–37</td>
<td>6–6.5</td>
<td>+4.3</td>
</tr>
<tr>
<td>5</td>
<td>Stratfin</td>
<td>14.4–21.5</td>
<td>4–5</td>
<td>+3.6</td>
</tr>
<tr>
<td>6</td>
<td>Thioredoxin-dependent peroxide reductase</td>
<td>15–21.5</td>
<td>7.5–8</td>
<td>+3.4</td>
</tr>
<tr>
<td>7</td>
<td>Flavin reductase</td>
<td>14.4–21.5</td>
<td>7–7.5</td>
<td>+3.0</td>
</tr>
<tr>
<td>8</td>
<td>Vimentin</td>
<td>36.5–42</td>
<td>5</td>
<td>+3.0</td>
</tr>
<tr>
<td>9</td>
<td>Rho GDI-2</td>
<td>14.4–21.5</td>
<td>5.5</td>
<td>+2.6</td>
</tr>
<tr>
<td>10</td>
<td>GST-pi</td>
<td>14.4–21.5</td>
<td>5.5–6</td>
<td>+2.5</td>
</tr>
<tr>
<td>11</td>
<td>MnSOD</td>
<td>19–21.5</td>
<td>8–9</td>
<td>+2.1</td>
</tr>
</tbody>
</table>

HSP27, heat shock protein 27; MnSOD, superoxide dismutase [Mn]; Rho GDI-2, Rho GDP-dissociation inhibitor 2; GST-pi, glutathione S-transferase pi.
3.3. Functional annotation of over-expressed proteins identified from OSCC tissues

One goal of this study was to discover novel components of the OSCC proteome in order to identify potentially new biomarkers for OSCCs. To gain a better understanding of the 11 proteins identified in this study, we carried out functional annotation analysis of these proteins. Grouping and naming of the identified proteins in the functional annotation analysis were done according to the Gene Ontology convention [26]. Information on the four functional categories, i.e., biologic process, cellular component, molecular function and cancer-related protein was collected from the 11 proteins (Table 4).

Table 4
Classification of identified proteins by functional category

<table>
<thead>
<tr>
<th>Category</th>
<th>Proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biologic process</td>
<td></td>
</tr>
<tr>
<td>Heat shock response</td>
<td>HSP27, αB-crystallin</td>
</tr>
<tr>
<td>Response to stimulus</td>
<td>HSP27</td>
</tr>
<tr>
<td>Cell motility</td>
<td>Rho GDI-2</td>
</tr>
<tr>
<td>Molecular function</td>
<td></td>
</tr>
<tr>
<td>Catalytic activity</td>
<td>GST-pi, flavin reductase, thioredoxin-dependent peroxide reductase, MnSOD</td>
</tr>
<tr>
<td>Ion binding</td>
<td>MnSOD</td>
</tr>
<tr>
<td>Structural molecular activity</td>
<td>Vimentin, myosin light chain 1, tropomyosin 2</td>
</tr>
<tr>
<td>Chaperon</td>
<td>Stratifin, HSP27</td>
</tr>
<tr>
<td>Cancer-related protein</td>
<td>MnSOD, HSP27</td>
</tr>
</tbody>
</table>

3.4. Detection of the gene expression levels of the over-expressed proteins

In order to determine whether observed over-expressed proteins in OSCC tissues occurred at the levels of gene expression, we performed real-time quantitative RT-PCR to evaluate the levels of mRNA of the 11 over-expressed proteins. These real-time quantitative RT-PCR data were calculated by using the $2^{-\Delta\Delta Ct}$ method.

The expression levels of αB-crystallin, myosin light chain 1 and MnSOD were the highest among 11 genes in all OSCC samples; the average increase over normal levels being 3.37-, 1.69- and 2.02-fold, respectively. The elevations of HSP27, Vimentin, and Rho GDI-2 in OSCC tissue were less obvious, with the average increase being over 0.79-, 1.07- and 1.04-fold (Fig. 3).

4. Discussion

Oral cancer remains one of the leading causes of death among men and therefore new protein markers have to be identified for its early detection. Identification of new biomarkers by proteomics currently relies on the 2-DE technology, picking out single spots, tryptic digestion and sequencing by MS. Once protein spots of interest have been identified as potential biomarkers, it is necessary to establish a more reliable assay. This is usually dependent on the development of a real-time quantitative RT-PCR or similar assay. The approach described in this study has the capability for improved screening and identification of over-expressed proteins in OSCC patients with oral cancer. In the present study, we quantified the protein levels in OSCC and the normal tissue pair with the goal to detect differences in the levels of proteins, which could serve as markers for the disease.

The number of proteins selected with over 2.0-fold induction consistent within 10 OSCC cases by our proteomic analysis consisted of 11 of the total approximately 800 proteins (Table 2). Even so, because the expression of these 11 proteins was consistently changed with OSCC, they may have relevance to the malignant alteration of oral mucosal epithelial cells. These seven genes have the potential of playing significant roles in...
elucidating the molecular mechanisms of the carcinogenesis and malignant behavior. At the mRNA level, therefore, it may be very difficult to find an mRNA that is universally up-regulated or down-regulated in all oral cancer patients (Fig. 3). The reason for the heterogeneous changes of mRNA level may be due to the fact that the biology of oral cancer is very complex at the molecular level and there is no or very few common changes at the mRNA and protein levels among very heterogeneous oral cancer patients.

Accumulating evidence has indicated that intracellular redox state plays important roles in cellular signal transduction and gene expression [27]. It has been suggested that ROS plays a role in all stages of carcinogenesis, including initiation, promotion, and progression [28]. In order to protect themselves from oxidative radical stress, cells have developed defense systems that comprise protein superoxide dismutases (SOD), catalase, glutathione peroxidases, and peroxiredoxins (PRX). SODs are antioxidant enzymes that catalyze the dismutation of O$_2^−$ free radicals in H$_2$O$_2$, thereby preventing the accumulation of this activated oxygen species. H$_2$O$_2$ can be further converted into H$_2$O and O$_2$ by catalase and/or glutathione peroxidase.

The manganese-dependent MnSOD is characteristic of aerobic organisms and composed of 4 homologous 24 kDa subunits [29]. While MnSOD has been reported to protect the cell from a variety of insults and to suppress apoptosis [30], this enzyme may also be deleterious and impede cell proliferation under some circumstances [31,32]. Thus, MnSOD appears to control multiple reactions, the orchestration of which will determine the fate of cells, in particular cancer cells [33]. Following the previous study, the up-regulation of MnSOD correlated with progression of disease from premalignant to invasive cancer reflects the cell defense effort in maintaining intracellular homeostasis [34]. Similar observations have been found in other cancers including the over-expression of MnSOD in human prostate cancer cell lines [35] and in buccal SCC [36]. The role of MnSOD in carcinogenesis is still controversial and unclear. In our study, MnSOD over-expressed in the protein level and correlated with gene expression. The elevation of MnSOD may imply that the antioxidant defense system has been stimulated in OSCC tissues.

Experimental evidence suggests that HSPs may promote tumorigenesis by suppressing apoptosis [37]. On the contrary, it seems that some members of HSPs play important roles in the immune response against cancer [38]. HSP27 is a stress-inducible cytosolic protein that ubiquitously present in many normal tissues [39]. Recent studies have shown that HSP27 may play a role in thermotolerance, cellular proliferation and apoptosis, estrogen response and molecular chaperoning [40–44]. Over-expression of HSP27 has been reported in many kinds of tumor tissues and found to be associated with the prognosis of astrocytic brain tumor, breast cancer, malignant fibrous histiocytoma, ovarian carcinoma, osteosarcoma, hepatocellular carcinoma and renal cell carcinoma [45–52]. Moreover, it was also reported to play a cytoprotective role in hormone-refractory prostate cancer [53]. Therefore, HSP27 will not be a specific tumor marker for OSCC. Among the several HSP proteins, HSP27 and HSP70 have been shown to have a strong association with cancer, showing, during carcinogenesis, alteration of their expression levels, either increasing or decreasing [54]. Moreover, the HSP27 was known to be a constitutive protein in many cell types, therefore it was only over-expressed in protein levels, but not in gene levels in OSCC tissues which is a common phenomenon (Table 2 and Fig. 3).

To our knowledge, MnSOD and HSP27 are both cancer-related proteins. Using proteomic and real-time quantitative RT-PCR analyses, we established initially that there is over-expression of MnSOD and HSP27 in OSCC tissues from oral cancer patients in this study. In order to understand the biological role of MnSOD and HSP27 in tumorigenesis of oral cancer, we will study their expression in oral cancer by immunohistochemistry (IHC) and western blot in samples of OSCCs and in oral relative cell lines, correlating its expression with clinicopathological features. Now, for further validation of MnSOD and HSP27 as the OSCC biomarkers, we used a larger cohort of samples and studies on the molecular mechanisms in OSCC carcinogenesis are being carried out in our lab. The current success in identifying a number of tumor-associated proteins proved that proteomic analysis could provide rich information to help understand the pathology of a disease in an integrated way. These extensive protein variations indicate that multiple cellular pathways were involved in the process of tumorigenesis, and suggest that multiple protein molecules should be simultaneously targeted as an effective strategy to counter the disease. These tumor-associated proteins can be further evaluated as potential biomarkers for clinical diagnosis and as targeted proteins for pathogenetic investigations.

Acknowledgments

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References


