Quantitative study of epigenetic signature in head >and neck squamous cell carcinoma

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Background/aim: The aberrant upregulation of Forkhead box protein M1 (FOXM1) plays a fundamental role in cancer initiation by perturbing stem cell differentiation. This study aims to investigate the role of FOXM1 in epigenetic modification and gene expression of target genes in primary human oral keratinocytes, squamous cell carcinoma cell lines, and head and neck squamous cell carcinoma (HNSCC) tissue biopsies.

Materials and methods: A genome-wide promoter methylation microarray was used to compare HNSCC cell line (n = 8), primary human oral keratinocytes (NOK; n = 8) transduced with FOXM1 and EGFP, and HNSCC tissue biopsies (n = 3). Seventeen Foxm1B-induced differentially methylated genes were shortlisted. An absolute quantitative polymerase chain reaction was used to validate the differential promoter DNA methylation of each candidate gene induced by FOXM1. These results were compared with the methylation status and altered gene expressions of candidate genes in a panel of genomic DNA and messenger RNA (mRNA) samples previously extracted from HNSCC tissue biopsies.

Results: The results were consistent with our hypothesis, showing that aberrant upregulation of FOXM1 expression in vitro primary NOK induces a global hypomethylation pattern similar to the HNSCC cell line and has an inverse correlation with in vivo mRNA expression levels of HNSCC tissue biopsy.

Conclusion: Such epigenetic changes have tremendous clinical potential as biomarkers for early cancer detection and therapeutic interventions.

Key words: Forkhead box protein M1, epigenetic modifications, gene expression
in the last few decades. Detection of HNSCC at an early stage can increase survival by up to 80%, hence suggesting that the early detection of stable markers such as DNA methylation markers can provide immense assistance in the diagnosis, treatment, and prediction of recurrence of HNSCC.

1.2. Epigenetics
Epigenetics refers to heritable and stable changes in gene expression without alteration in primary DNA sequences. These changes are brought about by different epigenetic mechanisms such as DNA methylation or methylation, phosphorylation, or acetylation of histone proteins around which the DNA wound itself to form chromatin. Epigenetic regulation controls the cellular RNA expression patterns. Epigenetics is involved in multiple processes such as gene silencing, X chromosome inactivation, paramutation, imprinting, the progress of carcinogenesis, and embryonic development (7,8). Any dysregulation in epigenetic mechanisms may lead to various human diseases, including autoimmune diseases, cancer, and neurological disorders (9). During cancer there is uncontrolled cell growth as cells escape the normal physiological regulation of proliferation, differentiation, and apoptosis. This takes place as a result of the silencing of tumor suppressor genes or abnormal activation of oncogenes. It has been established recently that there are factors other than genetic abnormalities, such as DNA sequence mutation and genomic instability, that lead to cancer predisposition. Here epigenetics comes into play. Research in this new field of science has shown that the early epigenetic alterations are a prerequisite to oncogenic genetic mutations that lead to cancer.

Of the various epigenetic mechanisms, DNA methylation has been intensely investigated. It involves the addition of a methyl group to the 5 position of the cytosine pyrimidine ring. During cell proliferation the DNA methylation pattern is maintained by DNA methyltransferase enzymes (DNMT1, DNMT3a, and DNMT3b) (10,11). DNA methylation plays a major role in normal cellular differentiation and development. Stable change in the methylation status of DNA can alter the gene expression so that the cells have the ability to retain their cellular pattern. Aberrant DNA methylation is associated with malignancies and can occur in 2 forms: global DNA hypomethylation and promoter DNA hypermethylation (12–14). DNA hypomethylation is associated with earlier stages of cancer. It can result in the reactivation of various cancer- and growth-related genes as well as chromosomal instability and genetic mutations, whereas hypermethylation in cancer cells occurs in the promoter regions of CpG islands (15,16) and results in the silencing of tumor suppressor genes. The loss of function of tumor suppressor genes leads to aberrant cell proliferation, lack of DNA repair, apoptosis evasion, and the continuation of the cell cycle even with the presence of damaged DNA, and all of these events lead to genomic instability that occurs due to loss of heterozygosity and chromosomal number abnormality. In normal cells the tumor suppressor genes are unmethylated and hence they are normally transcribed and can perform their functions. Promoter DNA methylation induces the loss of expression of tumor suppressor genes, predisposing transformation of normal cells into oncogenesis.

The other epigenetic mechanism is posttranslational modification of amino acids in histone proteins. Histones are proteins around which DNA wraps itself for compaction. Histone modification appears to work by maintaining and establishing the gene activity states as it controls DNA accessibility. DNA is highly accessible when histone H3 is methylated (H3K4me) and histone H4 is hyperacetylated (H4K16Ac), whereas DNA is inaccessible when histone 3 is methylated on lysine residue 27. The H2A family of histone proteins plays a major role in chromatin reprogramming; damage to this family changes its functions and results in genomic instability and cellular proliferation.

Any dysregulation in the abovementioned chief epigenetic mechanisms can lead to numerous diseases like cancer, autoimmune diseases, and neurological disorders (9). Epigenetic changes are reversible and precede genetic expression; therefore, they are thought to be an earlier mechanism, with an edge over genetics. Epigenetic changes have tremendous potential for the discovery of early cancer biomarkers, essential in screening, diagnosis, prevention, and treatment. For example, cancer can be prevented or treated by reversing cancer-specific epigenetic changes before irreversible mutations take place. Figure 1 illustrates a common alteration in cancer cells: hypermethylation of a tumor’s suppressor genes suppresses its transcription, leading to loss of its normal cellular functions.

1.3. Forkhead box M1 transcriptional factor
FOXM1 (previously called HFH-11B, TRIDENT, WIN, MPP2, and FOXM1b) is a protein that is encoded by the FOXM1 gene in humans. This protein is a member of the FOX family of transcription factors, which includes a minimum of 50 unique FOX genes. FOXM1 is present in 3 isoforms: FOXM1A/isoform1, FOXM1B/isoform3, and FOXM1C/isoform2. All 3 isoforms can bind to DNA, but only FOXM1B and FOXM1C act as transactivators and hence can activate their target genes by binding to 5’-A(C/T)AAA(C/T)AA-3’ (3). The cellular localization and regulation of FOXM1 is determined by posttranslational modification such as phosphorylation of FOXM1.

FOXM1 protein is highly expressed in actively dividing cells (17), whereas its levels are hardly detectable in terminally differentiated cells (18). This protein plays a key role in cell proliferation, cell growth, development,
differentiation, and longevity. Cells lacking the FOXM1 protein show multiple defects, such as chromosomal missegregation, polyploidy, an abnormal number of chromosomes, defects in cytokinesis, and DNA fragmentation. Hence, this suggests a fundamental role for FOXM1 in DNA repair (19) as well as maintaining genomic stability (2).

During embryogenesis FOXM1 expression is induced in a number of tissues, as it plays a vital role in the development of lungs, heart, blood vessels, and the liver. In adult tissues its levels are greatly decreased, but during organ injury and numerous cancers its expression level is high (20). During the cell cycle FOXM1 levels are found to be highest in the S phase and the G2/M phase (3,17). FOXM1 can regulate the expression of multiple cell cycle genes that control the G1/S transition phase, S phase and M phase progression, and G2/M phase and chromosomal segregation. FOXM1 itself is regulated by cell cycle-promoting factors such as cyclin B and cyclin D as well as antiproliferation signals that inhibit cell death, differentiation, and cell cycle arrest (3). Other than cell cycle transition, FOXM1 also plays important roles in DNA damage repair (19), apoptosis (21), tissue regeneration (22), angiogenesis (23), and metastasis (24).

1.4. FOXM1 role in human tumorigenesis

In addition to its key functions in the cell cycle and damaged DNA repair, FOXM1 plays a pivotal role in oncogenesis. FOXM1 is known as a human protooncogene. The first evidence of a link between FOXM1 and cancer was established in 2002, when it was found that in basal cell carcinoma there is abnormal upregulation of FOXM1 (25). Subsequent studies and microarray analyses have shown that FOXM1 is overexpressed in a number of human cancers. Furthermore, it has been established that during head and neck cancer FOXM1 expression is upregulated during the early stages (4). Epithelial stem cells having features like self-renewal, high migration capacity, drug resistance, and high clonogenic potential are susceptible to oncogenic selection and are thought to have a role in the development and maintenance of the majority of tumors. Studies have shown that FOXM1 plays a major role in the regulation of adult epithelial stem cell renewal and differentiation. Recent evidence supports the notion that aberrant upregulation of FOXM1 expression in undifferentiated human keratinocyte stem cells possessing high clonogenic potential leads to a disturbance of epithelial differentiation, leading to formation of hyperproliferative “precancer” stem cells (5). This finding indicates that abnormal upregulation of FOXM1 in normal stem cells leads towards a multistep oncogenic evolutionary pathway.

The role of FOXM1 is not only limited to the initiation of cancer; it also plays an important role in all stages of tumorigenesis, from early predisposition and tumor initiation (5) to cancer progression (4,26) and metastasis (3). There are a number of carcinogenic and environmental factors that activate FOXM1, resulting in uncontrolled cell proliferation and malignant transformation: for example, nicotine in tobacco (4), oxidative stress (27,28), UV light (29), and ionizing irradiation (19). These factors are reported to aberrantly increase FOXM1 expression levels, which induces genomic instability and epigenetic modification by activating downstream targets HELLs and CEP55 protein. HELLs is a helicase required in DNA strand separation, repair, replication, transcription, and epigenetic modification such as DNA methylation. CEP55 is a mitotic phosphoprotein required in cytokinesis (30). It has also been found that the gene locus of FOXM1 (12p13.3) is amplified in HNSCC. All these findings

Figure 1. DNA methylation in gene promoter region leads to repression of gene expression.
suggest that aberrant upregulation of FOXM1 expression results in oncogenic genomic instability and enhanced cellular proliferation leading to the development of cancer.

1.5. FOXM1, epigenetics, and HNSCC
Exposure to the various risk factors (e.g., UV light, tobacco, and ionizing radiations) associated with cancer may result in aberrant overexpression of oncogene FOXM1. It is also known that the loss of p53 function, c-MYC, RAS, and p16/Rb pathway inactivation in cancer cells lead to constitutive upregulation of FOXM1 (5). FOXM1 is known as a driver mutation, which acts during the early stages of oncogenesis by tumor initiation and predisposition (5). Epigenetic changes occur early in the development of cancer, followed by genetic mutations. FOXM1 initiates cancer by uncontrolled expansion of stem cells and by suppressing their differentiation (5). Aberrant expression of FOXM1 leads to genomic instability and epigenetic reprogramming by activating downstream targets CEP55 and DNMT1. CEP55 causes mitotic instability (4,31), and deregulation of HELLS may lead to altered genomic methylation due to downregulation of DNMT1 and chromatin remodeling (4). This all suggests that abnormal upregulation of FOXM1 brings epigenetic modifications in conjunction with genomic instability, which causes development of a heterogeneous population of abnormal cells that are more likely to acquire properties that lead to the development of cancer.

2. Materials and methods

2.1. Patient nucleic acid samples
The use of human tissue in this study was approved by Barts and the London NHS Trust; the School of Medicine and Dentistry, Queen Mary University of London; and the UK National Research Ethics Committee. All clinical samples, which were additional to diagnosis, were collected according to local ethics committee-approved protocols and written informed patient consent was obtained from all participants.

2.2. Cell culture and viral transduction
The primary normal human oral keratinocytes (OK355, HOKG, OK113, NOK, NOK1, NOK3, NOK16, and NOK376) used in this study were donated by healthy, disease-free individuals undergoing wisdom tooth extraction and were cultured as previously described (4,29). Oral SCC cell line SCC15 (32) is a well-established cell line cultured as described earlier (4,5,29).

In this experiment human primary oral keratinocytes were grown and subsequently transduced by the gene of interest. We used the FOXM1B human protooncogene and EGFP (an inert enhanced green fluorescent protein) as controls. In order to make infectious viral particles, retroviral packaging cells (Phoenix A) were first transduced with a retroviral DNA plasmid carrying the gene of interest. Those cells that were unable to be infected by virus were incapable of surviving the antibiotic selection, while the remaining transduced cells were cultured for viral production. A viral supernatant was obtained and was used to transfect the target primary human oral keratinocytes with FOXM1B and EGFP. Retroviral supernatant and transduction procedures were performed using our established protocols (4,5,29). Equal levels of EGFP and FOXM1B expression were achieved by serial retroviral supernatant titration experiment and subsequently EGFP plasmid copy numbers were confirmed by quantitative polymerase chain reaction (qPCR) using genomic DNA extracted from transduced cells according to our previously established method (29). The levels of ectopic FOXM1 expression in the primary keratinocytes were titrated to replicate levels found in cancer cells as reported previously (4,5,29). Transduced cells were cultured for 3–5 days to allow transgene expression prior to experiment.

2.3. Statistical analysis
Statistical analysis was performed with Microsoft Excel using the Student t-test (2-tailed). P < 0.05 was considered to be statistically significant.

2.4. Ethical approval
The human tissues used in this study were approved by Barts and the London NHS Trust; School of Medicine and Dentistry, Queen Mary University of London; and the UK National Research Ethics Committee.

2.5. Isolation of methylated genomic DNA

2.5.1. Fragmentation of gDNA
The samples used were DNA from normal oral keratinocytes treated with EGFP, cells infected by FOXM1B, and the HNSCC cell line (SCC15). This genomic DNA was fragmented by treating it with an enzyme according to the manufacturer’s protocol. The kit used was the Methyl Collector Ultra (catalogue no. 55005) provided by Active Motif Europe, Belgium.

2.5.2. Enriching methylated gDNA
The following protocol was used to enrich gDNA: a complete binding buffer was prepared, and a high-salt and a low-salt binding buffer were both produced by using reagents supplied in the kit (Table 1).

Next, 10 µL of magnetic beads was added to the required number of PCR tubes for the reaction and components shown in Table 2 were added to the tubes to complete the binding reaction.

The mixture was shaken and incubated for 1 h at 4 °C on a rotisserie shaker. The tubes were then placed on a magnetic stand so that the beads moved to one side and the supernatant that contained the unbound fraction was removed. To resuspend the beads, they were washed with 200 µL of binding buffer and the supernatant was...
removed. To recover the methylated DNA fragments from the beads they were resuspended in 100 µL of complete elusion buffer.

Elusion buffer for each reaction consisted of 2 µL of proteinase K and 98 µL of elusion buffer AM1. Next, it was incubated for 30 min at 50 °C, inverted every 10 min so that the beads were resuspended. Samples were returned to room temperature before the addition of 2 µL of proteinase K stop solution. The stop solution was warmed for 10 min at 37 °C prior to addition. Samples were mixed and then placed on a magnetic stand, the beads moved to one side and the supernatant removed and transferred to new tubes. The supernatant consisted of the methylated gDNA fragments.

2.5.3. DNA clean up
The DNA was cleaned in order to get rid of all protein contaminations necessary before PCR amplification. The protocol used was the phenol/chloroform extraction and ethanol precipitation.

2.6. Microarray and gene selection
A nonbiased genome-wide microarray promoter methylation profiling analysis was performed in order to obtain a list of differentially methylated genes present in the methylation-enriched gDNA and total gDNA that had been extracted and purified from normal primary oral keratinocytes transduced with either EGFP or FOXM1B and the HNSCC cell line (SCC15) as a positive control. The genes that were most significantly differentially methylated by FOXM1B were then identified and shortlisted. The list of candidate genes was further validated by using qPCR. This microarray analysis was performed using NimbleGen 720k array chips, which cover 22,532 promoter CpG islands proximal to transcriptional start sites. The service was provided by Roche NimbleGen, Inc., USA. Figure 2 illustrates the selection criteria of candidate genes for this project. The most significantly differentially methylated genes in control cells (NOKG) were identified. These were then compared with gene lists of NOKF and SCC15 cell lines. Genes that showed opposing methylation status between NOKG and NOKF were selected and compared with SCC15 to find the list of common genes between NOKF and SCC15.

2.7. List of target genes and primer sequence
The 17 target genes and primers are given in Tables 3, 4, and 5.

2.8. Real time quantitative PCR (RT-qPCR)
A LightCycler LC480 qPCR machine (Roche Diagnostics Ltd., UK) was used for absolute gene quantification. The SYBR Green (Roche Diagnostics) method was used. It was confirmed that the PCR primers used only produced a single peak. The 2 most stable reference housekeeping genes used were YAP1 and POLR2A.

2.9. qPCR workflow
Melting peak analysis was performed to verify the specificity of PCR primers. Ten-fold serial dilutions of the amplified products were prepared in t-RNA solution (Sigma). Absolute quantification for a single target gene was achieved by creating standard curves. These were then stored in the LC480 analysis software program for future sample analysis. Each PCR experiment consisted of

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**Table 1. Buffer conditions.**

<table>
<thead>
<tr>
<th>High-salt binding conditions</th>
<th>Low-salt binding conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reagent</td>
<td>One reaction</td>
</tr>
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<td>Binding buffer AM7</td>
<td>Binding buffer AM12</td>
</tr>
<tr>
<td>Protease Inhibitor cocktail</td>
<td>Protease Inhibitor cocktail</td>
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<tr>
<td>Total volume</td>
<td>Total volume</td>
</tr>
<tr>
<td>100 µL</td>
<td>100 µL</td>
</tr>
<tr>
<td>0.5 µL</td>
<td>0.5 µL</td>
</tr>
<tr>
<td>100.5 µL</td>
<td>100.5 µL</td>
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</table>

**Table 2. Reagents used.**

<table>
<thead>
<tr>
<th>Reagents</th>
<th>One reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnetic beads</td>
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<tr>
<td>Complete binding buffer (high or low salt)</td>
<td>70 µL</td>
</tr>
<tr>
<td>Fragmented genomic DNA</td>
<td>From 1 ng to 1 µg (in a final volume of 10 µL)</td>
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<tr>
<td>His-MBD2/MBD3L1 protein complex</td>
<td>10 µL</td>
</tr>
<tr>
<td>Total volume</td>
<td>100 µL</td>
</tr>
</tbody>
</table>
Figure 2. Selection of FOXM1B induced differentially methylated genes. The most significantly differentially methylated genes in control cells (yellow) were identified. These were then compared with a gene list of FOXM1B (blue) and SSC15 (orange) cell lines. Genes that showed opposing methylation status between control cell line and FOXM1B transduction cell line were selected and compared with SCC15 to find the list of common genes between FOXM1B-treated cell lines and SCC15.

Table 3. Hypermethylated genes.

<table>
<thead>
<tr>
<th>Size (bp)</th>
<th>Gene</th>
<th>Primer sequence</th>
</tr>
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<td>gTET3-F</td>
<td>aggtgcacagagttggagt</td>
</tr>
<tr>
<td></td>
<td>gTET3-R</td>
<td>gccatggctctgcttttg</td>
</tr>
<tr>
<td>76</td>
<td>gTAS2R60-F</td>
<td>ccatggatgccatgttgtgct</td>
</tr>
<tr>
<td></td>
<td>gTAS2R60-R</td>
<td>acattctgggattgtgtatgaa</td>
</tr>
<tr>
<td>95</td>
<td>gFSTL3-F</td>
<td>aaaaagtgcctcccttaggtgt</td>
</tr>
<tr>
<td></td>
<td>gFSTL3-R</td>
<td>cttggactttatcccccttggt</td>
</tr>
<tr>
<td>93</td>
<td>gPDGFB-F</td>
<td>tccacccctagtgccttcct</td>
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<tr>
<td></td>
<td>gPDGFB-R</td>
<td>aagggaaagccccccaaaat</td>
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<tr>
<td>102</td>
<td>gGN13-F</td>
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<td>gGN13-R</td>
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<td>94</td>
<td>gNXPH2-F</td>
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<tr>
<td></td>
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<td>89</td>
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<tr>
<td></td>
<td>gATF6B-R</td>
<td>acttccttgctccacactg</td>
</tr>
<tr>
<td>123</td>
<td>gFKBPL-F</td>
<td>tgctagggcagctcagtg</td>
</tr>
<tr>
<td></td>
<td>gFKBPL-R</td>
<td>cttttccaggttcccaagg</td>
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</table>
a standard concentration of 10^5 copies of target genes and housekeeping genes in order to find out the copy number of target genes in each unknown gDNA or cDNA sample. The LC480 software automatically calculates the gene copy number present in unknown samples by comparing it with the calibrated standard (10^5 copies) respective to that of previously stored standard curve for each target gene. A Microsoft Excel spreadsheet was used for data calculation and analysis.

3. Results

3.1. Gene expression quantification using qPCR
To quantify the exact mRNA copy number of target genes in the FOXM1B-transduced cells and cancer cell lines, we produced standard curves by using known DNA concentrations (copy number). Amplification curves and melting analyses were performed for all the target genes. Melting analysis determines PCR product specificity by showing only one peak, whereas amplification curves help to produce standard curves to determine PCR efficiency and curve fitting errors to allow accurate calculation of target gene concentrations in unknown samples, based on the Roche Light Cycler480 operating software internal algorithm. YAP1 and POLR2A were used as housekeeping genes, as they have been previously shown to be stably expressed in both normal and cancer cells (4).

3.2. Target genes table
In this study 17 target genes were investigated. Table 6 is a compilation of all the target genes. These details were taken from the standard curve produced for each candidate gene.

3.3. Correlation between promoter methylation and gene expression
It is known that differential promoter hypermethylation leads to silencing of gene expression whereas hypomethylation causes upregulation of gene expression. In this project we used an absolute qPCR to first validate
the differential methylation of genes in methylated gDNA collected from our control primary normal human oral keratinocytes, expressing either EGFP (NOKG) or FOXM1 (NOKF) SCC15 cell lines, and in methylated gDNA and cDNA collected from tissue biopsy of 3 pairs of normal margin and oral SCC cells. The absolute qPCR measured and compared the relative methylation and gene expression of the 17 candidate FOXM1B-induced differentially methylated genes that were identified from the microarray study. These genes showed the parallel differential gene expression in both cultured cells and in patient tumor tissues. According to our hypothesis, if the shortlisted 17 target genes were involved in oncogenesis, then their promoter methylation status would modify the gene expression in the cancer tissues as well. In order to support our hypothesis we performed absolute qPCR to measure the gene expression of these candidate genes in 3 pairs of normal margin and HNSCC biopsy tissues. Once their gene expression levels were obtained we next correlated them with the promoter DNA methylation status of these genes in FOXM1B-transduced cells.

3.4. DNA methylation vs. mRNA expression in NOKF
To validate if the candidate genes were indeed differentially methylated in FOXM1-transduced cells (NOKF), methylation (using gDNA) and gene expression (cDNA) were quantified for each gene in NOKG and NOKF. Differential methylation and gene expression were then calculated for each gene as a ratio of NOKF:NOKG and plotted as a correlation chart between gene expression and relative methylation. Figure 3 shows that there is an inverse correlation between promoter methylations and their respective gene expressions in FOXM1B-transduced cells, with a regression coefficient $R^2$ value of 0.564. Each data point with an error bar represents the mean ± SEM of 3 NOKF samples. A positive control gene, CDKN2A/p16, known to be hypermethylated by FOXM1B, was included in this study. This negative correlation is in agreement with our hypothesis. According to this graph, the methylation status of target genes from the microarray analysis is consistent with the qPCR results.

3.5. DNA methylation vs mRNA expression in SCC cell lines
Using the same validation strategy as above, the candidate genes were subsequently validated in SCC15 cell lines. Similarly, Figure 4 shows a negative relation between the promoter DNA methylation and relative mRNA expression in the SCC15 cell line with a regression coefficient $R^2$ of 0.502.

Table 6. Target gene standard curve metadata.

<table>
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<tr>
<th>Gene</th>
<th>Size (bp)</th>
<th>Tm</th>
<th>Primer dimer</th>
<th>Error</th>
<th>Efficiency</th>
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<td>PDGFB</td>
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<td>NXPH2</td>
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<td>ATF6B</td>
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<td>FKBPL</td>
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</table>
3.6. DNA methylation in NOKF vs. relative mRNA expression in HNSCC cell lines
We next investigated whether the candidate gene expression pattern was a common phenomenon across different HNSCC cell lines. There were 8 samples used (SCC15, 5PT, VB6, SqCC/Y1, SVpgC2a, SVFN4, SVFN5, and SVFN10). cDNAs were prepared from these 8 independent HNSCC cell lines and then compared with DNA methylation status for each candidate gene. Figure 5 shows the inverse correlation between the promoter methylation status in FOXM1B-induced cells, NOKF, and the corresponding gene expression in HNSCC cell lines. The regression coefficient $R^2$ was 0.849. This graph indicates that the candidate genes selected from FOXM1B-induced cells behaved the same in HNSCC cell lines.

3.7. Correlation between NOKF and tissue biopsies
Having established that the candidate differentially methylated genes showed inverse gene expression patterns in HNSCC cell lines, we next investigated if this trend persisted in in vivo HNSCC tumor biopsy tissues as well. First, the relative methylation status of each gene was compared between gDNA extracted from NOKF and HNSCC tissues. Three pairs of tissue biopsy samples were used. In Figure 6A there is a positive correlation between promoter methylation of NOKF and HNSCC tissue biopsies’ gDNA, with a correlation coefficient of $R^2 = 0.637$. This indicates that the methylation signature found in NOKF appears to be similar to that found in HNSCC tumor tissues.

To further validate that the differential methylation signature indeed resulted in inverse gene expression in the tumor tissues, the gene expressions of each of the candidate genes were quantified and compared between the 3 pairs
of normal margin and HNSCC tumor tissues. In Figure 6B an inverse correlation is shown between promoter methylation in NOKF and its corresponding mRNA expression in HNSCC tissues. The correlation coefficient \( R^2 \) is 0.921. This graph provides data that seem to be in agreement with our hypothesis that FOXM1 induces a cancer-like methylation signature. As stated previously, if the candidate genes were involved in oncogenesis, we hypothesized that their promoter methylation status would change the mRNA expression in tumor tissues.

### 3.8. Correlation between DNA methylation and relative mRNA expression in individual patient samples

Although the candidate genes showed differential gene expression in the HNSCC tumor samples, the methylation statuses of these genes in the tumor samples were not known. In order to investigate this, gDNA from the same 3 pairs of normal margin and tumor tissues were analyzed for differential methylation. Figures 7A–7C show 3 patient samples, A, B, and C. These graphs illustrate the inverse correlation of individual patient between promoter DNA methylation and relative mRNA expression.

Figure 7D shows the average association between the methylation status and the corresponding cDNA expression in all 3 patient samples. In the graph of patient A, all genes other than OR3A1 behaved as expected, hence giving an inverse relation value for \( R^2 \) of 0.972. The genes that were hypomethylated were highly expressed, such as, for example SPCSI and GALT8D1, whereas the hypermethylated genes showed low levels of expression of their respective genes, for example PDGFB and TET3.

The graph for patient B also shows a negative relation, with a regression coefficient \( R^2 \) value of 0.577. FKBPL showed an opposing DNA methylation status, which related well to its mRNA expression. Other hypermethylated genes, such as TAS2R60, PDGFB, FSTL3, and UTR-p16, showed downregulation of respective gene expression, whereas the hypomethylated genes such as SPCSI, OR3A1, GLT8D1, and FABP6 showed corresponding gene upregulation.

For patient C, an inverse relation was seen with regression coefficient \( R^2 \) of 0.760. In this graph the methylation statuses of a few genes were not consistent with the microarray results, as these genes showed opposing methylation statuses, for example FSTL3, PDGFB, TAS2R60, and OR3A1. Nevertheless, genes FSTL3, PDGFB, and TAS2R60 showed DNA hypomethylation, but these 3 genes correlated well to their respective gene expression, whereas OR3A1 showed DNA hypermethylation and corresponding gene expression downregulation.

In Figure 7D, all 3 patients’ data were averaged to obtain the combined result of the relationship between the promoter DNA methylation and respective mRNA expression levels. As expected, a negative correlation between methylation and gene expression was obtained, having a regression coefficient of 0.954.
4. Discussion

FOXM1, an important transcription factor for the regulation of cell cycle, cell division, genomic stability, aging, and development, has been found to be ubiquitously upregulated in many human cancers. It is known that FOXM1 expression is upregulated early
during HNSCC carcinogenesis (4). The understanding of the early mechanism of cancer initiation can help us find effective interventions for cancer prevention and cure. FOXM1’s role in cancer initiation by perturbing stem cell renewal has been recently established (5). Epigenetic modifications like DNA methylation and demethylation are also known to occur early during cancer (6). It is hence important to know whether FOXM1 has a role in epigenetic modification that initiates cancer formation. We hypothesize that upregulation of FOXM1 induces cancer-inducing epigenetic modifications such as DNA methylation and demethylation. In order to support our hypothesis we analyzed a list of differentially methylated genes that had been shortlisted after performing a genome-wide microarray promoter DNA methylation study. The promoter DNA methylation pattern and corresponding mRNA expression of each candidate gene is measured in normal primary oral keratinocytes with increased FOXM1 expression by using real-time absolute qPCR and is then compared to HNSCC cell lines, HNSCC biopsy cDNA samples, and a normal control.

An absolute qPCR was used to obtain a standard curve for each candidate gene so that accurate levels of promoter DNA methylation and the respective mRNA expression could be measured. First, to validate that the target genes were indeed differentially methylated in FOXM1-transduced cells, the promoter methylation and relative gene expression were quantified for each gene in control cells (NOKG) and NOKF. The data obtained showed that the microarray analysis was mostly consistent with the qPCR results, whereby the methylation status of each gene was inversely correlated with gene expression. An inverse relation was obtained with a correlation coefficient of $R^2 = 0.564$ (about 56%). Two genes, $ATF6B$ and $GNG13$, that were expected to be hypermethylated did not exhibit the strong status that was seen in microarray gene expression for $FOXM1$ and $SCC15$, but nevertheless their respective mRNA expressions were consistent with their DNA methylation status. This highlights the importance of validating microarray results using qPCR. A positive control gene, $p16$, which is known to be a tumor-suppressor gene, was included in this study. It was shown from the results that this gene was hypermethylated in NOKF and in the majority of HNSCC cell lines and tumor tissues and, correspondingly, its mRNA expression levels were suppressed. The $p16$ gene has a regulatory role in the cell cycle. It stabilizes the tumor suppressor gene $p53$, which controls apoptosis (32). Mutation of this gene increases the risk of developing cancer. It can cause downregulation of genes and impairment of the retinoblastoma pathway (33), of which CDK4 has a role, and $p16$ has a regulatory function over CDK4. Silencing of the gene $p16$ has been associated with numerous cancer cell lines. It is found to be inactive in HNSCC and is associated with advanced stages of this type of cancer (34). The silencing of $p16$ as a result of induced hypermethylation by FOXM1 may promote cancer development and progression.

Next, the candidate genes were subsequently validated in the SCC15 cell line using the same validation strategy as above. An inverse negative correlation between the promoter DNA methylation and relative mRNA expression in the SCC15 cell line, with a regression coefficient $R^2$ of 0.502 (about 50%) was obtained. Four genes, $PDGFB$, $FKBPL$, $OR3A1$, and $SPCS1$, showed opposing methylation status. However, the mRNA expression of these 4 genes correlates well with their respective promoter methylation status. For example, $SPCS1$ and $OR3A1$ showed hypermethylation with corresponding downregulation of gene expression. Similarly, $PDGFB$ and $FKBPL$ showed hypomethylation and upregulation of the relevant gene expression. The $FKBPL$ gene encodes a protein that plays a role in immunoregulation and basic cellular processes involving protein trafficking and folding. It is thought to have a role in induced radio resistance and also appears to have involvement in cell cycle (Gene ID 63943, NCBI 2011). The other $PDGFB$ gene encodes for a protein that is a member of the platelet-derived growth factor family. Mutations in this gene are associated with meningioma. Reciprocal translocations between chromosomes 22 and 7, at sites where this gene and that for COL1A1 are located, are associated with a particular type of skin tumor called dermatofibrosarcoma (Gene ID 5155, NCBI 2011).

To confirm whether the candidate gene expression pattern was a common phenomenon across different HNSCC cell lines, gene expression was measured in 8 independent HNSCC cell lines for correlation with DNA methylation status for each candidate gene. The results indicated that the candidate genes selected from FOXM1-induced cells showed a similar inverse correlation between methylation and gene expression. A coefficient of $R^2 = 0.845$ (about 85%) was obtained. This was true except for 2 genes, $GNG13$ and $ATF6B$, which showed promoter hypomethylation in NOKF and low levels of their corresponding gene expression in HNSCC cell lines. Their methylation status did not match their respective mRNA expression levels. The candidate differentially methylated genes showed inverse gene expression pattern in HNSCC cell lines. In order to investigate if the same trend persisted in HNSCC tumor biopsy tissues we compared the relative methylation status of each gene between gDNA extracted...
from NOKF and HNSCC tissue. A positive correlation with a coefficient of $R^2 = 0.637$ (about 64%) was obtained. OR3A1 showed promoter hypermethylation status in NOKF with a corresponding low expression of mRNA, whereas hypomethylation in HNSCC was consistent with gene expression. The data indicated that the methylation signature found in NOKF appear to be similar to that found in HNSCC tumor tissues. Next, the gene expressions of each of the candidate genes were quantified and compared between the 3 pairs of normal margin and HNSCC tumor tissues to validate that the differential methylation signature indeed resulted in inverse gene expression in the tumor tissues. There was an inverse correlation between promoter methylation in NOKF and its corresponding mRNA expression in HNSCC tissues, with a correlation coefficient of $R^2 = 0.921$ (about 92%). This shows that FOXM1 induces a cancer-like methylation signature.

To investigate the level of heterogeneity within each patient, an individual patient's methylation status and gene expression was studied for each candidate gene. In patient A, an inverse relation of $R^2$ value of 0.972 was obtained. The OR3A1 gene showed hypermethylation, which was the same as in NOKF, but it did not correspond with its respective gene expression. In patient B, a regression coefficient $R^2$ value of 0.577 was obtained. FKBPL showed opposing DNA methylation status, but it related well to its mRNA expression. In patient C, the methylation statuses of a few genes were not consistent with the microarray results. FSTL3, PDGFB, TAS2R60, and OR3A1 genes showed opposing methylation status but correlated well with their respective gene expression. These results demonstrated an interpatient heterogeneity with certain candidate genes. Nevertheless, the overall trends of methylation and gene expression were consistent with our hypothesis that FOXM1 induced a cancer-like methylation pattern that is found in tumor biopsy tissues. In the end, we averaged the data of the 3 patients to obtain a combined pattern that is found in tumor biopsy tissues. In the end, we averaged the data of the 3 patients to obtain a combined result of the relationship between the promoter DNA methylation and respective mRNA expression levels. As expected, a negative correlation between methylation and gene expression was obtained, having a regression coefficient of 0.954 (about 95%).

Modifications in genomic methylation status may be an enhancing factor in the oncogenesis process, and evidence shows that global DNA hypomethylation might contribute to genomic instability and increased mutation rate (35). Wide-spread global hypomethylation is accompanied with hypermethylation in a few genes and an increase in DNMT1, an enzyme responsible for regulating promoter methylation (36).

DNA methylation has emerged as a highly promising biomarker and is being actively studied in multiple cancers, and a large number of potential biomarkers have been identified. In prostate cancer the hypermethylation of the glutathione-S-transferase pi gene has emerged as a good diagnostic marker (37). This project can help us find good biomarkers for the early detection of HNSCC. In this study most of the candidate genes showed differential methylation status in both FOXM1-transduced cells and in HNSCC when compared to normal control cells. These genes may have a clinical potential for early diagnosis of cancer as well as prognosis and therapeutic treatments. By detecting the methylation status of various genes we can measure the level of genomic instability in cells, which may guide us into cancer prognostics. It is well known that tumor suppressor genes are hypermethylated and hence silenced in cancer, and the use of demethylating drugs, which could interact with enzymes such as DNA methyltransferase, can play a fundamental role in the therapeutic treatment of cancer. Procaine is one such drug that has been found to have growth inhibitory effects on human cancer cells. In the future we may be able to prevent cancer by arresting the reversible epigenetic modifications before irreversible mutation takes place.

In summary, this study shows that upregulation of FOXM1 leads to cancer-like epigenetic alterations in normal primary human oral keratinocytes. The differentially methylated genes selected from microarray analysis were validated using qPCR. Genes with promoter hypermethylation showed corresponding low levels of mRNA expression and hypomethylated genes showed elevated levels of relative gene expression. The same pattern was observed in HNSCC cell lines as well as in HNSCC tumor biopsy samples. We need further studies to confirm the mechanism of epigenetic modifications and the role of each gene in cancer initiation. Future proposed experiments include the following aims:

1. To investigate the mechanism by which FOXM1 causes differential promoter methylation of candidate genes.
2. To investigate the role of the differentially methylated genes in cancer initiation and progression.
3. To investigate whether the genes identified in this study could be used in clinical practice as early detection biomarkers and for cancer prognostics. Further clinical samples validation studies would be required to establish their clinical use.

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384
References


