Gene Expression Profiling Identified High-mobility GroupAT-hook 2 (HMGA2) as Being Frequently Upregulated in Esophageal Squamous Cell Carcinoma

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ABSTRACT

Background: Oesophageal cancer is one of the most deadly malignancies worldwide and oesophageal squamous cell carcinoma (ESCC) is the most frequent type.

Methods: We identified up-regulated genes from gene expression profiles of HKESC-4 cell line, its parental tumor tissues, non-tumoralesophageal epithelia and lymph nodes with metastatic carcinoma using Human Genome U133 plus 2.0 microarray.

Results: Four genes [High-mobility group AT-hook 2 *(HMGA2)*, paternally expressed 10 *(PEG10)*, SH3 and multiple ankyrin repeat domains 2*(SHANK2)* and WNT1 inducible signaling pathway protein 3*(WISP3)*] were selected for further validation with real-time quantitative polymerase chain reaction (PCR) in a panel of ESCC cell lines and clinical specimens. *HMGA2* was found to be overexpressed in the panel of ESCC cell lines tested. By using immunohistochemistry, HMGA2 was found to be up-regulated in 70% of ESCC tissues (21 out of 30 cases).

Conclusion: This study demonstrates successful use of gene microarray to identify and reveal HMGA2 as a novel and consistently overexpressed gene in ESCC cell lines and clinical samples.

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1. Introduction

Esophageal cancer ranks fifth as the most common cause of cancer-related deaths in men worldwide, causing about 400,000 deaths annually[1]. The incident rate is higher in Southern and Eastern Africa and Eastern Asia when compared to Western and Middle Africa and Central America[1]. This cancer comprises two major types, namely esophageal squamous cell carcinoma (ESCC) and adenocarcinoma, the former is more common and contributes to about 90% of cases in high-risk regions [1]. Patients with ESCC usually have poor prognosis largely because of late diagnosis of the disease[2]. Despite advances in surgical techniques combined with various treatment modalities, such as radiotherapy and chemotherapy, the overall 5-year survival rate remains at20-30%[3]. To alleviate this clinical situation, the development of new treatment modalities, diagnostic technologies and preventative measures is required, which cannot be accomplished without understanding the underlying mechanisms of esophageal carcinogenesis. The development of gene microarray technology allows comprehensive comparison of gene expression profiles in various pathophysiological processes, such as enabling the comparison of gene profiles between cancer and normal conditions. In the present study, we took advantage of this technology to identify differentially expressed genes in cancerous and non-cancerous conditions in esophagus, followed by further validation for the involvement of these genes in ESCC.

2. Materials and Methods

HKESC-4 cell line and clinical specimens for microarray.HKESC-4, a human ESCC cell line of Chinese origin, was established previously in our laboratory and the culture conditionsfor this cell line were as described elsewhere [4]. Cultured HKESC-4 cells were harvested at 80% confluency at passage 30 for extracting RNA. Parental tumoral tissue (T), from which HKESC-4 cell line was derived, and its corresponding non-tumoral epithelium (N) and lymph node containing metastatic carcinoma (LN) were isolated during esophagectomy and snap-frozen until their use for RNA extraction. The extracted RNA of the cell line (HKESC-4) and clinical specimens (T, N and LN) was subjected to gene microarray. Consent regarding the use of clinical specimens for this study was obtained from Institutional Review Board of The University of Hong Kong/Hospital Authority Hong Kong West Cluster (HKU/HA HKW IRB).

Clinical tissues and ESCC cell lines for quantitative real time polymerase chain reaction (*qPCR*). In addition to the clinical specimens for gene microarray as mentioned above, three non-tumoral tissues were obtained from esophageal epithelium at least 5 cm away from the tumorfrom other patients during surgical resection (Non-T). Apart from HKESC-4 cell line, HKESC-1, HKESC-2, HKESC-3 and SLMT-1 cell lines were included and used as described elsewhere [4-6]. These cells were cultured in minimal essential medium (Invitrogen,

Carlsbad, CA, USA) supplemented with 10% fetal bovine serum (Invitrogen) and 1% penicillin-streptomycin. Cell cultures were maintained in a humidified incubator at 37° C containing 5% carbon dioxide (CO₂). Monolayer cells at 80% confluency were harvested and used for RNA extraction.

RNA extraction for qPCR.RNA extraction was performed as described [7]. In brief, 1 ml TRIzol reagent (Invitrogen) was used to lyse tissues or cells for RNA extraction. Chloroform was used for phase separation. After centrifugation, the upper aqueous phase with RNA was collected and transferred to RNase-free tubes containing 0.5 ml isopropyl alcohol for RNA precipitation. RNA pellets were then washed with 75% RNase-free ethanol. Finally, RNA pellets were dissolved in RNase-free water. DNase I digestion was performed before the concentration of RNA was determined by measuring its absorbance at 260 nm and A₂₆₀:A₂₈₀ ratio

Gene microarray.RNA of HKESC-4 cells and clinical specimens were subjected to gene expression profiling using GeneChip Human Genome U133 Plus 2.0 Array (Affymetrix, Santa Clara, CA, USA). This GeneChip enables the analysis of over 47,000 human transcripts and variants. The whole procedure of RNA quality control, microarray labeling,GeneChip hybridization and data acquisition was performed at the Genome Research Centre, The University of Hong Kong, Hong Kong, PRC under standardized condition. The statistical analysis to identify differentially expressed genes was performed using MicroArray Suite software (Affymetrix).

Synthesis of complementary DNA (cDNA).RNA was reverse-transcribed into cDNAusing High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Invitrogen), according to the manufacturer's protocol. Briefly, DNase-treated RNA was diluted with RNase-free water to reach a final concentration of 250ng in 10µl. Diluted RNA was then mixed with 2µl 10X RT buffer, 0.8 µl 25X dNTP Mix (100 mM), 2µl10X RT Random Primers, 1µlMultiScribe Reverse Transcriptase (50 U/µl), 1 µlRNase Inhibitor and 3.2 µlnuclease-free water. The reaction was then incubated at 25°C for 10 minutes, followed by 37°C for 2 hours and 85°C for 5 seconds to inactivate the activities of the reverse transcriptase and to completely denature the template.

qPCR. The procedure for qPCR was followed as described elsewhere [8, 9]. In brief,qPCR was performed usingcDNA of each sample, gene-specific primers (Table 1) and Platinum SYBR Green qPCRSuperMix-UDG (Invitrogen), according to the protocol from the manufacturer. The reactions were run for 50 cycles at 94°C for 10 minutes, 95°C for 90 seconds and 72°C for 90 seconds in an ABI PRISM 7700 Sequence Detector (Applied Biosystems).Cycle threshold (C_T)values of each reaction were obtained using Sequence Detection System (SDS) Software Version 1.9.1. For each reaction, the expression of each

gene was normalized against the expression of the housekeeping gene \Box -actin. The relative expression of each gene was calculated based on a comparative C_T equation and is presented as the value of relative intensity.

Statistical analysis for qPCR data. Statistical analysis was performed as described elsewhere [10]. One-way ANOVA followed by Duncan's multiple range testwas used to determine statistical significance. Each sample was run in triplicates. The relative value is presented as the mean ± standard error of the mean (S.E.M.). *p*-Values less than 0.05 were considered statistically significant. Statistical analysis was performed with Statistical Package for the Social Sciences version 17 (IBM, New York, USA).

Immunohistochemistry. Immunohistochemistry was performed using the avidin-biotin method[11, 12].Five-micrometer paraffin sections were prepared on gelatin-coated glass slides. Sections were preheated at 60°C for 20 minutes, deparaffinized in xylene and rehydrated through graded alcohol. Antigen retrieval was carried out by heating the sections in 0.2M citrate buffer (pH6) in a microwave oven at 95°Cfor 5 minutes. After cooling for 30 minutes, the sections were treated with 0.3% hydrogen peroxide at room temperature for 30 minutes to block endogenous peroxidase activity. Non-specific binding sites on sections were blocked with 1XTBS with 2% bovine serum albumin (BSA; Sigma-Aldrich, St Louis, MO, USA) and 2% normal goat serum (Invitrogen) at room temperature for 30 minutes. HMGA2specific rabbit polyclonal antibody (1:100)(Santa Cruz Biotechnology, Santa Cruz, CA, USA) was applied to the sections and the reaction was incubated at room temperature for 30 minutes. After washing three times with 1X TBS for 5 minutes each, the sections were incubated with biotinylatedanti-rabbit secondary antibody (EnVision Systems; Dako, Glostrup, Denmark) at room temperature for 30 minutes. To visualize the signals, the sections were washed and stained with avidin-biotin complex and 3,3'-diaminobenzene (DAB) at room temperature.Lastly, the sections were counterstained with hematoxylin. The expressions of HMGA2 were evaluated by Dr. AK Lam, a qualified pathologist, under light microscope.

3. Results

Identification of differentially expressed genes using gene microarray. Gene microarray analysis revealed 3,081 genes, 4,027 genes and 3,590 genes having more than 2-fold induction in T, LN and HKESC-4 cells when compared to N. For down-regulated genes with less than 2-fold difference, 4,808 genes, 6,052 genes and 5,846 genes were found in T, LN and HKESC-4 cells in comparison with N. For those up-regulated genes in T, LN and HKESC-4, 43 of them had more than 10-fold induction (Table 2). Among them, 34 genes (*CAMK2A*, *CART1*, *CCNA1*, *COCH*, *FLJ33516*, *FMN2*, *FOXD1*, *FOXG1B*, *GPR*, *HMGA2*, *HOXC10*, *HOXC11*, *HTR2C*, *IMP-3*, *LOC163782*, *MAGEB2*, *MFAP2*, *MGC17986*,

MGC27005, *NBEA*, *NKX2-2*, *PCDHB5*, *PEG10*, *PFN2*, *POPDC3*, *PPFIA1*, *PRAME*, *SAGE1*, *SHANK2*, *SIX1*, *SLCO1B3*, *SYT1*, *TP53TG3* and *WISP3*) were first identified to be overexpressed in ESCC, while 13 of them (*DKK1*,*EGFR*, *EMS1*, *GAL*, *HCG4*, *LAMC2*, *MAGEA1*, *MAGEA4*, *MAGEA11*, *MMP13*, *PTHLH*, *ZIC* and *ZNF595*) have previously been reported to be overexpressed in ESCC (Table 3). Among the 34 newly identified genes, *HMGA2*, *PEG10*, *SHANK2* and *WISP3*were further selected for validation due to their potential involvement in tumorigenesis based on literature search.

Gene expression of HMGA2.High expression of *HMGA2* in N, T, LN and HKESC-4 observed in gene microarray was confirmed using qPCR, such that the relative intensities of *HMGA2* expression in N, T, LN and HKESC-4 obtained using these two methods were comparable (fold change by gene microarray: 1, 31, 63 and 18*versus*1, 26, 49 and 33 by qPCR). Moreover, increased gene expression of *HMGA2*was also detected in four ESCC cell lines (HKESC-1, HKESC-2, HKESC-3 and SLMT-1). Significantly higher gene expression on average of 20-fold of *HMGA2*was noted in ESCC cell lines when compared to N and Non-T (Figure 1A).

Gene expression of PEG10.An over-expression of PEG10 in ESCC observed in gene microarray was confirmed using qPCR. Therelative intensities of PEG10 in N, T, LN and HKESC-4 cells were 1, 20, 28 and 28 by gene microarray, while their relative intensities were 1, 45, 148 and 269 byqPCR, respectively. However, no significant difference in the average gene expression of PEG10 was detected in tested ESCC cell lines when compared to N and Non-T, despite their having an average of 100-fold induction in gene expression of PEG10 (Figure 1B).

Gene expression of SHANK2

High gene expression of *SHANK2* in ESCC observed in gene microarray was confirmed using qPCR. The relative intensities of *SHANK2* in N, T, LN and HKESC-4 cells were 1, 30, 20 and 20by gene microarray and 1, 21, 11 and 19 byqPCR, respectively. No significant difference in the gene expression of *SHANK2* was detected in ESCC cells when compared to N (Figure 1C).

Gene expression of WISP3.High gene expression of *WISP3* in ESCC detected using gene microarray was confirmed using qPCR. The relative intensities of gene expression of *WISP3* in N, T, LN and HKESC-4 cells were 1, 37, 53 and 26by gene microarray and 1, 152, 117 and 117byqPCR, respectively. A significant increase in the gene expression of *WISP3* was detected in HKESC-2 cells. An average of 100-fold higher gene expression of *WISP3* was found when compared to N and Non-T (Figure 1D).

Protein expression of HMGA2 in ESCC tissues.Immunohistochemical data show the localization of HMGA2 in the nuclei of the ESCC tissues(Figure 2). Overexpression of HMGA2 was found in 70% ESCC tissues (21 out of 30 cases) when the expression of HMGA2 was examined in 30 pairs of tumoral tissues and adjacent non-tumoral tissues. No detectable level of HMGA2 expression was observed in the non-tumoral tissues(data not shown).

4. Discussion

In the present study, to our knowledge, the gene microarray and real-time qPCR analysis showed the presence of *HGMA2*mRNA expression in human esophageal tissues for the first time. *HGMA2* mRNA overexpression was detected in ESCC cell lines compared with the corresponding morphologically non-tumoralesophageal epithelial tissues. The current findings showed that overexpression of *HMGA2* gene appearsto be a consistent feature in ESCC. The protein expression of HMGA2 was further validated in ESCC specimens by immunohistochemistry. Liu *et al.*have shown an elevated level of HMGA2 protein in ESCC tissues by comparing 150 pairs oftumor and adjacent non-tumor tissues in patients[13], for which this data are in line with our findings reported here. The immunostaining analysis showed that HMGA2 protein expression was localized in the nuclei of the ESCC cells. The majority of ESCC cases (21/30, 70%) were found to have significantly enhanced expression of HMGA2 compared with morphologically normal esophageal epithelium.

HMGA2 belongs to the HMGA family, which also contains two other members *HMGA1A* and *HMGA1B*. HMGA protein family members are small nuclear proteins. A prominent feature of the HMGA family is the three DNA-binding domains termed AT-hooks at the *N*-terminal region that bind the minor groove of AT-rich DNA sequences. These proteins play key roles in chromatin architecture and gene control by serving as generalized chromatin effectors, either enhancing or suppressing the ability oftranscriptional factors in the process of transcriptional regulation. HGMA2 expression was found to be restricted during embryogenesis, whereas it is absent or has low expression in normal adult tissues[14]. However, overexpression of HGMA2 has been reported in various types ofhuman cancer including of the pituitary [15], oral cavity [16], lung [17], breast [18], pancreas [19], and nerves[20]. In addition, HMGA2 protein was reported to be ectopically expressed at the invasive front of oral carcinomas and had a significant impact on tumor progression and patient survival [16]. Similarly, HMGA proteins were found to be expressed in lung carcinomas and their expressions were inversely associated with survival, providing a potentially useful marker for diagnosis and prognosis of lung cancer [21].

Overexpression of *HMGA2* gene leads to pituitary adenomas in mice. The mechanism has been described by Fedele*et al.*[22]. HMGA2 binds to the pRB A/B pocket domain, while it does not compete with the E2F1 protein. Conversely, E2F1 activation by HMGA2 occurs by displacing HDAC1 from the pRB/ E2F1 complex, resulting in enhanced acetylation of both E2F1 and DNA-associating histones, thereby promoting E2F1 activation [22]. It is well-known that pRB controls cell cycle progression through its interaction with the E2F family of transcription factors, whose activity is crucial for the expression of several genes required for cells to enter the S phase of the cell cycle[23]. By repressing E2F1 activity, pRB protein prevents cell from progressing beyond the G1 phase of the cell cycle. If the repression of E2F1 is relieved by phosphorylation or viral transformation of pRB[24, 25], resulting in the release of E2F1, the transcription of its target genes is activated[26]. This allows cells to progress toward S phase. The overexpressions of pRB[27] and E2F1[28] were also found in ESCC specimens.These findings are consistent with the mechanism in pituitary cancer described by Fedele*et al.*. This suggests that the pRB/E2F1 pathway involving *HMGA2* may also play a critical role in the pathogenesis of ESCC.

In summary, the gene microarray results show a comprehensive picture of the differential gene expression in ESCC. Thirty novel overexpressed genes were revealed in this study. The real-time qPCR results confirmed that *HMGA2* was up-regulated in all the ESCC cell lines. In addition, the protein expression of HMGA2 demonstrated a significantly higher incidence of overexpression in primary ESCCs than morphologically non-tumoralesophageal epithelium tissue. For the first time, the present findings showed that *HMGA2* was overexpressed in ESCC, and suggest that the activation of HMGA2 might be important in the pathogenesis of ESCC.

Table 1. Primer sequences of studied genes.

Gene	DNA sequences
HMGA2	5'-CAGCAGCAAGAACCAACC-3'
	5'-CAGTTTCCTCCTGAGCAG-3'
PEG10	5'-GGGTCTGTCATCGACTAC-3'
	5'-CTCGGTTGGATCTACCTG-3'
SHANK2	purchased from SuperArray Bioscience Corporation (Frederick, MD, USA)
WISP3	5'-CAGCAGCTTTCAACAAGCTACA-3'
	5'-TTCCCATCCCACATGTTCTG-3'
β-Actin	5'-GCTCGTCGTCGACAACGGCTC-3'
	5'-CAAACATGATCTGGGTCATCTTCTC-3'

			F	Relative intensit	У*
Gene Symbol	Position	Chromosome	Tumor	Lymph node	HKESC-4
·		location		metastasis	cells
CAMK2A	229163 at	1p36.13	24.08	17.26	20.94
CAMK2A	218309 at	1p36.13	22.7	17.76	23.08
CAMK2A	228302 x at	1p36.13	17.06	11.62	28.18
CART1	206837 at	12a21.3-a22	16.01	15.6	14.43
CCNA1	205899 at	13a12.3-a13	50.35	58.86	48.36
COCH	205229 s at	14a12-a13	14.14	13.1	11.59
DKK1	204602 at	10011.2	64.93	96.28	344.8
EGFR	201984 s at	7p12	22.85	31.67	30.82
EMS1	214073 at	11a13	17.43	17.63	21.22
FLJ33516	229160 at	Χα22.3	18.6	53.55	32.52
FMN2	223618 at	1043	16.56	22.98	22.86
FMN2	1555471 a at	1043	11 93	14 46	17.5
FOXD1	206307 s at	5a12-a13	15 44	20.53	13.95
FOXG1B	206018 at	14012-013	18.82	24.46	30.44
GAL	200010_at	110131	41.56	46 58	41 16
	278/6 at	15014	12 01	11 //	13.8/
	206685 at	6n21 2	11 //	21.19	7 004
	200005_at	12015	21.22	62.65	17.04
	200025_5_at	12013 2	22.06	22.05	21 12
	210909_at	12413.3 2a21.1	12.90	15.05	01.13
	229400_at	2431.1	20.12	10.00	21.27
	214004_at	2431.1	20.12	10.00	23.37
	207307_at	7p11	21.95	10.02	20.37
		/pii	21.00	30.90	30.02
	202267_at	1,20-1	21.33	28.72	11.3
	229125_at	1p32.1	14.43	100.0	39.89
	207325_x_at	Xq28	89.08	192.8	129.5
	210503_at	Xq28	36.73	43.59	25.86
MAGEA4	214254_at	Xq28	30.04	47.19	28.42
MAGEB2	206218_at	Xp21.3	101.3	111.9	113.1
MFAP2	203417_at	1p36.1-p35	47.36	25.34	14.93
MGC17986	1552946_at	19013.33	19.44	21.97	18.82
MGC27005	156/912_s_at		345.1	662	441.5
MGC27005	235700_at	Xq26.3	227.8	410.3	305.3
MMP13	205959_at	11q22.3	206.5	82.06	46.82
NBEA	239010_at		24.72	19.12	21.34
NKX2-2	206915_at	20pter-q11.23	20.36	20.32	19.78
PCDHB5	223629_at	5q31	13.68	19.72	25.47
PEG10	212094_at	7q21	19.58	27.79	27.8
PFN2	204992_s_at	3q25.1-q25.2	14.39	20.15	17.83
POPDC3	219926_at	6q21	20.75	26.54	28.86
PPFIA1	210236_at	11q13.2	14.19	17.12	15.62
PRAME	204086_at	22q11.22	11.47	17.7	13.24
PTHLH	1556773_at	12p12.1-p11.2	42.7	120.4	67.42
PTHLH	206300_s_at	12p12.1-p11.2	41.34	76.86	48.18
PTHLH	211756_at	12p12.1-p11.2	35.64	76	34.72
SAGE1	220793_at	Xq26	15.89	20.5	56.04
SHANK2	213308_at	11q13.2	29.9	19.83	19.99
SHANK2	213307_at	11q13.2	19.41	14.38	15.04
SIX1	228347_at	14q23.1	20.37	23.38	19.25
SLCO1B3	206354_at	12p12	13.48	27.82	30.46
SYT1	203999_at	12cen-q21	21.44	20.41	21.4
TP53TG3	220167 s at	16p13	16.8	12.91	10.12

Table 2. Genes with >10-fold higher expression in esophageal squamous cell carcinoma tissues and cells studied

WISP3	210861_s_at	6q22-q23	36.77	52.97	26.4
ZIC1	206373_at	3q24	61.6	70	97.5
ZNF595	227952_at		20.42	39.94	31.02

* Relative to morphologically normal esophageal epithelium

Gene Symbol	Description	Reference
DKK1	Overexpression of DKK1 gene in the distal squamous	[29]
	esophageal mucosa in patients with esophagitis	
EGFR	Overexpression of EGFR in ESCC and its correlation with depth	[30]
	of tumorinvasion	
EMS1	Association of amplification and overexpression of EMS1 with	[31]
	lymph node metastasis in ESCC	
GAL	Distribution of galanin(GAL) immunoreactive nerve bundles and	[32]
	scattered nerve fibres in esophageal carcinoma	
HCG	High expression of HCG expression in patients with lymph node	[33]
	metastasis and its correlation with infiltration and metastasis	
LAMC2	Co-expression of LN-5 gamma2 (LAMC2) and EGFR is closely	[34]
	related to the progression and poor prognosis of ESCC	
PTHLH	High level of serum parathyroid hormone-related protein	[35]
	(PTHLH) in esophageal carcinoma	
MAGE-A,	High expression of MAGE-A, MMP13 and zinc finger proteinsin	[36]
MMP13,	ESCC	
ZNF595		

Table 3. Genes found to have overexpression in ESCC in previous studies.

Figure 1.



N	Morphologically normal esophageal epithelium collected from the same patient from which the HKESC-4 was derived.
Т	Tumor esophageal epithelium collected from the same patient from which the HKESC-4 was derived.
LN	Lymph node metastasis of ESCC collected form the same patient from which HKESC-4 was derived.
Non-T	Another three non-tumoral esophageal tissues from esophageal cancer patients.

С

Figure 2.



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Figure legends

Figure 1.Gene expressions of *HMGA2* (A), *PEG10* (B), *SHANK2* (C) and *WISP3* (D) relative to β-actin have been shown in clinical tissues and ESCC cell lines. The value of N is arbitrarily set to 1 for comparison.

Figure 2.Immunohistochemicalstaining of HMGA2 in ESCC tissues. Original magnification, 200X(A, C and E); 400X (B, D and F).