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Identification of biomarkers for lymph node metastasis in early-stage cervical cancer by tissue-based proteomics

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Background: Pelvic lymph node metastasis (PLNM) is the key to determining the treatment and prognosis of early-stage cervical cancer (CC, I–IIst). The aim of this study was to identify biomarkers for PLNM of CC, I–IIst.

Methods: Two-dimensional fluorescence difference gel electrophoresis and matrix-assisted laser desorption/ionisation-time-offlight mass spectrometry (MALDI-TOF/TOF MS) were used to identify differentially expressed proteins in primary CC, I–IIst tissue with (n = 8) and without (n = 10) PLNM. The expression levels of three differential proteins (FABP5, HspB1, and MnSOD) were validated using western blotting and immunohistochemistry. An independent cohort of 105 CC, I–IIst patients was analysed to assess the correlation of FABP5, HspB1, and MnSOD with clinicopathologic factors and clinical outcomes.

Results: Forty-one differential proteins were identified. Upregulation of FABP5, HspB1, and MnSOD in CC, I–IIst with PLNM was confirmed and was significantly correlated with PLNM. FABP5, HspB1, and MnSOD were significant predictors of PLNM in univariate analysis. FABP5, HspB1, and lymphovascular space invasion (LVSI) were independent predictors of PLNM in multivariate analysis. Survival curves indicated that CC, I–IIst patients with FABP5, HspB1, and MnSOD upregulation had poor prognosis.

Conclusions: FABP5, HspB1, and MnSOD may be potential biomarkers for PLNM of CC, I–IIst and may have important roles in the pathogenesis of PLNM.

Globally, cervical cancer is the third most common cancer in women (Jemal *et al*, 2011). Pelvic lymph node metastasis (PLNM) is a critical factor for determining the individualised treatment and prognosis of cervical cancer, particularly in early-stage cervical cancer (CC, I–IIst) (Gien and Covens, 2009). The primary treatment for CC, I–IIst is either surgery or chemoradiotherapy. For patients with low risk of PLNM, surgical treatment (radical hysterectomy/conisation/trachelectomy plus lymph node (LN) dissection) can offer hope of fertility or function preservation for young patients and may avoid the long-term complications of chemoradiotherapy (Kobayashi *et al*, 2006; Rob *et al*, 2008; Cao *et al*, 2013). For patients with high risk of PLNM, primary chemoradiotherapy should be recommended. However, because it is difficult to clinically detect LN metastasis efficiently and reliably, some patients receive suboptimal treatment (first unnecessary

surgery with adjuvant chemoradiotherapy) (Selman *et al*, 2008). Combining surgery and chemoradiotherapy leads to a higher morbidity, such as urological complications and lymph oedema (Landoni *et al*, 1997). Therefore, before the first treatment, stratification of CC, I–IIst patients based on PLNM is crucial to planning the optimal individualised treatment.

Imaging modalities, including computed tomography (CT), magnetic resonance imaging (MRI), and positron emission tomography (PET) scans are routinely used in preoperative examinations of PLNM. Nonetheless, these imaging modalities have relatively low sensitivity (Choi *et al*, 2006; Chou *et al*, 2006) for assessing PLNM in CC, I–IIst. Recently, laparoscopic lymphadenectomy was recommended before fertility-preserving surgery (Vercellino *et al*, 2012). However, lymphadenectomy is both invasive and associated with intraoperative and postoperative

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morbidity (van de Lande *et al*, 2012). Sentinel LN (SLN) biopsy is considered to be feasible for detecting the status of LN in earlystage cervical cancer (Plante *et al*, 2003). SLN biopsy can not only enhance the sensitivity and specificity of the diagnosis of PLNM but also reduce the extension of pelvic lymphadenectomy. Although it is less invasive than open and laparoscopic surgery, SLN biopsy still requires a general anaesthesia. At the present time, there are no known non-invasive and highly accurate detection methods that may be used to assess PLNM in CC, I–IIst before the first treatment. Therefore, an urgent need to identify biomarkers for PLNM in CC, I–IIst exists.

Previous studies for identifying PLNM-associated biomarkers in cervical cancer focused mainly on the transcriptomic (Huang et al, 2012) and genomic levels (Kim et al, 2008; Huang et al, 2011). However, alterations at the protein level can reflect cellular changes more accurately. Advances in bioinformatics and high-throughput technologies have provided powerful tools for unravelling the proteomic profile during PLNM progression. A recent study investigated sera from CC, I-IIst patients using surface-enhanced laser desorption/ionisation-time-of-flight mass spectrometry (SELDI-TOF-MS), and found that a model based on three differentially expressed peaks could determine LN status (Van Gorp et al, 2012). However, this study failed to identify the proteins represented by the three differentially expressed peaks. In contrast, our study was performed with primary CC, I-IIst tissue. Analysis of CC, I-IIst tissue has several potential advantages: first, there is an inherent link between the differential proteins and the tumour itself; second, we can take into account the contributions of the tumour microenvironment by testing protein expression in tissue, because the tumour microenvironment has a vital role in cancer metastasis (Langley and Fidler, 2011), and finally, primary cervical cancer tissue can be easily acquired by transvaginal biopsy before surgery. To our knowledge, there are no reports on the use of tissue-based proteomics to identify biomarkers associated with PLNM of CC, I-IIst.

The aim of this study was to identify biomarkers for PLNM of CC, I–IIst. We applied two-dimensional fluorescence difference gel electrophoresis (2D-DIGE) and matrix-assisted laser desorption/ ionisation-time-of-flight mass spectrometry (MALDI-TOF/TOF MS) to identify differential proteins between primary CC, I–IIst tissue with and without PLNM. Three differential proteins were further validated on an independent tissue cohort to evaluate their reliability in predicting PLNM of CC, I–IIst.

MATERIALS AND METHODS

Tissue sample collection. The study was approved by the Ethical Committee of the First Affiliated Hospital of Sun Yat-sen University (Guangzhou, China). Informed consent was obtained from each patient. Eighteen patients with early-stage cervical squamous cell cancer who underwent radical hysterectomy and pelvic lymphadenectomy at the First Affiliated Hospital of Sun Yat-sen University from January 2011 to September 2012 were recruited. None of these patients received preoperative radiation or chemotherapy, and none was documented as having other diseases per systematic examination before surgery. Based on the postoperative pathological examinations of pelvic LN (PLN), the CC, I–IIst cases were assigned into PLNM group (n = 8) and non-PLNM (NPLNM) group (n = 10). The clinicopathological characteristics of the tumour samples are shown in Table 1. There was no difference between the two groups in clinicopathological factors. To obtain tumour tissue, while on ice, each tissue sample was assessed and dissected by an experienced pathologist immediately after tumour resection. The pathologist then confirmed that each dissected tumour tissue block was composed of at least 80% tumour cells by frozen section examination. Dissected tissue samples were snap-frozen in liquid nitrogen with the shortest possible delay and were then stored at -80 °C until use.

For immunohistochemistry (IHC) validation, an independent tissue cohort with 105 early-stage cervical squamous cell cancer

No.	Age (years)	FIGO stage	Tumour size (cm)	Number of positive PLN	Number of removed PLN	LVSI	Differentiation	Pathological type	Stromal invasion
Early-	stage cervical o	ancer with	PLNM						
1	41	lla1	3	2	34	Negative	G2	Scc	> 1/2
2	56	lb1	2.5	3	36	Negative	G2	Scc	>1/2
3	42	lla2	5	2	32	Negative	G3	Scc	>1/2
4	58	lla1	3.5	2	40	Negative	G3	Scc	< 1/2
5	39	lb2	5	1	47	Negative	G2	Scc	>1/2
6	58	lb1	3	4	41	Negative	G2	Scc	> 1/2
7	67	lla2	5	13	43	Positive	G2	Scc	> 1/2
8	35	lb1	3.5	7	35	Negative	G3	Scc	<1/2
Early-	stage cervical o	cancer with	out PLNM						
1	40	lla2	5	0	35	Negative	G2	Scc	>1/2
2	64	lb1	2	0	34	Negative	G2	Scc	< 1/2
3	53	lla1	3.5	0	41	Negative	G2	Scc	< 1/2
4	32	lb1	3	0	37	Negative	G2	Scc	> 1/2
5	42	lb2	5	0	43	Negative	G2	Scc	> 1/2
6	42	lb1	3	0	48	Negative	G3	Scc	< 1/2
7	49	lla1	3.5	0	30	Negative	G2	Scc	> 1/2
8	57	lb1	3	0	51	Positive	G2	Scc	>1/2
9	48	lla2	4.5	0	46	Negative	G3	Scc	> 1/2
10	52	lla1	3	0	48	Negative	G3	Scc	< 1/2

Abbreviations: FIGO=the International Federation of Gynaecology and Obstetrics; LVSI=lymphovascular space invasion; PLN=pelvic lymph node; PLNM=pelvic lymph node metastasis Scc=squamous cell carcinoma.

samples (22 PLNM, 83 NPLNM) was used (Supplementary Table S1). This cohort of paraffin-embedded tissue material was collected from the archives of the pathology department at the First Affiliated Hospital of Sun Yat-sen University. Surgeries were performed between 2003 and 2007. None of the patients received radiotherapy and/or chemotherapy before surgery. All specimens were handled according to legal and ethical standards.

Protein sample preparation. Tissue ($\sim 100 \text{ mg}$) was mechanically homogenised in lysis buffer (2 M thiourea, 7 M urea, 30 mM Tris, 4% w/v CHAPS) on ice. These homogenates were sonicated intermittently and then separated by centrifugation at 20, 000 r.p.m. for 40 min at 4°C. The supernatant was collected. To purify the protein extract and to quantify the final protein concentration, a 2-D clean-up kit (Amersham Biosciences, Buckinghamshire, UK) and a 2-D Quant Kit (GE Healthcare, London, UK) were used sequentially according to the manufacturers' instructions. Equal amounts of proteins extracted from the same group were pooled for DIGE analysis.

Protein sample labelling with CyDye. CyDye DIGE fluors (Cy2, Cy3, and Cy5) were used to label the protein extracts following the manufacturer's protocol (GE Healthcare). The internal standard pool was generated by combining equal amounts of extracts from all samples, labelled with Cy2. Protein extracts from the PLNM and NPLNM groups were labelled with Cy3 and Cy5, respectively. The labelling reaction was performed on ice for 30 min in darkness, and was then quenched with 10 mM lysine for 10 min on ice under dark conditions. The labelled samples were then mixed and prepared for the following steps.

Two-dimensional electrophoresis. After rehydration, the labelled protein mixture for each gel was applied to an immobilised pH gradient (IPG) strip (24 cm, pH 3-10 NL; GE Healthcare). Isoelectric focusing (IEF) was performed following the manufacturer's instructions with an Ettan IPGphor II System (GE Healthcare). After IEF, the proteins were reduced with equilibration buffer containing 2% w/v dithiothreitol (DTT) for 15 min. The proteins were then alkylated with equilibration buffer containing 2.5% w/v iodoacetamide for 15 min. The second dimension of electrophoresis was conducted using an Ettan DALT-VI system (GE Healthcare).

Gel image acquisition and analysis. The Typhoon 9400 imager (GE Healthcare) was used to visualise labelled proteins at excitation/emission wavelengths of 488/520 nm (cy2), 532/580 nm (cy3), and 633/670 nm (cy5). DeCyder software (V6.0, GE Healthcare) was used to analyse the 2-D DIGE gel images. Protein spots with significant differences in abundance (increased or decreased more than > 1.5-fold) were selected for further analysis.

Spot picking and enzymatic digestion. Using an Ettan Spot Handling Workstation (GE Healthcare), the selected protein spots were automatically picked up and treated. The protein spots were washed with $25 \text{ mM NH}_4\text{HCO}_3$ and 50% acetonitrile and were then digested with $8 \text{ ng } \mu l^{-1}$ trypsin solution (mass spectrometry grade; Promega, Madison, WI, USA) in $25 \text{ mM NH}_4\text{HCO}_3$ for 16 h at 37° C. After digestion, the tryptic peptides were extracted with 60% (v/v) acetonitrile and 0.1% (v/v) trifluoroacetic acid (TFA). Finally, the tryptic peptides were dissolved in 4 mg ml⁻¹ 4-hydroxy-alphacyanocinnamic acid matrix in 70% (v/v) acetonitrile and 0.1% (v/v) TFA and were spotted on the target plate.

MALDI-TOF/TOF MS analysis. In positive ion reflector mode, an Ultraflex III MALDI-TOF/TOF MS (Bruker, Karlsruhe, Baden-Württemberg, Germany) was used to perform MS analysis. Four of the most intense ion signals were automatically chosen as precursors for MS/MS acquisition, excluding the matrix ion signals and trypsin autolysis peaks. MASCOT Version 2.2 (Matrix Science, London, UK) and Biotools software (Version 3.2, Bruker) were used to search the MS/MS spectra combined peptide mass fingerprint in the NCBInr database. The following search criteria were used: Homo

sapiens, trypsin cleavage, oxidation of methionines allowed as variable modification, one missed cleavage allowed, carbamidomethylation as fias modification, peptide mass tolerance at 75 ppm, and fragment tolerance at 0.5 Da. A successfully identified protein had the following parameters: ion score confidence interval (CI%) for PMF and MS/MS data \geq 95%, distinct sequences identified in MS/MS analysis \geq 2 and peptide count (hit) \geq 4.

Differential proteins categorisation and network modelling. The differential proteins were assorted using the protein analysis through evolutionary relationships (PANTHER) system (http:// www.pantherdb.org). The PANTHER ontology is a highly controlled vocabulary categorised according to biological process, protein class, and molecular function.

To establish an interaction map among the differential proteins, we used Pathway Studio 5.0 software (Ariadne Genomics, Rockville, MD, USA), a text-mining tool that can construct protein interaction networks and pathways. It includes protein–protein interactions, pathway components, proteins and their cellular processes, and functional classes. In this study, we selected the shortest path analysis.

Western blotting. Western blotting was performed on the same set of samples used in the DIGE analysis. Briefly, 50 μ g pooled protein extracts of each group were fractionated by 10% SDS-PAGE and transferred to a PVDF membrane as previously described (Li *et al*, 2013). The signal was detected with an enhanced chemiluminescence (ECL) plus kit (Milipore, Bedford, MA, USA). Anti-manganese superoxide dismutase (anti-MnSOD) (1:2000), anti-heat shock protein B1 (anti-HspB1) (1:1000), anti-fatty acid binding protein 5 (anti-FABP5) (1:1000), and anti-glycerol-3-phosphate dehydrogenase (anti-GAPDH) (1:1000) were used. All of these antibodies were rabbit polyclonal antibodies and were purchased from Abcam, Inc. (Cambridge, MA, USA).

Immunohistochemistry. Immunohistochemistry staining was performed using various primary antibodies including anti-

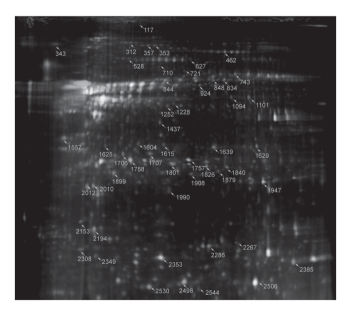


Figure 1. Representative DIGE gel image of differentially expressed proteins in primary early-stage cervical cancer with and without PLNM. Proteins extracted from the two groups were labelled with Cy3 (green, PLNM) and Cy5 (red, NPLNM). An internal standard (a mixture of PLNM and NPLNM CC, I–IIst samples) was labelled with Cy2 (blue). The yellow arrow indicates the significantly differentially expressed protein points in the PLNM and NPLNM groups. The number in the figure corresponds to the number shown in Table 2. The full colour version of this figure is available at *British Journal of Cancer* online.

MnSOD (1:600), anti-HspB1 (1:1200), and anti-FABP5 (1:600) as previously described (Pang *et al*, 2010). These antibodies are the same as those used in western blotting. In negative controls, primary antibodies were omitted. Two experienced pathologists assessed the immunostaining in a blinded fashion as described

Table 2. Differential proteins between early-stage cervical cancer with and without PLNM identified by MALDI-TOF/TOF MS

previously (Cheng *et al*, 2008). The results were scored by adding the staining intensity (0 = no, 1 = weak, 2 = moderate, 3 = strong staining) and the staining area (0 = no, 1 = <30%, 2 = between 30% and 60%, 3 = between 60 and 100% stained cells). For statistical analysis, the immunostaining scores (ranging from 0 to 6)

Master no.	Accession no.	Mr	PI	Mascot score	Coverage (%)	Fold change (PLNM/ NPLNM)	Protein name
117	P12110	109709	5.85	96	14%	2.19	COL6A2 Isoform 2C2 of Collagen alpha-2(VI) chain
312	P02679	50 0 92	5.7	68	10%	- 4.05	FGG Isoform Gamma-A of Fibrinogen gamma chain
343	Q05707	92740	4.70	98	18%	4.67	COL14A1 93 kDa protein
353	P02679	50 0 92	5.7	124	18%	- 6.04	FGG Isoform Gamma-A of Fibrinogen gamma chain
357	P02679	50 0 92	5.7	77	10%	- 8.02	FGG Isoform Gamma-A of Fibrinogen gamma chain
462	P02787	79294	6.81	359	46%	1.53	TF Serotransferrin
528	BOYJC5	26 956	4.68	70	26%	- 5.45	VIM Vimentin variant 4
527	Q9UBG3	53730	5.73	84	21%	3.68	CRNN Cornulin
710	B4E3Q9	101 217	5.35	86	15%	- 1.52	VCL cDNA FLJ59659, highly similar to Vinculin
721	Q96AX9	82 990	7.41	71	6%	- 2.49	MIB2 E3 ubiquitin-protein ligase MIB2 isoform 5
743	P02675	56 577	8.54	80	18%	- 3.2	FGB Fibrinogen beta chain
334	B4DRR0	58088	8.09	210	36%	- 3.53	KRT6A cDNA FLJ53910, highly similar to Keratin, type II cytoskeletal 6
344	P13647	62 568	7.59	335	40%	- 2.32	KRT5 Keratin, type II cytoskeletal 5
348	B4DRR0	58 088	8.09	368	47%	- 2.86	KRT6A cDNA FLJ53910, highly similar to Keratin, type II cytoskeletal 6
924	B4DRR0	58 088	8.09	388	53%	- 7.89	KRT6A cDNA FLJ53910, highly similar to Keratin, type II cytoskeletal 6
1094	B4E1D3	50 4 36	8.22	148	24%	- 5.97	FGB fibrinogen beta chain isoform 2 preproprotein
1101	B4E1D3	50 4 36	8.22	251	43%	- 5.66	FGB fibrinogen beta chain isoform 2 preproprotein
1228	B4DW52	38 950	5.19	84	13%	1.59	ACTB cDNA FLJ55253, highly similar to Actin, cytoplasmic 1
1252	B4E1D3	50 4 36	8.22	192	39%	- 2.02	FGB fibrinogen beta chain isoform 2 preproprotein
1437	P17706	41 363	5.98	69	10%	2.46	PTPN2 tyrosine-protein phosphatase non-receptor type 2
1557	P67936	28619	4.67	251	42%	2.16	TPM4 Isoform 1 of Tropomyosin alpha-4 chain
604	B4E3A4	40 1 1 6	5.23	169	21%	- 3.84	cDNA FLJ57283, highly similar to Actin, cytoplasmic 2
1615	P02679	50 0 92	5.7	109	17%	- 4.69	FGG Isoform Gamma-A of Fibrinogen gamma chain
1625	P30101	13739	6.78	131	33%	2.52	PDIA3 14 kDa protein
1629	P04075	39851	8.3	63	11%	2.05	ALDOA Fructose-bisphosphate aldolase A
1639	E7EQV7	58718	6.6	333	28%	- 7.69	KRT6C Uncharacterized protein
1706	P14618	58 538	7.6	103	13%	2.89	PKM2 Isoform M1 of Pyruvate kinase isozymes M1/M2
1707	P49761	74267	9.94	73	8%	- 4.3	CLK3 Isoform 4 of Dual specificity protein kinase CLK3
1757	P04792	22 826	5.98	305	42%	15.22	HSPB1 Heat shock protein beta-1
1758	P60709	42 0 5 2	5.29	197	39%	2.06	ACTB Actin, cytoplasmic 1
1801	P13647	23 267	4.82	343	49%	- 3.88	KRT5 Keratin, type II cytoskeletal 5
1826	P02768	23 4 1 4	5.93	164	47%	1.6	ALB 23 kDa protein
1840	E7EUE8	56065	5.95	247	31%	- 3.51	KRT6A Uncharacterized protein
879	P04792	22 826	5.98	238	51%	5.11	HSPB1 Heat shock protein beta-1
1899	B4DRR0	58 0 8 8	8.09	251	29%	- 2.44	KRT6A cDNA FLJ53910, highly similar to Keratin, type II cytoskeletal 6
1908	E9PH29	26 107	7.04	74	14%	2.27	PRDX3 thioredoxin-dependent peroxide reductase, mitochondrial isoform b
1947	P04179	24 878	8.35	169	44%	2.57	SOD2 Superoxide dismutase [Mn], mitochondrial
1990	P09211	23 569	5.43	171	43%	2.27	GSTP1 Glutathione S-transferase P
2010	E7EUE8	56 065	5.95	216	23%	- 2.57	KRT6A Uncharacterized protein
2012	B4DNH8	2182	5.98	75	37%	- 1.53	cDNA FLJ59138, highly similar to Annexin A2
2153	B4DNG6	18212	4.99	90	44%	2.68	ANXA5 ANXA5 Uncharacterized protein
2194	G5E9R0	13 907	5.24	246	54%	1.51	ACTB 14 kDa protein
2267	P25398	14 905	6.81	85	25%	2.8	RPS12 40S ribosomal protein S12
2286	Q01469	15 497	6.6	84	57%	7.49	FABP5 Fatty acid-binding protein, epidermal
2308	P20674	16923	6.3	100	11%	3.46	COX5A Cytochrome c oxidase subunit 5A, mitochondrial
2349	P09382	15 048	5.34	311	62%	1.6	LGALS1 Galectin-1
2353	P06702	13 291	5.71	101	90%	- 2.47	S100A9 Protein S100-A9
2385	O94964	185 512	6.15	63	7%	2.84	KIAA0889 Isoform 2 of Uncharacterized protein KIAA0889
2498	P31949	11847	6.56	65	28%	1.51	S100A11 Protein S100-A11
2506	P05109	10885	6.51	194	67%	- 2.17	S100A8 Protein S100-A8
2530	Q9H628	7191	9.06	64	48%	1.53	RERGL 7 kDa protein
2544	F8VYY1	7837	9.81	62	31%	2.6	CDK4 Uncharacterized protein

Abbreviations: MALDI-TOF/TOF MS = matrix-assisted laser desorption/ionisation-time-of-flight mass spectrometry; Mr = relative molecular mass; NPLNM = no pelvic lymph node metastasis; PI = isoelectric point; PLNM = pelvic lymph node metastasis.

were evaluated and a cut-off was determined. The samples were divided accordingly into low- and high-staining groups. For HspB1, a staining score of 4 was defined as the cutoff. For MnSOD and FABP5, a staining score of 3 was defined as the cutoff.

Statistical analysis. Statistical analyses were performed using SPSS 13.0 (SPSS Inc., Chicago, IL, USA) and MedCalc statistical software (MedCalc Software, Mariakerke, Belgium). The relationships between the FABP5, HspB1, and MnSOD expression levels and clinicopathological factors were evaluated using the Pearson χ^2 test or Fisher's exact test. Logistic regression analyses (univariate and stepwise multivariate) were used to determine variable predictability of PLNM. Receiver operating characteristic (ROC) analysis was used to test the discriminatory power of measured markers. The Kaplan–Meier method was used to construct relapse-free probability curves and overall survival curves, and the curves were compared by the log-rank test. P < 0.05 was considered to be statistically significant.

RESULTS

Identification of differential proteins in primary early-stage cervical cancer with and without PLNM by proteomic analysis. Protein extracts from tissue homogenates were subjected to DIGE analysis. Figure 1 shows a typical DIGE gel image. After visual review, 52 differential protein spots with at least a 1.5-fold discrepancy between the two groups were selected for MALDI-TOF/TOF MS analysis. In total, 41 differentially expressed proteins (26 upregulated and 15 downregulated) were successfully identified (Table 2). Some proteins appeared as multiple spots in DIGE, likely due to their isoforms and/or modifications.

Differential proteins categorisation and network modelling. The differential proteins were classified according to biological processes, molecular functions, and protein classes using the PANTHER system (Figure 4). The proteins were classified into 23 groups based on protein classes, including signalling molecules (14%), calcium-binding proteins (8%), cell adhesion molecules (6%), oxidoreductases (4%), and so on. Most of the proteins participated in the metabolic (15%), cellular (14%), and cell communication processes (12%). The proteins were also involved in other biological processes, such as the immune responses, cell adhesion, the generation of energy, apoptosis and cellular component organisation. Regarding molecular function, the proteins were divided into five categories: (a) catalytic activity (34%), (b) binding (30%), (c) structural molecule activity (21%), (d) receptor activity (6%), and (e) motor activity (3%).

To explore the associations between differentially expressed proteins, we constructed a protein network with Pathway Studio

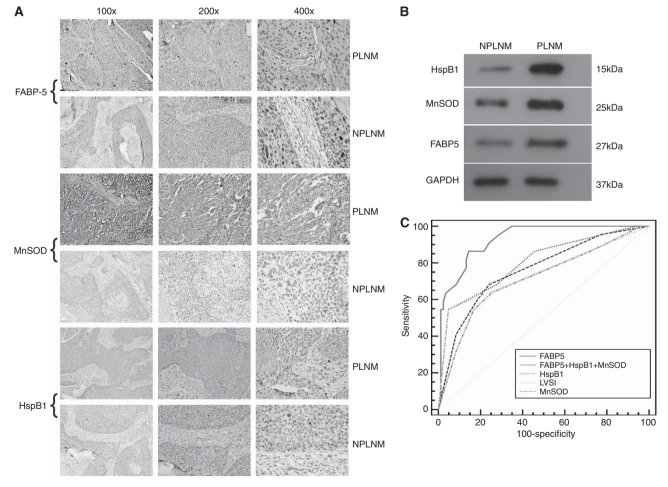


Figure 2. Validation of FABP5, HspB1, and MnSOD expression in early-stage cervical cancer with and without PLNM, and their discriminatory power as proteomic biomarkers. (A) Representative results of immunohistochemical staining of FABP5, HspB1, and MnSOD in early-stage cervical cancer with and without PLNM. (B) Representative results of western blotting of FABP5, HspB1, and MnSOD in early-stage cervical cancer with and without PLNM. (B) Representative results of western blotting of FABP5, HspB1, and MnSOD in early-stage cervical cancer with and without PLNM. GAPDH is used as an internal loading control. (C) ROC curves of FABP5, HspB1, MnSOD, and LVSI as markers to differentiate PLNM in early-stage cervical cancer (HspB1 AUC = 0.803; MnSOD AUC = 0.713; FABP5 AUC = 0.761; LVSI AUC = 0.749; the combination of FABP5, HspB1, and MnSOD AUC = 0.929).

5.0 software. Complex interactions between mitochondrial, nuclear, cytoplasmic, and extracellular proteins were present in the prediction network (Supplementary Figure S1). Further research is needed to explore the role of these proteins and their interaction pathways in the pathogenesis of PLNM in CC, I–IIst.

Validation of differential proteins by western blotting. To further confirm the proteomic results, western blotting was performed to examine the expression levels of FABP5, HspB1, and MnSOD in the same pooled protein extracts used in the DIGE analysis. The expression levels of the three proteins were higher in the PLNM group than in the NPLNM group (Figure 2B). This validation result was consistent with the DIGE analysis.

Validation of differential proteins by IHC. Immunohistochemical staining was performed to examine the expression levels of FABP5, HspB1, and MnSOD in an independent tissue cohort of primary CC, I–IIst. Semi-quantitative analysis showed that the PLNM group had significantly higher expression of FABP5, HspB1, and MnSOD than the NPLNM group (Table 3, Figure 2A). Next, the correlation of FABP5, HspB1, and MnSOD with clinicopathological factors was analysed (Table 3). HspB1 was significantly correlated with PLNM (P<0.001). MnSOD was significantly correlated with PLNM (P=0.006), lymphovascular space invasion (LVSI) (P=0.01), and stromal invasion (P=0.002). FABP5 was significantly correlated with PLNM (P<0.001) and LVSI (P=0.001). No significant correlations were identified between the expression of the three proteins and other clinicopathological characteristics, including age, tumour size, tumour differentiation, International Federation of Gynaecology and Obstetrics (FIGO) stage, parametrial invasion, and vaginal invasion.

Correlations of FABP5, HspB1, and MnSOD expression with PLNM. Univariate logistic regression analysis revealed that tumour size (P = 0.003), LVSI (P < 0.001), stromal invasion (P = 0.006), MnSOD (P = 0.011), HspB1 (P = 0.001), and FABP5 (P < 0.001) were significant predictors of PLNM (Table 4). Stepwise multivariate logistic regression analysis indicated that LVSI (P = 0.001), HspB1 (P = 0.013), and FABP5 (P = 0.018) were independent predictors of PLNM in patients with CC, I–IIst (Table 4).

	Ν		HspB1		MnSOD			FABP5		
	105	Low (0–4) 67	High (5–6) 38	<i>P</i> -value	Low (0–3) 69	High (4–6) 36	<i>P</i> -value	Low (0–3) 70	High (4–6) 35	<i>P</i> -value
Age (years)	-	-	-	0.761	-	-	0.918	-	-	0.121
≼35	21	14	7	-	14	7	-	17	4	-
> 35	84	53	31	-	55	29	-	53	31	-
FIGO stage	-	-		0.205	-	-	0.745	-	-	0.076
la2	12	9	3	-	10	2	-	12	0	-
lb1	37	28	9	-	24	13	_	24	13	-
lb2	22	13	9	-	14	8	-	15	7	-
lla1	16	8	8	-	10	6	-	10	6	-
lla2	18	9	9	-	11	7	-	9	9	-
Tumour size (cm)	-	-	-	0.059	-	-	0.586	-	-	0.256
≼4	65	46	19	-	44	21	-	46	19	-
>4	40	21	19	-	25	15	-	24	16	-
PLNM	-	-	-	< 0.001	-	-	< 0.001	-	-	< 0.001
Positive	22	7	15	-	10	12	-	6	16	-
Negative	83	60	23	-	58	25	-	64	19	-
LVSI	-	-	-	0.287	-	-	0.01	-	-	0.001
Positive	16	8	8	_	6	10	_	5	11	_
Negative	89	59	30	-	63	26	-	65	24	_
Differentiation	-	-	-	0.366ª	-	-	0.555ª	-	-	1ª
G1	6	4	2	-	3	3	-	4	2	-
G2	37	27	10	-	26	11	-	25	12	-
G3	62	36	26	-	40	22	-	41	21	-
Stomal invasion	-	-	-	-	-	0.002	-	-	-	0.096
≤1/2	48	35	13	-	39	9	-	36	12	_
> 1/2	57	32	25	-	29	28	-	34	23	
Parametrial invasion	-	-	-	1	-	-	1	-	-	0.064
Positive	9	6	3	-	6	3	-	3	6	-
Negative	96	61	35	-	63	33	-	67	29	-
Vaginal invasion	-	-	-	0.948	-	-	0.184	-	-	0.37
Positive	19	12	7	-	10	9	_	11	8	_
Negative	86	55	31	_	59	27	_	59	27	_

Table 3. Associations of FABP5, HspB1, and MnSOD expression levels with clinicopathological factors in early-stage cervical cancer

Abbreviations: FIGO = the International Federation of Gynecology and Obstetrics; LVSI = lymphovascular space invasion; PLNM = pelvic lymph node metastasis ^aP value from Fisher's exact test.

Table 4. Univariate and multivariate logistic regression analyses of factors
associated with pelvic LN metastasis

Variables	HR (95% CI)	P-value
Univariate analysis, n=	105	
Age (years)	0.976 (0.919–1.037)	0.428
Tumour size (cm)	1.768 (1.219–2.564)	0.003
Differentiation		0.963
G1 (reference)	1	
G2	1.379 (0.140–13.556)	0.783
G3	1.327 (0.142–12.366)	0.804
FIGO stage		0.116
l (reference)	1	
ll	2.174 (0.826–5.718)	
LVSI		< 0.001
Negative (reference)	1	
Positive	23.700 (6.402–87.741)	
Stromal invasion		0.006
≤1/2 (reference)	1	
> 1/2	5.077 (1.582–16.293)	
Parametrial invasion		0.348
Negative (reference)	1	
Positive	2.026 (0.464–8.849)	
Vaginal invasion		0.214
Negative (reference)	1	
Positive	2.019 (0.666–6.123)	
MnSOD		0.011
Low (reference)	1	
High	3.551 (1.342–9.399)	
HspB1		0.001
Low (reference)	1	
High	5.59 (2.020–15.468)	
FABP5		< 0.001
Low (reference)	1	
High	6.750 (2.413–18.880)	
Multivariate analysis ^a , i	n = 105	
LVSI		0.001
Negative (reference)	1	
Positive	17.512 (3.181–96.420)	
Stromal invasion		0.074
≤1/2 (reference)	1	
> 1/2	4.038 (0.872–18.701)	
HspB1		0.013
Low (reference)	1	
High	6.379 (1.480–27.482)	
FABP5		0.018
Low (reference)	1	
High	5.829 (1.355-25.075)	
MnSOD		0.053
Low (reference)	1	
High	4.053 (0.980–16.769)	

Gynaecology and Obstetrics; HR = hazard ratio; LVSI = lymphovascular space invasion. ^aFor the stepwise multivariate analysis, backward LR method was used to drop insignificant variables. Variables entered for analysis were the following: age, tumour size, differentiation, FIGO stage, LVSI, stomal invasion, parametrial invasion, vaginal invasion, FABP5, HspB1, and MnSOD. To assess the discriminatory power for the three proteins to predicting PLNM in CC, I–IIst, ROC curves were constructed (Figure 2C). The three proteins were used as covariates to create a logistic regression model with PLNM as the dependent variable. As a result, we could further evaluate the feasibility of the combination of FABP5, HspB1, and MnSOD to predicting PLNM. The area under the ROC curve (AUC), sensitivity, specificity, positive predictive value (PPV), and negative predictive value (NPV) for FABP5, HspB1, MnSOD, and the combination of the three proteins are shown in Table 5. The cut-points (by maximising the Youden index) of FABP5, HspB1, MnSOD, and the combination of the three proteins were 3, 5, 3, and 0.2448 (predicted probability from the logistic regression model of combining the three proteins), respectively. When all three proteins were considered, the discriminatory power was strong (AUC = 0.929).

Correlations of FABP5, HspB1, and MnSOD expression with postoperative relapse and survival. At the end of the study, 34 patients (32.38%) experienced disease recurrence, 31 patients (29.53%) had died of CC, I–IIst, and 74 patients (70.48%) were still alive. The relapse-free probability curves indicated that the relapse rate increased significantly with upregulation of FABP5, HspB1, and MnSOD (Figure 3). The survival curves indicated that the overall survival rate decreased significantly with upregulation of FABP5, HspB1, and MnSOD (Figure 3).

DISCUSSION

PLNM is an important prognostic factor in CC, I–IIst, and PLN status is crucial to decision-making about personalised treatment of CC, I–IIst. Currently, it is difficult to diagnose PLNM accurately and efficiently in usual assessment methods used for CC, I–IIst before first treatment. Therefore, reliable biomarkers for clinical diagnosis and better insights into the pathogenesis of PLNM in CC, I–IIst are needed. Our results suggest that FABP5, HspB1, and MnSOD may predict PLNM in CC, I–IIst patients and may also provide information useful to help further elucidate the molecular mechanisms underlying PLNM in CC, I–IIst.

In this study, first, we identified 41 differential proteins between primary early-stage cervical squamous cell cancer tissue with and without PLNM using DIGE-based proteomics. These proteins participate in processes of malignancy progression and metastasis, such as oxidative stress, energy metabolism, cytoskeleton reorganisation, and signal transduction. Second, three differential proteins (FABP5, HspB1, and MnSOD) were validated by western blotting and IHC, indicating that the proteins identified by the proteomic approach were actually differentially expressed proteins. Finally, correlations of the expression levels of the three proteins with clinicopathological factors and clinical outcomes were evaluated in an independent tissue cohort. Our data indicated that FABP5, HspB1, and MnSOD were significantly correlated with PLNM and poor prognosis.

FABP5 is a member of the fatty acid binding protein family with functions in fatty acid uptake, metabolism, and transportation. FABP5 is highly expressed in several types of tumours, such as prostate (Adamson *et al*, 2003), oral (Fang *et al*, 2010), and hepatocellular carcinomas (Fujii *et al*, 2005). In addition, accumulative studies have shown that upregulation of FABP5 is correlated with tumour metastasis (Fang *et al*, 2010; Jeong *et al*, 2012). Previous studies reported that overexpression of FABP5 induces metastasis by upregulating vascular endothelial growth factor (VEGF) (Jing *et al*, 2001) and matrix metalloproteinase 9 (MMP-9) (Fang *et al*, 2010), which has important roles in the metastatic progression. To our knowledge, there are currently no published reports on the expression of FABP5 in cervical cancer. This study, for the first time, showed that FABP5 was higher expressed in PLNM CC, I–IIst

Proteomic markers for pelvic lymph node metastasis

Factors	Sensitivity (95% CI) (%)	Specificiy (95% Cl) (%)	PPV (95% CI) (%)	NPV (95% CI) (%)	AUC (95% CI)
HspB1	54.55 (32.2–75.6)	95.18 (88.1–98.7)	75 (47.6–92.7)	88.8 (80.3–94.5)	0.803 (0.714–0.874
FABP5	68.18 (45.1–86.1)	75.9 (65.3–84.6)	42.9 (26.3–60.6)	90 (80.5–95.9)	0.761 (0.668–0.839
MnSOD	63.64 (40.7–82.8)	74.7 (64.0–83.6)	40 (23.9–57.9)	88.6 (78.7–94.9)	0.713 (0.616–0.797
HspB1 + FABP5 + MnSOD	86.36 (65.1–97.1)	85.54 (76.1–92.3)	61.3 (42.2–78.2)	95.9 (88.5–99.2)	0.929 (0.862-0.970

Abbreviations: 95% CI=95% confidence interval; AUC=the area under the receiver operating characteristic curve; NPV=negative predictive value; PPV=positive predictive value.

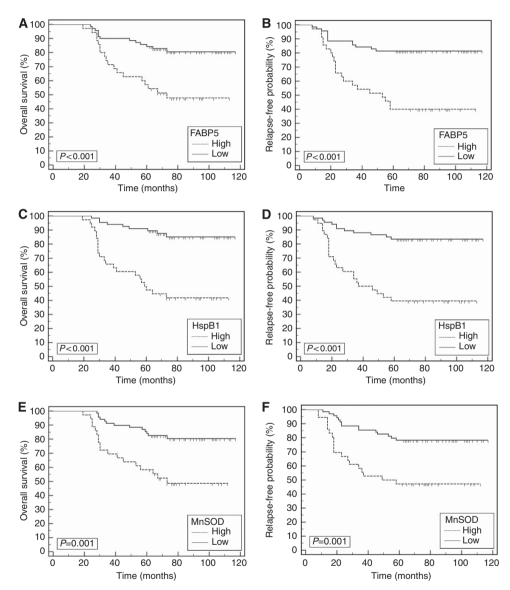


Figure 3. Kaplan–Meier survival plots for early-stage cervical cancer according to the expression levels of FABP5, HspB1, and MnSOD. (A) FABP5 expression and overall survival (log-rank test, P < 0.001). (B) FABP5 expression and relapse-free probability (log-rank test, P < 0.001). (C) HspB1 expression and overall survival (log-rank test, P < 0.001). (D) HspB1 expression and relapse-free probability (log-rank test, P < 0.001). (E) MnSOD expression and overall survival (log-rank test, P = 0.001). (F) MnSOD expression and relapse-free probability (log-rank test, P = 0.001).

tissue than NPLNM CC, I–IIst tissue and that its upregulation was significantly correlated with PLNM (P<0.001) and poor prognosis (P<0.001). Moreover, the results showed that FABP5 was an independent predictor (P=0.018) for CC, I–IIst patients with PLNM in multivariate analysis. Thus, FABP5 may be a biomarker for predicting PLNM of CC, I–IIst. It seems that FABP5 upregulation has a role in the development of PLNM.

HspB1 acts as a molecular chaperone whose main function is to prevent the aggregation of misfolded proteins and to restore denatured proteins. Its functions can be triggered by different types of stress such as inflammation, oxidative stress, and malignant transformation (Ciocca and Calderwood, 2005). HspB1 is overexpressed in several types of malignancies (Huang *et al*, 2010; Yu *et al*, 2010) and has been correlated with poor prognosis (Yu *et al*, 2010).

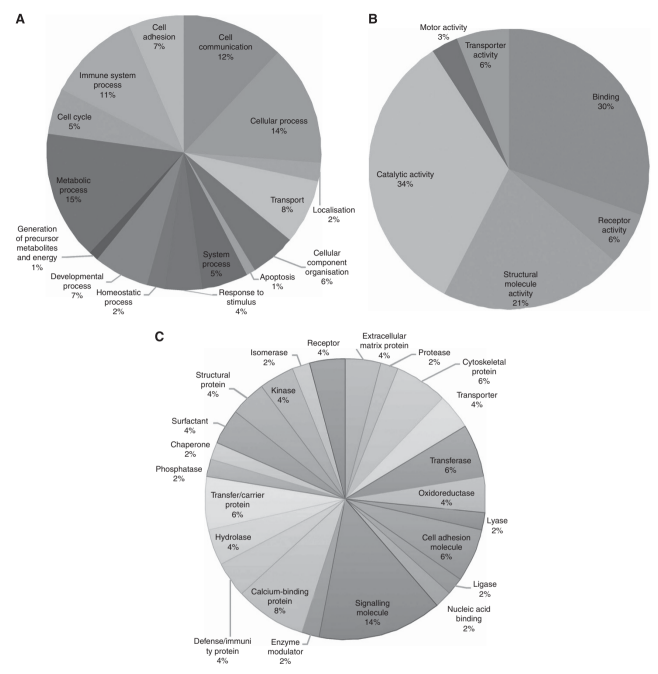


Figure 4. Classification analysis of differentially expressed proteins in early-stage cervical cancer with and without PLNM. (A) Classification of the differential proteins according to biological process. (B) Classification of the differential proteins according to molecular function. (C) Classification of the differential proteins according to protein class.

Moreover, it has been reported that upregulation of HspB1 is associated with tumour metastasis (Nagaraja *et al*, 2012). In cervical cancer, the correlation of HspB1 overexpression with carcinogenesis has been shown (Ono *et al*, 2009; Lomnytska *et al*, 2011), but has not yet been associated with PLNM. Our study revealed a significant association of HspB1 expression with PLNM (P < 0.001) and that HspB1 was an independent predictor of PLNM in multivariate analysis (P = 0.013). Thus, our data suggest that HspB1 may be a biomarker of PLNM for CC, I–IIst and that upregulation of HspB1 may be involved in the development of PLNM in CC, I–IIst. Additionally, our data showed that patients with HspB1 upregulation had a poor prognosis. However, a previous study showed that cervical cancer patients (FIGO stage Ib1-IIIb) with low (n = 2) or high HspB1 (n = 4) expression levels had poorer prognoses than patients with moderate protein expression levels (n=8) (Lomnytska *et al*, 2011). The different results between these two studies may be due to different sample cohorts and sample quantities (105 samples versus 14 samples).

MnSOD (also called SOD₂) is localised to the mitochondria and can catalyse the superoxide radical (O_2^-) to H_2O_2 and hydroperoxides. H_2O_2 and hydro-peroxides then promote oxidative stress, which contributes to tumour progression and metastasis (Sotgia *et al*, 2011). MnSOD has attracted the attention of many investigators because of its overexpression in various types of human malignancies such as colorectal (Nozoe *et al*, 2003), gastric (Malafa *et al*, 2000), and oral squamous cell carcinomas (Liu *et al*, 2010), and it is often associated with metastatic progression (Malafa *et al*, 2000; Liu *et al*, 2010). Additionally, some studies have shown that MnSOD-dependent upregulation of MMPs (including MMP-1 and MMP-9) may contribute to the increased invasive and metastatic capacities of tumours (Nelson et al, 2003). In cervical cancer, a correlation of MnSOD upregulation with different stages of cancer was shown (Termini et al, 2011), but there is no report of studying its association with PLNM. Our study showed a significant correlation between MnSOD expression and PLNM (P = 0.006), which is consistent with previous studies in other tumours (Nozoe et al, 2003; Liu et al, 2010). Additionally, we found that MnSOD was a significant predictor of PLNM (P = 0.011) in the univariate analysis. In the multivariate analysis, MnSOD trended towards being an independent predictor of PLNM (P=0.053), but the differences did not reach statistical significance. This result may be due to the study's relatively small sample size, which affects statistical power. Therefore, our data suggest that MnSOD may be a biomarker for PLNM of CC, I-IIst, and upregulation of MnSOD may support the development of PLNM in CC, I-IIst.

Surgery as a treatment option for CC, I-IIst can offer hope of fertility or function preservation and avoid the damage of ovary and vagina function. Therefore, surgical treatment is a much better option for patients without PLNM and with desire for fertility or function preservation. For patients with PLNM and with desire for function preservation, surgical treatment also can be considered because the normal ovaries can be transposed during the surgery, so that the ovaries will not be damaged by postoperative radiotherapy. For patients with PLNM and without desire for function preservation, chemoradiotherapy should be recommended as the first treatment to avoid the morbidity caused by combining surgery and chemoradiotherapy. In the present study, the combination of FABP5, HspB1, and MnSOD had a good discrimination power (AUC = 0.929), high sensitivity and specificity (86.36% and 85.54%, respectively), high NPV (95.7%), and moderate PPV (61.3%) for the identification of CC, I-IIst patients with PLNM. These test characteristics are based on the decision to use 0.2448 (by maximising the Youden index) as a cut-point. Because of the high NPV, these proteins can be helpful in the clinical decision-making process. If a patient has a NPLNM diagnosed by the combination of these proteins (cutpoint = 0.2448, NPV = 95.7%), then the clinician can plan surgery (fertility preservation or function preservation) for patients, which depends on the patient's desire. If a patient has a PLNM diagnosed by the combination of these proteins (cut-point = 0.654, by maximising PPV to 92.3%), then the clinician can recommend chemoradiotherapy or function preservation surgery (plus postoperative adjuvant therapy) for patients, which depends on the patient's desire. As primary cervical cancer tissues can be easily obtained by transvaginal biopsy for detecting the expression levels of these proteins before treatment, these proteins can be applied to evaluate PLNM for CC, I-IIst patients before the first management.

Consistent with previous studies (Pallavi *et al*, 2012), our results revealed that LVSI was also an independent predictor of PLNM. When comparing PLNM prediction capacities, we observed that the AUCs of HspB1 (0.803) and FABP5 (0.761) were higher than the AUC of LVSI (0.749). Additionally, when combining FABP5, HspB1, and MnSOD, we found that the discriminatory power was significantly improved (Figure 4). The AUC of LVSI (0.749) was much lower than that of the combination of the three proteins (0.929). These findings suggest that FABP5, HspB1, and MnSOD may be promising biomarker candidates for identifying CC, I–IIst patients with PLNM.

Previous studies have shown that stromal invasion is an important predictor of PLNM in cervical cancer (Gauthier *et al*, 1985; Ayhan *et al*, 1991). In our study, univariate analysis showed that stromal invasion was a significant predictor of PLNM (P = 0.006). However, in multivariate analysis, it was not an independent predictor of PLNM (P = 0.074). This may be due to the small sample size, which is a main limitation of our study.

The reason for the small sample size was that only patients without preoperative adjuvant therapy and with complete follow-up data were eligible to be enrolled in our study. Moreover, the incidence of PLNM in CC, I–IIst is low. Therefore, the sample size of this study was relatively small. Another limitation of the present study is that only one pathological type of cervical cancer (squamous cell cancer) was studied. This was because of the need for homogenous samples in proteomic analysis. On the other hand, the prevalence of other pathological types of cervical cancer is low so that we could not collect enough tumour samples for the study. Thus future studies with larger sample sizes and different pathological types need to be performed.

In summary, we identified 41 differentially expressed proteins between primary CC, I–IIst tissues with and without PLNM using DIGE-based proteomics. We further found that three differentially expressed proteins (FABP5, HspB1, and MnSOD) are potential biomarkers for PLNM of CC, I–IIst, and may contribute to the pathogenesis of PLNM. These findings may help clinicians to design optimal individualised treatment plans before first treatment.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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