Abstract. Within the past decade, the incidence of breast cancer in Taiwan has been rising year after year. Breast cancer is the first most prevalent cancer and the fourth leading cause of cancer-related deaths among women in Taiwan. The early stage of breast cancer not only have a wider range of therapeutic options, but also obtain a higher success rate of therapy than those with advanced breast cancer. A test for tumor markers is the most convenient method to screen for breast cancer. However, the tumor markers currently available for breast cancer detection include carcinoembryonic antigen (CEA), carbohydrate antigen 15.3 (CA15.3), and carbohydrate antigen 27.29 (CA27.29) exhibited certain limitations. Poor sensitivity and specificity greatly limits the diagnostic accuracy of these markers. This study aims to identify potential tumor markers for breast cancer. At first, we analyzed genes expression in infiltrating lobular carcinoma, metaplastic carcinoma, and infiltrating ductal carcinoma of paired specimens (tumor and normal tissue) from breast cancer patients using microarray technology. We selected 371 overexpressed genes in all of the three cell type. In advanced breast cancer tissue, we detected four genes MMP13, CAMP, COL10A1 and FLJ25416 from 25 overexpressed genes which encoded secretion protein more specifically for breast cancer than other genes. After validation with 15 pairs of breast cancer tissue and paired to normal adjacent tissues by membrane array and quantitative RT-PCR, we found MMP13 was 100% overexpressed and confirmed to be a secreted protein by Western blot analysis of the cell culture medium. The expression level of MMP13 was also measured by immunohistochemical staining. We suggest that MMP13 is a highly overexpressed secretion protein in breast cancer tissue. It has potential to be a new tumor marker for breast cancer diagnosis.

Introduction

New cases (910,000) of breast cancer appear globally every year, and 376,000 people die as a result (1). In 2006, breast cancer is estimated to account for 31% new onset female cancers (2). In Taiwan, breast cancer has the highest incidence among all female cancers (3) and is ranked fourth in all cancer deaths (4). In the Taiwanese breast cancer patients the first occurrence of this disease is often at a young age. According to statistics, more than 50% of total breast cancer diagnosed annually in Taiwan derives from patients under 50 years old, and this proportion is higher than that observed in Western population (5,6).

Early diagnosis of breast cancer can provide patients a wider range of therapeutic options as well as a higher success rate of therapy that lowers mortality. Through proper treatment, the 10-year survival rate can reach 60%, and the stage I survival rate can be as high as 80%. Survival rate in stage 0 is close to 100% (7). These data indicate that early diagnosis and treatment of breast cancer is the key to higher survival rates.

Currently, mammography, MRI (magnetic resonance imaging), and tumor markers are the most commonly used diagnostic tools for breast cancer but they have inherent limitations. Mammography is widely used as a convenient screening tool in women over 50 years, but it does not have high sensitivity. Therefore, <50% of patients are diagnosed...
Results from this study demonstrated that MMP13 is a secretion protein. Finally, using immunohistochemical (IHC) staining, we proved that MMP13 was highly overexpressed in the breast tissue of general breast cancer patients (100%).

Of the 25 genes, MMP13, CAMP, COL10A1, and FLJ25416 were selected. We discovered 25 genes which encode secretion proteins that are more specific for breast cancer than other genes. We performed microarray analysis using the Agilent Oligo array. As a result, 371 genes with expression levels in tumor tissue 2-fold higher than normal tissue in all three cell type groups were selected. We discovered 25 genes which encode secretion proteins from 371 genes by employing bioinformatics studies. Of the 25 genes, MMP13, CAMP, COL10A1, and FLJ25416 are more specific for breast cancer than other genes. We utilized membrane array and RT-PCR in 15 pairs of tissues of different cell types (infiltrating lobular carcinoma, metastatic carcinoma, and infiltrating ductal carcinoma) and performed microarray analysis using the Agilent Oligo array. Predicting secretion protein. We used three types of breast cancer and paired them with the corresponding normal tissues to perform three microarray technology studies. After analysis with GeneSpring Biological data analysis software, we found that 371 genes were overexpressed 2-fold or more in the three groups of breast cancer tissues. The overexpressed gene groups of the breast tumors were further analyzed with Swiss-Prot, Secreted protein database (SPD) (14), Signal P3.0 (15), and pTARGET (16) to predict and screen the ones that were secretion proteins. The selected ones represent breast cancer tumor marker candidate genes.

Virtual Northern analysis. The functions of Cancer Genome Anatomy Project (CGAP) of National Center for Biotechnology Information (NCBI), genetic information of the breast cancer tumor marker candidate genes, such as EST data and SAGE data, were used as adjunctive confirmation, to further select the ones with the most specificity for breast cancer.

Total RNA extraction and first strand cDNA synthesis. Total RNAs were extracted from breast cancer patient's tissue and cell line with ISOGENTM (Nippon Gene, Toyama, Japan) and QiAmp® RNA Blood Mini Kit (Qiagen Inc., Valencia, CA) according to the manufacturer's instructions (17).

Clinical tissue sample collection. All paired samples including tumor and tumor-free tissues were obtained by clinical pathologist from 45 patients diagnosed of breast cancer at Chung-Ho Memorial Hospital of Kaohsiung Medical University during 2003-2007. Tumor grading was carried out and confirmed by pathologists. Samples were further used for microarray analysis, establish primary culture cell lines, real-time PCR and membrane array analysis.

Oligonucleotide microarray. The oligonucleotide array contains 22,500 elements designed for expression profiling (Human 1A V2, Agilent Technologies, Palo Alto, CA, USA), for which over 18,000 well-characterized, full-length human genes were defined. First-strand cDNA targets for hybridization were made by reverse transcription of the mRNA isolated from both the tumor and paired normal tissues from the same case by using SuperScript II RT (Gibco-BRL, Gaithersburg, MD, USA) in the presence of either Cy3- or Cy5-labeled dUTP (Amersham Pharmacia Biotech, Piscataway, NJ, USA). The targets were dried to 18 μl by a SpeedVac™ concentrator (Thermo Electron Co., Waltham, MA, USA), and 3.6 μl 20X SSC, 1.8 μl 10 mg/ml poly-A and 0.54 μl 10% SDS were added. Then, the mixture was heated to 100°C for 2 min proceeding to hybridization reaction on Human 1A Oligo Microarray V2 array slides (Agilent Technologies) in an incubator at 63°C for 12-16 h. After being sequentially washed with 1X SSC, 0.2X SSC and 0.05X SSC, hybridized microarray slides were scanned and fluorescence signals were detected by using an Axon GenePix 4000A dual-color confocal laser scanner (Axon Instruments, Union, CA, USA). Subsequent quantification analysis was performed using the commercial software GenePix Pro 3.0™ (Axon Instruments). The acceptance criterion for a gene signal was a signal-to-noise ratio of ≥2. If either the Cy3 or Cy5 signal of a specific spot passed the criterion, the flag of its ratio was counted to be ‘TRUE’. The elements with the ‘TRUE’ flag were analyzed with GeneSpring GX7 (Silicon Genetics, Redwood, CA, USA).

Materials and methods

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The differentially expressed elements were analyzed by the two-sided statistical tolerance interval (95%).

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Recombinant RNasin® Ribonuclease Inhibitor (Promega). The reaction mixtures with RNA were incubated at 42˚C for longer than 2 h, heated to 95˚C for 5 min, and then stored at -80˚C until analysis.

Real-time polymerase chain reaction (RT-PCR). Two microliters of each cDNA sample were used for each reaction. Sequences of the oligonucleotide primers (Table I) were designed according a PCR primer selection program based on primer 3 at http://frodo.wi.mit.edu/cgi-bin/primer3/prim3www.cgi. Each RT-PCR reaction mixture contained 1X PCR buffer (10 mmol/l Tris-HCl, pH 8.3, 50 mmol/l KCl, 2 mmol/l MgCl₂), 50 μmol/l dNTP, 0.1 μmol/l sense and antisense primers for target genes, and 0.01 μmol/l sense and antisense primers for ß-actin. PCR products were analyzed in 3% agarose gel containing 0.5 μg/ml ethidium bromide. The signals on UV transilluminator for each target gene and ß-actin expression levels were scanned with a computing laser densitometer (Alpha Inotech, San-Leandro, CA) to calculate the relative mRNA density. Real-time PCR was performed in a Rotor-Gene 2000 thermocycler (Corbett Research, Inc.). The reaction mixture contained 2 μl of 20 mM dNTP, 2 μl of 30 mM MgCl₂, 2 μl of 20X SYBR-Green, 2 μl of 1 μM forward primer, 2 μl of 1 μM reverse primer, 4 μl of nuclease free water, 2 μl of 80-100 ng/ml cDNA, and 2 μl of 1 U/μl polymerase. PCR conditions were as follows: 35 cycles of denaturation at 95˚C for 20 sec, annealing at 60˚C for 20 sec, and extension at 74˚C for 40 sec. PCR products (i.e. synthesized dsDNA) were quantified by measuring the fluorescent intensity at the end of each amplification cycle. For each sample, real-time PCR analysis was repeated in three independent experiments to ensure the reproducibility of results. We used ß-actin and Oryza sativa sequence as positive and negative controls, and DDwater as blank control.

Membrane array. The procedure of the membrane-array method for the detection of multiple genes simultaneously was performed based on our previous work (18-20). Visual OMP3 (Oligonucleotide Modeling Platform, DNA Software, Ann Arbor, MI) was used to design probes for each target gene and ß-actin, and the latter served as an internal control (Table II). The probe selection criteria included strong mismatch discrimination, minimal or no secondary structure, the signal strength at the assay temperature, and lack of cross-hybridization. The oligonucleotide probes were then synthesized according to the designed sequences, purified, and controlled before being grafted onto the subtracts. The newly synthesized oligonucleotide fragments were dissolved in distilled water to a concentration of 20 mM, applied to a BioJet Plus 3000 nl dispensing system (BioDot, Irvine, CA), which blotted the four target oligonucleotides, and the ß-actin

<table>
<thead>
<tr>
<th>Gene</th>
<th>Sequence Product (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAMP</td>
<td>Forward 5'-CAGCAGGGCAAATCCTGTATATTCCT-3' 241</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-CTACCCGCTTCTGGACCT-3'</td>
</tr>
<tr>
<td>COL10A1</td>
<td>Forward 5'-CCTCACTTTGAATGGGACGACAGGG-3' 228</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-TGCAAGTCTTTCAGTATGGAAGG-3'</td>
</tr>
<tr>
<td>MMP13</td>
<td>Forward 5'-GTTCTCTCCTTGGGTAAGGCT-3' 140</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-GTGATCTCTTGGACATGGAAGG-3'</td>
</tr>
<tr>
<td>FLJ25416</td>
<td>Forward 5'-GCTCCCACGAGCATCTTGTTTTCAAT-3' 190</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-AGGCCCACCTCTCAGTGATGAAACT-3'</td>
</tr>
<tr>
<td>ß-actina</td>
<td>Forward 5'-GCATCCACGAAACTACCTTC-3' 183</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-CAGGAGGAGGATGATCTTG-3'</td>
</tr>
</tbody>
</table>

*ß-actin primers were added as internal controls to correct for the differences in different cells.

Table I. Primer sequences of target genes.

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Oligonucleotide sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>COL10A1</td>
<td>5'-AGATATTGTGAAAGGCGCATACAGGCCTAGAGTGATGCACCCATAGA-3'</td>
</tr>
<tr>
<td>CAMP</td>
<td>5'-CTCTATAGGCAGGGCCACAGCTCTTGGATGATGACCTGAT-3'</td>
</tr>
<tr>
<td>MMP13</td>
<td>5'-AATAAGGCTCAGACAGCCACTCTGATCGATGAGATGGT-3'</td>
</tr>
<tr>
<td>FLJ25416</td>
<td>5'-TGACTCTTTGAACAAATCAACTTCAGTCCCAGATTAGGTCAGTGATG-3'</td>
</tr>
</tbody>
</table>

Table II. Oligonucleotide sequences of target genes.
control sequentially (0.05 μl per spot and 1.5 mm between spots) on SuPerCharge nylon membrane (Schleicher and Schuell, Dassel, Germany) in triplicate. Dimethyl sulfoxide (DMSO) was also dispensed onto the membrane as a blank control (Fig. 1A). After rapid drying and cross-linking procedures, the preparation of hypoxia- and glycolysis-associated gene expression membrane array was accomplished (21).

Preparation of digoxigenin (DIG)-labeled cDNA targets and hybridization. First-strand cDNA targets for hybridization were made by reverse transcription of the mRNA from the tumor and corresponding normal tissues of breast cancer patients in the presence of DIG-labeled UTP (Roche Diagnostics GmbH, Penzberg, Germany) using SuperScript II reverse transcriptase (Gibco-BRL). The membrane array needs prehybridization at 42˚C for 2 h in a hybridization oven (Autoblot Bellco, Vineland, NJ, USA), and subsequent blocking before hybridization. The lifts were covered with the ExpressHyb Hybridization Solution (BD Biosciences, Palo Alto, CA, USA) containing DIG-11-UTPlabeled cDNA probes, and then incubated with an anti-DIG alkaline phosphatase conjugated antibody (Roche Diagnostics) at a dilution of 1:2500 (75 mU/ml) in 1X blocking buffer (100 mM Maleic acid, 150 mM NaCl, pH 7.5). We incubated the arrays for hybridization at 42˚C overnight in a humid shaking chamber. After washing each one three times for 10 min in 20 ml washing buffer, the arrays were exposed to excitation light. For signal detection, the membranes we incubated for 15 min without shaking in a chromogen solution containing nitroblue-tetrazolium and 5-bromo-4-chloro3-indolyl-phosphate (NBT/BCIP).

Statistical analysis of membrane array. All data were analyzed using the Statistical Package for the Social Sciences Version 11.5 software (SPSS Inc., Chicago, IL). Data are presented as means ± SE and P-values were determined by unpaired Student’s t-test. Furthermore, the two-sided Pearson χ² test was used to analyze the differences in the overexpression of the genotypes between different age groups, gender and stages. A P-value of <0.05 was considered statistically significant.

ATCC cell lines. Five different kinds of cell lines were purchased from ATCC (American Type Culture Collection, ATCC, Rockville, MD, USA) including four human breast cancer cell lines [ZR-75-1(CRL-1500), MDA-MB-231 (HTB-26), T-47D(HTB-133) and MCF-7(HTB-22)], two human colorectal cancer cell lines [CCL-228(SW480) and CCL-227(SW620)], one normal human cell line [CRL-9609 (BEAS-2B)], two human lung cancer cell lines [CRL-5800 (NCI-H23) and CRL-5807(NCI-H358)] and one human cervical cancer cell line [CRL-13011(HeLa NR1)]. ZR-75-1(CRL-1500), T-47D(HTB-133), MCF-7(HTB-22), CRL-9609(BEAS-2B), CRL-5800(NCI-H23) and CRL-5807(NCI-H358) and CRL-13011(HeLa NR1) cell lines were cultured with RPMI-1640, 10% FBS, 1.5 gl sodium bicarbonate, 10 mM HEPES, 1.0 mM sodium pyruvate and 1% antibiotic-antimycotic cell cultivating solution at 37˚C and 5% CO₂; MDA-MB-231(HTB-26), CCL-228(SW480), CCL-227(SW620) was cultured with Leibovitz’s L-15 medium, 10% FBS, and 1% antibiotic-antimycotic of cell cultivating solution at 37˚C and 100% air.

Establishment of primary culture cell lines. Fresh samples collected from the operating room were immersed in 50 ml of phosphate-buffered saline (PBS), in a centrifuge tube and maintained at 4˚C, followed by transport to the laboratory for immediate cultivation. The time from the removal of sample from the patient in the operating room to the start of the
cultivation process did not exceed 4 h. Operational procedures referred to Freshney (23), mainly involving enzyme digestion and they were gradually adjusted based on process of the study and experience. After testing under different conditions, the following standard procedures were utilized in the study: 0.00625 g of collagenase (Sigma C-9891) was dissolved in 12 ml Hank's balanced salt solution, then prepared through a 0.22 μm filter. Sample tissues were first washed with PBS twice under the laminar flow hood, then placed into a 6-well culture plate and cut into pieces with a No.10 surgical blade. After cutting to a size of about 1 mm², the tissues were washed with collagenase solution and transferred to a 50 ml centrifuge tube, and placed in room temperature to react for 45-50 min, depending on the size of the tissue sample. A 50-ml centrifuge was prepared and 10 ml of RPMI-1640 and 10% FBS cell culture solution were added. After waiting 45-50 min for the collagenase to act, the solution was filtered through 0.5 mm of molecular pores into the prepared 10 ml RPMI-1640, and 10% FBS cell culture solution was added to stop the action of collagenase. The solution was then centrifuged at 1,000 rpm for 10 min. The clear solution on top was discarded, and then the RPMI-1640 and 10% FBS cell culture solution was used to dispense the cells. Finally, cells were transferred to a T25 tissue culture flask. The flask was placed into a fixed-temperature CO2 incubator. The growth criteria for the cell line were 37°C, 5% CO2, and pH 7.4. This primary culture cell line displayed biological characteristics of slow growth, and the subculture required a significantly long time, which corresponded with the postulations of Fournier et al. (22). To minimize the mixture of fibroblasts during cell line cultivation, we employed different speeds in cell adherence to gradually reduce the amount of fibroblasts (23). Fibroblasts have the characteristic of rapid adherence (after subculture with pancreatin for 30 min, adherence begins). Therefore, in each subculture process, the tissue culture flask was first placed in a 37°C, 5% CO2 fixed temperature CO2 incubator for cultivation. Afterwards, the culture solution containing unadhered epithelial cells was transferred to a new tissue flask. This method significantly reduces the amount of fibroblasts.

Western blot analysis. Cells were placed in a T75 culture dish and grown to 80% confluence. Next, the cell culture solution was discarded, and the cells were washed three times with PBS. Afterwards, 8 ml of PBS culture solution was added, and placed into a 5% CO2 incubator at 37°C for cultivation. After 48 h, the top clear solution was collected, and then concentrated with an Amicon Ultra-4 PLGC Centrifugal Filter Unit (Blossom Biotechnologies Inc., Taiwan). After concentration, the clear cell solution on top was immediately stored at -70°C. After 4 μl of reducing dye was added into 20 μl of the concentrated clear cell solution sample, it was poured onto a specifically made 10% SDS electrophoresis gel for electrophoresis. The electric current was 40A. After electrophoresis, 95 V was used to blot the protein on the gel onto nitrocellulose paper (NC paper). Skim-milk was used to fill the gaps in the NC paper. Target protein antibodies were added under room temperature and shaken for 1 h for specificity linking. After washing off non-specific bound antibodies with PBS, secondary antibodies of the target HRP were added and shaken for 1 h for immune reactions. After washing off extra antibodies, the substrate, Immobilon Western Chemiluminescent HRP Substrate (Millipore, Mississauga, ON), was added to achieve color effects.

**Immunohistochemical stain (IHC).** The biopsy samples of breast cancer tissues were baked on slides at 60°C for 1 h. Then, the biopsy samples were soaked in xylene for 10 min, and repeated twice. Afterwards, it was immersed in 100% of absolute alcohol for 5 min, and repeated twice. Then, it was soaked in 85 and 75% alcohol, respectively, for 5 min each and then in distilled water for 5 min each for removal of wax. De-waxed biopsy samples were soaked in 10 mM of sodium citrate, and placed in a microwave for 5 min. Three repeated antigen retrieval procedures were then performed. When the tissue slides were cooled to room temperature, 3% of H2O2 was used to remove intrinsic catalase. Afterwards, the slides were washed with distilled water followed by blocking serum (normal rabbit serum 20X) being dropped on it for 1 h. Then, first antibody, secondary antibody of the target protein, and Streptavidin-HRP (Dako, #K0675) were added separately and allowed to act for 1 h. After washing off the antibody at the end of the specific bond, DAB substrate was added (20 ml of DAB+1 ml sub-strate buffer solution; Dako, #K3468). The slides were observed at all times under the microscope to prevent over dyeing. Finally, they were washed with TBST for 5 min three times, and the nuclei were stained with hematoxylin, using 1% NH3OH. The final step to make it transparent included passing the slide through 75, 85, 95 and 100% alcohol at 3-min intervals each to remove water. After passing through xylene for 5 min for secondary transparency, filter paper was used to dry the tissue biopsy samples and covered with a coverslip.

**Results**

**Microarray analysis of clinical breast cancer tissues pairs.** We utilized three pairs of breast cancer and tumor-free tissues of different cell types (infiltrating lobular carcinoma, metaplastic carcinoma, and infiltrating ductal carcinoma). All results from the experiments underwent standardized analysis and validation. Then, we used biological data software, GeneSpring, hierarchical clustering was performed in the three experimental groups to initially assess gene expressions of all genes on the chip. After further analysis and validation, 371 genes displayed overexpression and potentially serving as biological markers in breast cancer.

**Identification of candidate genes by bioinformatics.** The 371 overexpressed genes validated in the breast tissues, were analyzed with biological software such as Swiss-Prot secreted protein database (SPD) (14), Signal p3.0 (15), and pTARGET (16), and results indicated that a total of 25 genes contained a structural sequence of secretion protein (Table III). Furthermore, with the use of Virtual Northern functional software from NCBI, the EST and SAGE data of the 25 candidate genes were assessed for each gene's expression in different cancerous tissues. The genes, MMP13, CAMP, COL10A1 and...
FLJ25416 possessed the highest specificity for breast cancer. Therefore, these four genes were selected as the candidate target genes for this study.

**Gene validation in clinical breast cancer tissues by membrane array and RT-PCR.** To validate the expression of the above-mentioned four genes in the breast tissue of clinical breast breast cancer patients, both membrane array and RT-PCR were conducted to detect the mRNA expression level from the 15 paired tissues. Based on results of membrane array hybridization analysis (Fig. 1B), if the gene presented a color response of >2-fold between the breast cancer and normal tissue, it was defined as overexpression. We found four genes with overexpression in breast cancer tissues (Table IV). MMP13 showed 100% overexpression in 15 breast cancer paired tissues, and was the highest among the four genes. The same finding was seen in RT-PCR. This indicated a high degree of consistency and correlation between the results of RT-PCR analysis and that of membrane array hybridization.

**MMP13 protein was overexpressed in human breast cancer tissue.** To further validate whether MMP13 is indeed a secretion protein, other than cultivating 4 lines of ATCC human breast cancer cell lines and 2 lines of primary culture cell lines of breast cancer, we also performed Western blot
analysis for MMP13 protein detection in 2 lines of ATCC human colorectal cell lines, 2 ATCC human lung cancer cell lines, and 1 ATCC human cervical cancer cell line. MMP13 (pro form) and MMP13 (active form) protein of 60 kDa of molecular weight were detected on the clear cell solution (Fig. 2). The test results of all cell culture solutions, we found that all the different cell lines could detect MMP13 except lung normal cell line (Table V). Interestingly, MMP13 is indeed a secretion protein and it is highly expressed in breast cancer and colon cancer cell lines. At the same time, we utilized MMP13 antibody to perform IHC stain in the collected tissue biopsies of breast cancer patients, to investigate MMP13 expression in breast cancer tissues. The antibody used was the same as that in Western blot analysis. A positive

Table V. Results of MMP13 secretion in 17 cell lines.

<table>
<thead>
<tr>
<th>Breast cancer</th>
<th>Colorectal cancer</th>
<th>Lung normal</th>
<th>Lung cancer</th>
<th>Cervix cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>HTB26 (+)</td>
<td>SW480 (+)</td>
<td>CRL9609</td>
<td>CRL5800</td>
<td>CRL13011 (+)</td>
</tr>
<tr>
<td>HTB133</td>
<td>SW620 (+)</td>
<td></td>
<td>CRL5807 (+)</td>
<td></td>
</tr>
<tr>
<td>CRL1500 (+)</td>
<td>CP1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MCF7</td>
<td>CP2</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>BP1 (+)</td>
<td>CP3</td>
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<td>BP2 (+)</td>
<td>CP4 (+)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>CP5</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

66.67% (4/6) 42.86% (3/7) 0% (0/1) 50% (1/2) 100% (1/1)

+ indicates the cell line secretions expressed MMP13 protein. The percentage at the bottom of the table indicates positive rate, or expression of MMP13 in each cancer cell line. BP, breast cancer primary culture; CP, colon cancer primary culture.
yellowish-brown appearance after tissue dyeing indicated that MMP13 was 100% overexpressed in all breast cancer tissues (Fig. 3).

Discussion

In recent years, many studies have proven that early diagnosis is one of the most effective methods to lower the incidence and mortality of breast cancer. Compared to traditional cellular diagnosis or pathological dyes, the clinical test method of tumor markers is more convenient, simpler, less expensive and less invasive for early diagnosis of breast cancer. In addition, the public readily accepts it. However, the current tumor markers widely used in breast cancer screening all have their individual restrictions with the primary issue being low specificity and sensitivity (24). In this study, we selected four candidate markers the MMP13, CAMP, COL10A1, and TLJ25416 from the initial gene group presenting overexpression. Among these, evidence indicated that MMP13 was the most commonly overexpressed gene in the tumor tissue of clinical breast cancer patients (100%). Therefore, MMP13 has tremendous potential to serve as a biological marker to assist in the clinical diagnosis of breast cancer.

The human MMPs family includes 24 members (25,26). This protein family mainly functions as proteolytic enzymes, and they can dissolve extracellular matrix (27,28). This family of proteins not only participates in normal physiological functions, such as wound healing and modulating growth factors and enzyme activation pathways (28-31), it also plays an important role in cancer infiltration, metastasis and angiogenesis (32-35). MMP13 (collagenase-3), MMP1 (collagenase-1) and MMP8 (collagenase-2) are all collagens. These three types of collagenase all have very high structural homogeneity. They can break down collagen types I, II, III, and V (31). Of these, MMP13 is most effective in breaking down type II collagen (36). Nielsen et al pointed out that in the process of breast cancer turning from ductal carcinoma in situ to invasive ductal carcinoma, MMP13 will break down basement membranes of tissues to form an invasive cancer (37). A consensual theory may be reached for this biochemical characteristic based on the results of this study, which demonstrate that MMP13 was only expressed in the highly invasive breast cancer cell line, MDA-MB-231, but not in the less invasive MLF-7 breast cancer cell line (38-40). This result also conforms to those gathered by Selvarumugan and Partridge (41).

In addition, many studies have discussed the relationship between MMP13 and other cancers. For example, MMP13 is overexpressed in colon cancer tissues (42,43). In a head and neck squamous cell carcinoma study, MMP13 was found to be related to the invasiveness of cancer cells, and could be used to evaluate patient prognosis (44). This result was also confirmed in oral squamous cell carcinoma where MMP13 was determined to be a potential OSSC tumor marker (45). MMP13 increased its overexpression in tissues after metastasis in prostate cancer patients. Therefore, it can be a biomarker for prostate cancer diagnosis, treatment monitoring, and prognosis determination (46). Of note, although MMP8 and MMP1 have homogeneous structures, MMP8 was postulated to play a protective role in a study related to lymph node metastasis (47). The age of onset of breast cancer patients in Taiwan is about 10 years earlier than patients in the Western nations, and in these young patients, the breast cancer is more invasive. Whether these differences relate to the expression of MMP13 and MMP8 in breast cancer tissues require further studies.

In the future, we will measure the MMP13 protein in the plasma of breast cancer patients and normal individuals, find the relationship between MMP13 concentration and clinical breast cancer symptoms, and clinically evaluate the sensitivity and specificity of MMP13 protein in breast cancer diagnosis. From results obtained in this study, we believe that MMP13 has the potential to become a new breast cancer tumor marker, when accompanied by current clinical screening methods, may increase the rate of early breast cancer diagnosis.

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