Abstract. We have established 3 cell lines ORL-48, -115 and -136 from surgically resected specimens obtained from untreated primary human oral squamous cell carcinomas of the oral cavity. The in vitro growth characteristics, epithelial origin, in vitro anchorage independency, human papillomavirus (HPV) infection, microsatellite instability status, karyotype and the status of various cell cycle regulators and gatekeepers of these cell lines were investigated. All 3 cell lines grew as monolayers with doubling times ranging between 26.4 and 40.8 h and were immortal. Karyotyping confirmed that these cell lines were of human origin with multiple random losses and gains of entire chromosomes and regions of chromosomes. Immunohistochemistry staining of cytokeratins confirmed the epithelial origin of these cell lines, and the low degree of anchorage independency expressed by these cell lines suggests non-transformed phenotypes. Genetic analysis identified mutations in the p53 gene in all cell lines and hypermethylation of p16 INK4a in ORL-48 and -136. Analysis of MDM2 and EGFR expression indicated MDM2 overexpression in ORL-48 and EGFR overexpression in ORL-136 in comparison to the protein levels in normal oral keratinocytes. Analysis of the BAT-26 polyadenine repeat sequence and MLH-1 and MSH-2 repair enzymes demonstrated that all 3 cell lines were microsatellite stable. The role of HPV in driving carcinogenesis in these tumours was negated by the absence of HPV. Finally, analysis of the tissues from which these cell lines were derived indicated that the cell lines were genetically representative of the tumours, and, therefore, are useful tools in the understanding of the molecular changes associated with oral cancers.

Introduction

Oral cancer is the 8th and 13th most common cancer worldwide for males and females, respectively (1), and 80% of these cases occur in the Asian region. Amongst the multiethnic population of Malaysia, oral cancer is a disease most commonly seen in the Indian ethnic group. The age standardized rate (ASR) of oral cancer occurrence in Malaysian male Indians was comparable to that of Indian males from high-incidence regions in the Indian subcontinent, but the ASR for female Indians in Malaysia was markedly higher (2). Despite advances in cancer diagnosis and treatment, there has been little improvement in the 5-year survival rate of oral cancer patients in the last few decades.

While oral cancer is generally associated with smoking or betel quid chewing habits, the genetic alterations identified in smoking-related oral cancers differ from those observed in oral cancers associated with betel quid chewing (3). However, although its aetiological factors are well-established, the mechanism through which oral cancer develops is still largely unknown, partly due to the lack of in vitro and in vivo models of oral carcinogenesis. As a result, most studies focus on oral cancer tissues, which are pivotal in any research setting and represent real changes in the tumour (4,5), but they are by definition limited. In addition, genetic manipulations to examine gene function as well as the up- and down-stream effects of such changes are restricted when working with tissue. Cell lines which are representative of particular cancers, on the other hand, make useful laboratory models to study genes and proteins implicated in the initiation and progression of cancers. For example, molecular targets regulated by epidermal growth factor (EGF) and TGFβ were identified through experiments on oral cell lines established from either the primary tumour or a lymph node metastasis.
Materials and methods

Tumour specimen and tissue collection. Cell lines were derived from surgically resected tissue specimens from 3 different oral squamous cell carcinoma patients receiving treatment at the Dental Faculty and University Malaya Medical Centre, Kuala Lumpur Hospital and Tengku Ampuan Rahimah Hospital, respectively. These patients had received no treatment prior to surgery. Informed consent was obtained from all individuals, and this project was approved as part of a main study on ‘Oral Cancer and Precancer in Malaysia’ by the Medical Ethics Committee, Faculty of Dentistry, University Malaya and endorsed by the Ministry of Health, Malaysia (Medical Ethics Clearance no. DP OP0306/0018/L). Information on the patients and specimens from which the cell lines were derived are tabulated in Table I. Tissues were collected in sterile 30-ml universal bottles containing 10 ml MEM medium (Cambrex, USA) supplemented with 10% fetal bovine serum (FBS), antibiotic solution (200 iu/ml penicillin and 200 μg/ml streptomycin) and fungizone (5 μg/ml) immediately upon excision of the tumour.

Primary culture of the human oral squamous cell carcinoma cell lines and normal oral keratinocytes. The tissues were soaked in absolute ethanol for 20-30 sec and then washed twice with phosphate-buffered saline (PBS) under sterile conditions in the laboratory. Tissues were minced, washed in culture media twice and re-suspended in α-MEM containing 20% (v/v) FBS, 200 iu/ml penicillin, 200 μg/ml streptomycin, 0.4 ng/ml EGF, 2 μg/ml hydrocortisone and 2 mM L-glutamine and seeded into 60-mm tissue culture dishes. All cultures were incubated in a humidified atmosphere of 5% CO₂ at 37°C. The culture medium was changed after 72 h. From this point, cells were grown in α-MEM with the above supplements along with 10% (v/v) FBS. Host fibroblast contamination was removed by trypsinisation (0.2%), and cells were passaged routinely every week. Normal oral mucosa tissues were washed in ethanol and PBS as described above and incubated in 2.5 ml of 1% trypsin type III (Sigma) containing 200 iu/ml penicillin and 200 μg/ml streptomycin at 4°C for 16 h followed by incubation at 37°C for 30 min. Five milliliters of α-MEM/FBS was added and the mixture was pipetted vigorously to disaggregate the cells within the tissue. The cells were pelleted and washed in PBS twice and re-suspended in keratinocyte serum-free medium (KSFM, Invitrogen) containing 200 iu/ml penicillin, 200 μg/ml streptomycin, 0.4 ng/ml EGF, 25 μg/ml bovine pituitary extract (BPE) and 30 μM calcium chloride and seeded into 25-cm² culture flasks.

Growth curves. Growth curves were generated for all cells by taking triplicate cell counts with a haemacytometer each time the cells were subcultured. The cell counts were used to calculate the population doubling of the cells.

Immunostaining for cytokeratin. Keratin was examined in all the cultured cells. The cells were grown on glass cover slips, washed twice with PBS and fixed with acetone for 10 min. After 2 PBS washes, the cells were incubated with anti-
cytokeratin monoclonal antibody (1:250 in PBS; Dako Cytomation). To exclude the possibility that the established cell lines were fibroblasts, a separate coverslip with the respective cells was incubated with anti-fibroblast monoclonal antibody (1:250 in PBS; Dako Cytomation). The coverslips were processed using the Dako Cytomation Envision kit following the manufacturer's instructions, and all incubations were carried out at room temperature. The presence of target proteins was visualized using diaminobenzidine (DAB) as a chromogen, and the cells were counter-stained with Mayer's hematoxylin and mounted with DPX mountant. The cells were examined under a light microscope.

Assays of malignancy. The capacity of the established cell lines to grow in semi-solid media in vitro was monitored by soft agar assays as described elsewhere, with minor modifications (8). α-MEM (2.5 ml) supplemented with 10% FBS, 0.4 ng/ml EGF, 2 μg/ml hydrocortisone and 2 mM L-glutamine containing 1% low-gelling agarose (Sigma A 4018) was allowed to set in each 4-ml dish and was overlaid with 2 ml of media containing 0.4% low-gelling agarose and 1% FBS, 0.4 ng/ml EGF, 2 μg/ml hydrocortisone and 2 mM L-glutamine. Triplicate cultures were incubated for each cell line. Two milliliters of media was added on day 7, and after 14 days the number of colonies (>30 cells) was counted in 10 random microscopic fields (x20 objective), and the percentage of cells expressing anchorage independence (colony-forming efficiency; CFE) was calculated.

Immunoblotting. Keratinocytes were grown to semi-confluence in 100-mm culture dishes, following which, the cells were trypsinized, pelleted and lysed in cold RIPA buffer (100 mM NaCl, 10 mM Tris-HCl pH 7.6, 1 mM EDTA, 0.2 mg Aprotinin, 0.1 mM PMSF, 5 mM DTT and 1% Triton X-100) for 20 min on ice. This was spun down at 13,000 rpm for 10 min at 4˚C, and the supernatant containing total cell protein was stored at -70˚C for further analysis. Total protein concentration was determined using the BCA Protein Assay Kit (Pierce, Rockford, IL) and corrected for loading. The blots were probed with primary antibodies (anti-MLH-1 monoclonal antibody (1:250; BD Pharmingen), anti-human MSH-2 (1:250; BD Pharmingen), anti-p53 (1:1000; Santa Cruz Biotechnology), anti-MDM2 (1:250; BD Pharmingen), anti-EGFR (1:200; Clone 31G7, Zymed) for 1 h. Following incubation with the primary antibody, the blots were washed with PBS/0.1% Tween-20 and probed with the secondary antibody conjugated to HRP for 1 h at room temperature, followed by washing with PBS/0.1% Tween-20. To normalize for loading, the blots were probed with anti-actin monoclonal antibody (1:1000; Chemicon) for 1 h at room temperature and processed as described above. The target proteins were detected by enhanced chemiluminescence method (Pierce) and visualized using the ChemiImager™ Imaging Systems (Alpha Innotech).

Analysis of microsatellite instability (MSI). MSI status was determined by screening the polyadenine sequence BAT-26, by PCR amplification of the repeat sequence. PCR was performed with primers 5’ TGACTACTTTGACCTCAGCC 3’ and 5’ AACCATTCAACATTATTACC 3’ as described elsewhere (9). The amplified products were separated on 12% polyacrylamide gel and stained with ethidium bromide.

Methylation specific polymerase chain reaction (MSP) and homozygous deletion of p16INK4a and p14ARF. Genomic DNA was isolated from ORL-48, -115 and -136 cells using the QIAamp DNA mini kit (Qiagen, USA). Methylation of p16INK4a and p14ARF was investigated using the CpG Genome DNA modification kit (Intergen). Following DNA modification, PCR was performed using primers described elsewhere either targeting for wild-type DNA, methylated DNA or unmethylated DNA (10,11). PCR was also performed on unmodified DNA samples. PCR was performed in a 50-μl reaction mixture containing 10 mM Tris-HCl pH 9.0, 50 mM KCl, 1.5 mM MgCl2, and 0.1% Triton X-100, 200 μM dNTPs, 2.5 U TaqDNA polymerase (Promega, Madison, WI, USA) and 100 ng of genomic DNA. PCR amplification was carried out in the GeneAmp PCR System 9700 (Applied Biosystems, USA). The PCR conditions were 95˚C for 2 min, followed by 30 cycles of 95˚C denaturing for 30 sec, 62-68˚C annealing for 30 sec (primer-dependent) and 72˚C elongation for 30 sec. The final extension was conducted for 4 min at 72˚C. Methylated DNA included in the kit was used as a positive control in the DNA modification procedure and PCR. For homozygous deletion analyses of p16INK4a, both exons 1 and 2 were amplified simultaneously with the β-actin gene. PCR conditions were as described above and primer sequences were as previously reported (11). Exon 1B of the p14ARF gene was amplified under the same conditions, and β-globin was coamplified with the 5’ region whilst β-actin was coamplified with the 3’ region of exon 1B of the p14ARF gene, and the amplicons were electrophoresed on a 4% agarose gel for deletion assessment. Primer sequences used were as previously published (12).

Mutational analysis of TP53 and epidermal growth factor receptor (EGFR). Genomic DNA was isolated as described above. Exons 4-10 of the TP53 gene were amplified using PCR. The TP53 primers were designed in the intronic region using the Primer Express software (Applied Biosystems, USA) or as previously published (12). Primer sequences were: exon 4, sense 5’ ACTTCCTGAAAACAACGTTCCT 3’ and antisense 5’ CAGGATAGTGCTCCATGG 3’; exon 9, sense 5’ GAGTATTGCTCCAGATTCA 3’ and antisense 5’ GGCATTGTGAGTGTAGACT 3’; and exon 10, sense 5’ GGTTCTGTGTATATACCTAC 3’ and antisense 5’ ATGAGAAGTGCTCATGG 3’. PCR was performed in a 50-μl reaction mixture containing 10 mM Tris-HCl pH 9.0, 50 mM KCl, 1.5 mM MgCl2, and 0.1% Triton X-100, 200 μM dNTPs, 2.5 U TaqDNA polymerase (Promega) and 100 ng of genomic DNA. PCR amplification was carried out in the GeneAmp PCR System 9700. The PCR conditions were 95˚C for 2 min, followed by 30 cycles of 95˚C denaturing for 30 sec, 62-68˚C annealing for 30 sec (primer-dependent) and 72˚C elongation for 30 sec. The final extension was conducted
for 4 min at 72˚C. Exons 18-21 which encode the tyrosine kinase domain of the EGFR gene were amplified as above with PCR conditions as follows: 95˚C for 2 min, followed by 30 cycles of 95˚C denaturing for 30 sec, 52-58˚C annealing for 30 sec (primer-dependent) and 72˚C elongation for 30 sec. The final extension was conducted for 4 min at 72˚C. PCR primers used were as reported by Lynch and colleagues (13). TP53 and EGFR PCR products were purified using either the QIAamp PCR purification kit (Qiagen, Germany) or the QIAamp gel extraction kit (Qiagen). Purified products were sequenced using the DYEnamic ET Terminator cycle sequencing kit (Amersham Biosciences) and the primers used in the cycle sequencing reactions were similar to those used in PCR. Both DNA strands were sequenced, and the cycle sequencing products were electrophoresed on the MegaBACE™ 500 automated sequencer (Amersham Biosciences). Sequences for the individual exons were checked for mutations by comparing them to known normal sequences from GenBank.

HPV analysis. The presence of HPV was determined by PCR using a combination of the general primers GP5+ and GP6+ which amplify the L1 region from all types of HPV genomes (14). PCR was performed in a 50-μl reaction mixture containing 10 mM Tris-HCl pH 9.0, 50 mM KCl, 1.5 mM MgCl₂, 0.1% Triton X-100, 200 μM dNTPs, 2.5 U TaqDNA polymerase (Promega) and 100 ng of genomic DNA. PCR amplification was carried out in the GeneAmp PCR System 9700. The PCR conditions were 94˚C for 2 min, followed by 30 cycles of 94˚C denaturing for 30 sec, 40˚C annealing for 2 min and 72˚C elongation for 1.5 min. The final extension was conducted for 7 min at 72˚C. HPV DNA clones which were kind gifts from Professor de Villiers were included as positive controls for the PCR.

Karyotyping of cell lines. ORL-48, -115 and -136 cell lines were treated with uridine (0.4 mM) and fluorodeoxyuridine (5 μM) for 16 h at 37˚C, followed by colcemid (10 μg/ml) for 5 h at 37˚C. The adherent cells were removed with a cell scraper and treated with hypotonic KC1 (0.075 M) for 30 min followed by fixation in fresh methanol:glacial acetic acid (3:1, v/v) for 10 min at room temperature. Slides were stained by the trypsin-Giemsa technique and at least 20 metaphases were evaluated for each cell line.

Results

Behaviour of the in vitro cultures. All three oral squamous cell carcinoma (OSCC) cell lines grew as monolayers in culture with the population doubling ranging between 26.4 and 40.8 h. ORL-48, -115 and -136 cells are immortal and currently at passage 28, 20 and 24, respectively. The OSCC cells are typically polygonal and the morphological features of the cultured cells are shown in Fig. 1a. These OSCC cell lines stained positive for cytokeratin confirming its keratinocyte origin (Fig. 1b). Soft agar assays demonstrated that all 3 cell lines expressed a low degree of anchorage independency (CFE <6%).

Expression of MDM2 and EGFR. Overexpression of MDM2 has previously been shown to be a frequent event in oral cancers (12). Elevated levels of MDM2 protein in
comparison to normal oral keratinocytes were observed in ORL-48 (Fig. 2). The expression of EGFR is prominent in a variety of solid tumours including head and neck squamous cell carcinomas and is associated with a more aggressive disease, resistance to therapy, increased metastasis and poor clinical outcome. However, overexpression of EGFR in comparison to normal oral keratinocytes was only observed in ORL-136 and not in ORL-48 and -115 (Fig. 2).

Analysis of microsatellite instability. Since losses of MLH-1 and MSH-2 expression are known causes of inactivated mismatch repair function, the expression of these genes was analyzed. All 3 cell lines demonstrated the expression of MLH-1 and MSH-2 repair proteins (Fig. 2) suggesting that these enzymes are functional. In addition, the polyadenine sequence BAT-26 was analyzed using PCR to determine the microsatellite instability status of our cells. None of the cell lines demonstrated microsatellite instability either at the BAT-26 polyadenine region (Fig. 3) or by the loss of MLH-1 and MSH-2 expression.

Analyses of cell cycle regulators and tumour suppressors. Methylation and gene deletion are mechanisms of inactivating the tumour suppressors \( p16^{INK4a} \) and \( p14^{ARF} \). Gene deletion and the methylation status of both genes were investigated in ORL-48, -115 and -136. \( p16^{INK4a} \) and \( p14^{ARF} \) genes were amplified simultaneously either with \( \beta\)-actin or \( \beta\)-globin. Control cells are known to have both \( p16^{INK4a} \) and \( p14^{ARF} \) intact.

Figure 3. Analysis of the polyadenine repeat sequence BAT-26. The BAT-26 region was amplified and separated in a denaturing polyacrylamide gel and stained with ethidium bromide. Amplicons from the cell lines (CL) were compared to the same region amplified from peripheral blood DNA (PB) of the same individual, and no alterations in the BAT-26 repeat sequence was found in any of the cell lines. DNA extracted from a normal oral keratinocyte cell line (ORL-52) was included.

Figure 4. Agarose gel electrophoresis demonstrating that \( p16^{INK4a} \) and \( p14^{ARF} \) genes were intact in ORL-48, -115 and -136. \( p16^{INK4a} \) and \( p14^{ARF} \) genes were amplified simultaneously either with \( \beta\)-actin or \( \beta\)-globin. Control cells are known to have both \( p16^{INK4a} \) and \( p14^{ARF} \) intact.

Figure 5. Sequence analysis of the \( TP53 \) gene. i) Electropherogram of the \( TP53 \) gene showing 2 different mutations in ORL-48. (a) c.876delA in exon 8 and (b) p.Glu336X in exon 10 (sequence in antisense direction). ii) Electrophorogram of the \( TP53 \) gene showing the p.Cys176Phe alteration in ORL-136. iii) Electrophorogram of the \( TP53 \) gene showing the c.336-338delCTT in-frame deletion in ORL-115. All mutations were confirmed by sequencing an independent PCR amplicon.
ORL-48, a deletion of an adenine was seen in nucleotide 876 in exon 8, and a heterozygous mutation at nucleotide 1006 resulted in a stop codon in exon 10 (Fig. 5). In addition to the mutations observed in ORL-48, a polymorphism resulting in an arginine to proline amino acid change was detected in exon 4 which may explain the slight alteration in the molecular weight of the TP53 protein (Fig. 2). A 3-base in-frame deletion in exon 4 (nt 336-338) was detected in ORL-115, while a G to T transition at nucleotide 527 in exon 5 was identified in ORL-136 resulting in a cysteine to phenylalanine amino acid change in codon 176 (Fig. 5). Mutations in exons 18-21 of the EGFR gene were undetected in these cell lines.

HPV infection. HPV infection has been implicated in oral carcinogenesis. HPV infection in ORL-48, -115 and -136 was evaluated using PCR. None of the cell lines were infected with HPV.

Chromosome analyses. Chromosomal analysis of ORL-48 and -115 showed human female karyotype whilst ORL-136 showed a human male karyotype. All cell lines had an abnormal number of chromosomes ranging between 64 and 82 with random gains or losses of entire chromosomes or regions of the chromosomes (Fig. 6). The composite karyotype of each cell line is detailed in Table II.

Discussion

While the difficulties in developing oral cancer cell lines have been well-reported, the importance of these cell lines in the understanding of oral cancer biology is also well-recognized. Several previously proposed reasons explaining the difficulties in establishing viable cultures include a limited tumour stem cell population and unfavorable culture conditions introduced by 3T3 feeder cells (15,16). Here we described the establishment and characterization of 3 OSCC cell lines established from individuals with and without habits associated with oral cancer. In our laboratory, the cultures appeared to be more viable when 3T3 feeder cells were not used, therefore we resorted to establishing our cultures without the use of feeder layers. These cell lines grew as monolayers of polygonal cells and were shown to be immortal. Although the keratinocytes were more viable without the feeder cells, the lack of these cells encouraged the growth of host fibroblasts which had to be frequently removed with mild trypsinisation. We confirmed the human origin of these cells by karyotyping and demonstrated that the cells had near triploid and tetraploid number of chromosomes and that they gained and lost entire chromosomes or regions of chromosomes (Table II), a common characteristic of solid tumours (17). The epithelial origin of these cells was confirmed by cytokeratin expression, and the contamination

Table II. Karyotype of ORL-48, -115 and -136 cell lines.

<table>
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<th>Cell line</th>
<th>Karyotype</th>
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<tr>
<td>ORL-48</td>
<td>64-73.XXX,+X,der(1)t(1;2)(p10;q10),+del(1)(p22),+del(3)(p13),+del(4)(q21),+der(4)hsrc(4)(q21q31),-8,+i(9)(q10),-10,-10,-11,-15,-14,+i(17)(q10),+20,-22,+mar[15]</td>
</tr>
<tr>
<td>ORL-115</td>
<td>70-80.XX,X,+der(1)t(1;12)(p11;q11),+del(1)(p13),+2,+3,+5,+8,der(8)(8;7)(q24;7),-9,+i(10)(q10),+11,+11,+11,-12,-13,add(13)(q34),+14,+16,+17,+18,+19,+20,+20,+21,-22,+mar1,+mar2[cp10]</td>
</tr>
<tr>
<td>ORL-136</td>
<td>74-82.XX,YY,add(2)(p25),del(6)(q21),-7,-8,+i(8)(q10),der(10)t(3;10)(p10;q10),del(10)(p10),+11,+13,+14,+15,+17,+18,+19,+20,+20,+22,-22,-22,+mar1,+mar2[cp18]</td>
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Figure 6. Karyotype analysis of ORL-48 cells showing a female karyotype with random gains and losses of regions and whole chromosomes.
of fibroblast cells in these cultures was negated by negative staining with an anti-fibroblast monoclonal antibody (data not shown). Anchorage independency has been reported to be indicative of a transformed phenotype by which cells can form tumours in vivo (18). None of these cell lines formed colonies in semi-solid agarose suggestive of non-transformed phenotypes, in contrast to previous reports of other OSCC cell lines readily exhibiting anchorage independent growth (19,20). It is well-recognized that cancers develop as a result of the accumulation of sequential genetic alterations. As the TP53 and retinoblastoma (RB1) pathways are central in the control of cellular proliferation and senescence, it is not surprising that alterations within these pathways are frequently detected in cancers. We have examined our cell lines for the common genetic alterations reported in genes known to play major roles in the carcinogenesis process. Genetic characterization showed TP53 mutations in all 3 cell lines resulting either in a stop codon (in ORL-48) or amino acid changes (ORL-136). Two of these mutations (p.Glu336X and p.Cys176Phe) have previously been reported in head and neck tumours or cell lines (21-25). MDM2, a negative regulator of TP53 that has been reported in head and neck tumours or cell lines (21-25), has also been demonstrated to be over-expressed in oral cancers (5,12,27). Surprisingly, amongst our cell lines, only ORL-48 demonstrated overexpression of the MDM2 protein (26), which shares a similar locus to p14ARF previously reported to extend cellular lifespan and confer immortality in fibroblasts (30,31,32).

HPV infection has been implicated in a fraction of head and neck cancers (33), and frequently, HPV infection is detected in tumours with an intact TP53 pathway (34). However, the absence of HPV infection in all 3 cell lines suggests that HPV did not play a role in the pathogenesis of oral cancers in these particular individuals, indicating that the growth advantage of these tumours had already been provided by the inactivation of TP53. EGFR has been reported to be overexpressed in head and neck cancers and has been associated with poor clinical outcome (35-37). In view of this, EGFR has been targeted for the treatment of cancers, including head and neck (38-40). While EGFR overexpression is a common event in many cancers of epithelial origin, mutations within the tyrosine kinase domain of this gene were also unobserved in the cell lines. Microsatellite instability analyses indicated that all 3 cell lines were microsatellite stable with expression of both DNA mismatch repair enzymes (MLH-1 and MSH-2) intact. Similar genetic analyses were performed on the tissues from which these cell lines were derived with similar results, indicating that these cell lines closely resembled the tissues (data not shown). These cell lines which are genetically representative of the tissues from which they are derived will serve as useful tools in investigating the mechanisms of oral cancer development.

Acknowledgements

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References