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Tumour-suppressive *microRNA-29s* inhibit cancer cell migration and invasion by targeting laminin–integrin signalling in head and neck squamous cell carcinoma

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Background: Our recent studies of microRNA (miRNA) expression signatures demonstrated that *microRNA-29s* (*miR-29s; miR-29a/b/c*) were significantly downregulated in head and neck squamous cell carcinoma (HNSCC) and were putative tumoursuppressive miRNAs in human cancers. Our aim in this study was to investigate the functional significance of *miR-29s* in cancer cells and to identify novel *miR-29s*-mediated cancer pathways and responsible genes in HNSCC oncogenesis and metastasis.

Methods: Gain-of-function studies using mature *miR-29s* were performed to investigate cell proliferation, migration and invasion in two HNSCC cell lines (SAS and FaDu). To identify *miR-29s*-mediated molecular pathways and targets, we utilised gene expression analysis and *in silico* database analysis. Loss-of-function assays were performed to investigate the functional significance of *miR-29s* target genes.

Results: Restoration of *miR-29s* in SAS and FaDu cell lines revealed significant inhibition of cancer cell migration and invasion. Gene expression data and *in silico* analysis demonstrated that *miR-29s* modulated the focal adhesion pathway. Moreover, laminin γ^2 (*LAMC2*) and α 6 integrin (*ITGA6*) genes were candidate targets of the regulation of *miR-29s*. Luciferase reporter assays showed that *miR-29s* directly regulated *LAMC2* and *ITGA6*. Silencing of *LAMC2* and *ITGA6* genes significantly inhibited cell migration and invasion in cancer cells.

Conclusion: Downregulation of *miR-29s* was a frequent event in HNSCC. The *miR-29s* acted as tumour suppressors and directly targeted laminin–integrin signalling. Recognition of tumour-suppressive miRNA-mediated cancer pathways provides new insights into the potential mechanisms of HNSCC oncogenesis and metastasis and suggests novel therapeutic strategies for the disease.

Head and neck squamous cell carcinoma (HNSCC) is the sixth most common cancer in the world and \sim 500 000 cases are diagnosed every year (Jemal *et al*, 2010). In spite of considerable advances in multimodality therapy, including surgery, radio-therapy and chemotherapy, the overall 5-year survival rate for patients with HNSCC is only 40–50% (Leemans *et al*, 2011).

Local tumour recurrence and distant metastasis after conventional therapy appear to be major contributing factors for restricted survival of HNSCC patients. Therefore, understanding the molecular pathways of metastasis accompanying HNSCC would help to improve diagnosis, approaches to therapy and prevention of the disease.

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The discovery of noncoding RNAs in the human genome was an important conceptual breakthrough in the postgenome sequencing era (Mattick, 2004). The microRNAs (miRNAs) constitute a class of small, noncoding RNA molecules (19–22 nucleotides in length) that regulate protein-coding gene expression by repressing translation or cleaving RNA transcripts in a sequence-specific manner (Bartel, 2004). The miRNAs are unique in their ability to regulate multiple protein-coding genes. Bioinformatic predictions indicate that miRNAs regulate > 30% of the protein-coding genes in the human genome (Lewis *et al*, 2005).

A growing body of evidence suggests that miRNAs contribute significantly to HNSCC progression, development and metastasis (Esquela-Kerscher and Slack, 2006). In cancer pathways, normal regulatory mechanisms are disrupted by altered expression of tumour-suppressive or oncogenic miRNAs. Therefore, identification of differentially expressed miRNAs is an important step to understand human oncogenesis. Genome-wide miRNA expression signatures have been used to rapidly and precisely identify aberrant miRNA expression in the HNSCC. Based on recent reports, including our own signatures, several down- or upregulated miRNAs have been discovered in the HNSCC (Avissar *et al*, 2009; Hui *et al*, 2010; Kikkawa *et al*, 2010; Nohata *et al*, 2011).

Our expression signature of HNSCC demonstrated that the microRNA-29 family (miR-29s) was downregulated in cancer tissues (Kikkawa et al, 2010; Nohata et al, 2011). The miR-29s include three members, miR-29a, miR-29b and miR-29c, in the human genome. The miR-29b includes two members, miR-29b-1 and miR-29b-2 (Wang et al, 2013). These miRNAs form clusters on two human genome loci: miR-29b-1 and miR-29a in 7q32 and miR-29b-2 and miR-29c in 1q32 (Kriegel et al, 2012). Although aberrant expression of miR-29s was reported in several types of human cancers, the expression status of miR-29s varied according to the cancer types (Fabbri et al, 2007; Sengupta et al, 2008; Gebeshuber et al, 2009; Han et al, 2010; Mott et al, 2010; Xiong et al, 2010; Zhao et al, 2010; Li et al, 2011; Fu et al, 2012; Presneau et al, 2013). Therefore, the aim of this study was to investigate the functional significance of the miR-29s in cancer cells and to identify novel miR-29s-regulated cancer pathways and responsible genes in HNSCC oncogenesis and metastasis.

Genome-wide gene expression analysis of miR-29a transfectants and in silico database analysis showed that the focal adhesion pathway was a promising candidate target pathway. In this regard, the laminin-integrin signalling is critically important. It occurs in the basal lamina and influences cell differentiation, migration and adhesion as well as proliferation and cell survival. Interestingly, laminin $\gamma 2$ (coded by LAMC2), a member of laminin-332, and $\alpha 6$ integrin (coded by ITGA6) were listed as candidate genes in the focal adhesion pathway. Moreover, an interaction between laminin-332 and $\alpha 6\beta 4$ integrin triggers a number of signalling cascades, promoting both cell migration and cancer cell survival (Marinkovich, 2007; Kariya et al, 2009). We investigated the functional significance of LAMC2 and ITGA6 in HNSCC and showed that these genes were regulated by miR-29s. The identification of novel cancer pathways regulated by tumoursuppressive miR-29s provides new insights into the potential mechanisms of HNSCC oncogenesis and metastasis.

MATERIALS AND METHODS

Clinical HNSCC specimens. A total of 23 pairs of primary HNSCC and corresponding normal epithelial samples were obtained from patients with HNSCC at Chiba University Hospital (Chiba, Japan) from 2005 to 2011. The samples considered normal were free of cancer cells by pathologic examination. The patients' backgrounds and clinicopathological characteristics are summarised in Supplementary Table 1. The patients were classified according to the 2002 Union for International Cancer Control (UICC) TNM staging criteria before treatment. Written consent for tissue donation for research purposes was obtained from each patient before tissue collection. The protocol was approved by the institutional review board of the Chiba University. The specimens were immersed in RNAlater (Qiagen, Valencia, CA, USA) and stored at -20 °C until RNA was extracted.

RNA isolation. Total RNA was isolated using TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's protocol. The RNA concentrations were determined spectrophotometrically, and molecular integrity was checked by gel electrophoresis. The quality of RNA was confirmed using an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA).

HNSCC cell culture. The following human HNSCC cell lines were used: SAS (derived from a primary lesion of tongue SCC) and FaDu (derived from a primary lesion of hypopharyngeal SCC). Both cell lines were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum in a humidified atmosphere containing 5% CO₂ at 37 °C.

Quantitative real-time RT-PCR (qRT-PCR). First-strand cDNA was synthesised from $1 \mu g$ of total RNA using a High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA, USA). Gene-specific PCR products were assayed continuously using a 7900-HT Real-Time PCR System according to the manufacturer's protocol. The initial PCR step consisted of a 10 min hold at 95 °C, followed by 40 cycles consisting of 15 s denaturation at 95 °C and 1 min annealing/extension at 60 °C. The expression levels of miR-29a (Assay ID: 002112), miR-29b (Assay ID: 000413) and miR-29c (Assay ID: 000587) were analysed by TaqMan quantitative real-time PCR (TaqMan MicroRNA Assay; Applied Biosystems) and normalised to RNU48 (Assay ID: 001006). TaqMan probes and primers for LAMC2 (P/N: Hs01043711_m1), ITGA6 (P/N: Hs01041011_m1) and GUSB (P/N: Hs99999908_m1) as an internal control were obtained from Applied Biosystems (Assay-On-Demand Gene Expression Products). The $\Delta \Delta Ct$ method was adopted and applied to calculate the relative quantities of subject genes. All reactions were performed in triplicate, and included negative control reactions that lacked cDNA.

Mature miRNA transfection and small interfering RNA treatment. The following mature miRNAs species were used in this study: mirVana miRNA mimic for hsa-miR-29a-3p (product ID: MC12499), hsa-miR-29b-3p (product ID: MC10103) and hsa-miR-29c-3p (product ID: MC10518) (Applied Biosystems). The following siRNAs were used: Stealth Select RNAi siRNA; si-*LAMC2* (Cat no. HSS105965), si-*ITGA6* (Cat no. HSS179959) (Invitrogen) and negative control miRNA/siRNA (P/N: AM17111, Applied Biosystems). The RNAs were incubated with OPTI-MEM (Invitrogen) and Lipofectamine RNAiMax reagent (Invitrogen) as described previously (Ichimi *et al*, 2009). Transfection efficiency of miRNA in cell lines was confirmed based on downregulation of *TWF1* (*PTK9*) mRNA following transfection with *miR-1* as previously reported (Ichimi *et al*, 2009).

Cell proliferation, migration and invasion assays. Cells were transfected with 10 nM miRNA or siRNA by reverse transfection and plated in 96-well plates at 3×10^3 cells per well. After 72 h, cell proliferation was determined with the XTT assay, using the Cell Proliferation Kit II (Roche Molecular Biochemicals, Mannheim, Germany) as previously reported (Chiyomaru *et al*, 2010).

A cell migration assay was performed using BD Falcon Cell Culture Inserts (BD Biosciences, Franklin Lakes, NJ, USA) that contained uncoated Transwell polycarbonate membrane filters

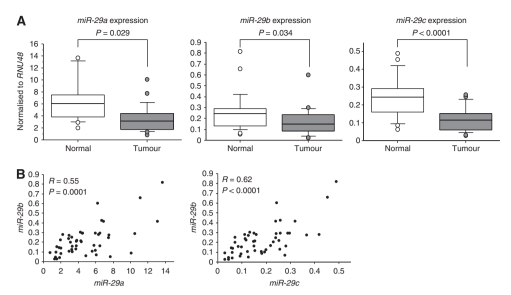


Figure 1. Expression levels of miR-29s in HNSCC clinical specimens. (A) Expression levels of miR-29a, miR-29b and miR-29c in HNSCC clinical specimens were measured by qRT–PCR. The RNU48 was used for normalisation. (B) Correlations between miR-29a-miR-29b and miR-29b-miR-29c were determined in HNSCC specimens.

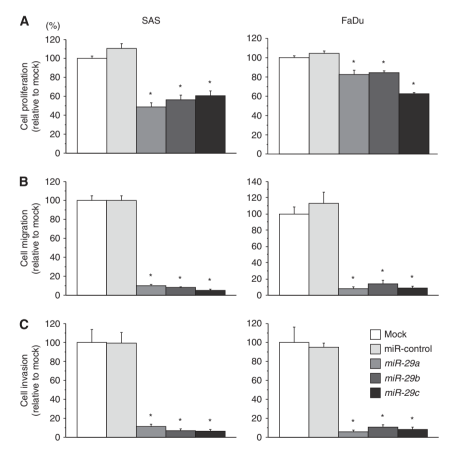


Figure 2. Effect of *miR-29s* transfection on cell proliferation, migration and invasion in SAS and FaDu. (A) Cell proliferation 72 h after transfection with *miR-29s* (10 nM) was determined with the XTT assay. (B) Cell migration activity 48 h after transfection with *miR-29s* (10 nM) was determined with the migration assay. (C) Cell invasion activity 48 h after transfection with *miR-29s* (10 nM) was determined with the Matrigel invasion assay. **P* < 0.005.

with 8 μm pores in 24-well tissue culture plates. Cells were transfected with 10 nm miRNA or siRNA by reverse transfection and plated in 10 cm dishes at 8 \times 10⁵ cells. After 48 h, the cells were collected and 2 \times 10⁵ cells were added to the upper chamber of

each migration well and were allowed to migrate for 48 h. After gentle removal of the nonmigratory cells from the filter surface of the upper chamber, the cells that migrated to the lower side were fixed and stained with Diff-Quick

(Sysmex Corporation, Kobe, Japan). The number of cells that migrated to the lower surface was determined microscopically by counting four areas of constant size per well.

A cell invasion assay was carried out using modified Boyden chambers containing Transwell-precoated Matrigel membrane filter inserts with $8\,\mu m$ pores in 24-well tissue culture plates at 2×10^5 cells per well (BD Biosciences). All experiments were performed in triplicate.

In silico analysis, genome-wide gene expression analysis and miR-29s target search. To identify miR-29s target genes, we used in silico analysis and genome-wide gene expression analysis. First, we screened genes using TargetScan Release 6.2 (http://www.targetscan.org/). These genes were then categorised into KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways using GENECODIS analysis (Tabas-Madrid *et al*, 2012) (http://genecodis. cnb.csic.es/). To identify upregulated genes in HNSCC, we analysed a publicly available gene expression data set in GEO (Accession number: GSE9844). To gain further insight into which genes were affected by miR-29s, we performed genome-wide gene expression analysis using miR-29a transfection of SAS. SurePrint G3 Human GE 8×60 K Microarray (Agilent Technologies) was used for expression profiling of miR-29a transfectants in comparison with negative control miRNA transfectants.

Western blotting. Cells were harvested 72 h after transfection and lysates were prepared. From each lysate, 50 μ g of protein was separated on a Mini-PROTEAN TGX gel (Bio-Rad, Hercules, CA, USA) and transferred to PVDF membranes. Immunoblotting was performed with mouse LAMC2 antibody (1:500, MAB19562, Merck Millipore, Billerica, MA, USA) and rabbit ITGA6 antibody (1:1000, ab97760, Abcam, Cambridge, UK) with GAPDH antibody (1:1000, ab8245, Abcam) used as an internal control. The membrane was washed and incubated with anti-mouse IgG HRP-linked antibody (7076, Cell Signaling Technology, Danvers, MA, USA) or anti-rabbit IgG HRP-linked antibody (7074, Cell Signaling Technology). Complexes were visualised with an Immun-Star WesternC Chemiluminescence Kit (Bio-Rad), and the expression levels of these genes were evaluated by ImageJ software (ver.1.44; http://rsbweb.nih.gov/ij/).

Plasmid construction and dual-luciferase reporter assay. The partial wild-type sequences of the *LAMC2* 3'-untranslated region (3'-UTR) or those with deleted *miR-29s* target sites (positions 611–627 and 1430–1436 of *LAMC2* 3'-UTR) were inserted between the *XhoI–PmeI* restriction sites in the 3'-UTR of the *hRluc* gene in psiCHECK-2 vector (C8021; Promega, Madison, WI, USA). The same was done with the partial wild-type sequences of the *ITGA6* 3'-UTR or those with deleted *miR-29s* target site (positions 1570–1576 of *ITGA6* 3'-UTR). The sequences of the oligonucleotides were described in the Supplementary Document. The SAS cells were transfected with 5 ng of vector and 10 nm of *miR-29s* using Lipofectamine 2000 (Invitrogen). The activities of firefly and *Renilla* luciferases in cell lysates were determined with a dual-luciferase assay system (E1910; Promega). Normalised data were calculated as the quotient of *Renilla*/firefly luciferase activities.

Statistical analysis. The relationships between two groups and the numerical values obtained by real-time RT–PCR were analysed using the paired *t*-test. Spearman's rank test was used to evaluate the correlation between the expressions of *miR-29s*. The relationship among more than three variables and numerical values were analysed using the Bonferroni adjusted Mann–Whitney *U*-test. All analyses were performed using Expert StatView (version 4, SAS Institute Inc., Cary, NC, USA).

Expression of *miR-29s* **in clinical HNSCC specimens.** The chromosomal locations of *miR-29s* in the human genome are shown in Supplementary Figure 1. These miRNAs are clustered at two different human genomic loci, *miR-29b-1* and *miR-29a* at 7q32.3 and *miR-29b-2* and *miR-29c* at 1q32.2.

To validate our past miRNA profiling results, we evaluated *miR-29s* expression in 23 clinical HNSCC specimens. The expression levels of *miR-29a*, *miR-29b* and *miR-29c* were significantly lower in tumour tissues than in corresponding adjacent noncancerous epithelia (P = 0.029, P = 0.034 and P < 0.0001, respectively, Figure 1A). Spearman's rank test showed positive correlation between the expression of *miR-29a* and that of *miR-29b* (R = 0.55 and P = 0.0001, Figure 1B). Similarly, the expression of *miR-29c* was positively correlated with that of *miR-29b* (R = 0.62 and P < 0.0001, Figure 1B).

Effect of restoring *miR-29s* on cell proliferation, migration and invasion activities in HNSCC cell lines. To investigate the functional effects of *miR-29s*, we performed gain-of-function studies using miRNA transfection of SAS and FaDu cell lines. The XTT assay demonstrated that cell proliferation was significantly inhibited in *miR-29s* transfectants in comparison with the mock or miR-control transfectant cells. Specifically, we observed the following growth, expressed as a percentage of the mock: (1) SAS–Mock, 100.0 ± 2.4 ; miR-control, 110.8 ± 4.9 ; *miR-29a*, 48.6 ± 4.2 ; *miR-29b*, 56.4 ± 5.0 ; *miR-29c*, 60.7 ± 4.9 ; (2) FaDu–Mock, 100.0 ± 1.9 ; miR-control, 104.5 ± 2.6 ; *miR-29a*, 82.3 ± 4.4 ; *miR-29b*, 84.5 ± 1.8 ; *miR-29c*, 62.7 ± 0.7 , with *P*<0.005 for both (Figure 2A).

The migration assay demonstrated that cell migration activity was significantly inhibited in *miR-29s* transfectants in comparison

Table 1. Candidate target genes for miR-29s in focal adhesion pathway				
	GEO expression data (GSE9844)			Gene expression analysis of SAS
Gene symbol	Entrez gene ID	Log ₂ ratio (tumour/ normal)	P-value	Log ₂ ratio (<i>miR-29a/</i> miR-control)
LAMC2	3918	3.27	3.91E-06	- 1.18
COL4A6	1288	2.60	2.12E-06	- 0.87
COL4A1	1282	2.26	2.11E-04	- 1.78
ITGA6	3655	2.07	7.10E-06	- 1.62
COL11A1	1301	1.97	1.88E-03	- 0.75
COL5A2	1290	1.74	3.87E-04	- 3.18
COL4A2	1284	1.63	1.22E-03	- 1.76
COL1A2	1278	1.45	6.95E-04	- 0.61
COL6A3	1293	1.35	5.72E-03	- 0.80
COL5A1	1289	1.23	1.69E-03	- 1.56
COL3A1	1281	1.18	9.15E-03	- 0.78
COL4A5	1287	1.14	5.52E-04	- 0.96
COL5A3	50509	0.76	9.15E-03	- 0.70
THBS2	7058	0.76	6.85E-02	- 1.14
CAV2	858	0.55	5.15E-02	- 0.81
CDC42	998	0.54	7.60E-03	- 0.70
Abbreviations: $GEO = Gene$ Expression Omnibus; miR-29 = microRNA-29; SAS = head and				

neck squamous cell carcinoma cell line.

miR-29s target laminin-integrin in HNSCC

with the mock or miR-control transfectant cells. Specifically, we observed the following migration activities, expressed as a percentage of the mock: (1) SAS–Mock, 100.0 ± 5.3 ; miR-control, 100.2 ± 4.8 ; *miR-29a*, 10.2 ± 1.3 ; *miR-29b*, 7.9 ± 0.9 ; *miR-29c*, 5.2 ± 0.8 ; (2) FaDu–Mock, 100.0 ± 8.7 ; miR-control, 113.4 ± 13.8 ; *miR-29a*, 8.3 ± 4.4 ; *miR-29b*, 14.0 ± 4.0 ; *miR-29c*, 8.6 ± 2.3 , with *P* < 0.005 for both (Figure 2B).

The Matrigel invasion assay demonstrated that cell invasion activity was significantly inhibited in *miR-29s* transfectants in comparison with the mock or miR-control transfectant cells. Specifically, we observed the following migration activities, expressed as a percentage of the mock: (1) SAS-Mock, 100.0 ± 13.6 ; miR-control, 99.2 ± 11.4 ; *miR-29a*, 11.1 ± 2.8 ; *miR-29b*, 6.9 ± 1.8 ; *miR-29c*, 6.4 ± 1.8 and (2) FaDu-Mock, 100.0 ± 16.3 ; miR-control, 94.9 ± 4.4 ; *miR-29a*, 5.8 ± 1.8 ; *miR-29b*, 10.9 ± 2.0 ; *miR-29c*, 8.3 ± 2.2 , with P < 0.005 for both (Figure 2C).

Selection of candidate genes targeted by *miR-29s*. To identify genes targeted by *miR-29s*, we used *in silico* analysis and genomewide gene expression analysis. Our strategy for selection of *miR-29s*-targeted genes is shown in Supplementary Figure 2. First, we screened *miR-29s*-targeted genes using the TargetScan database and identified 2627 genes (Supplementary Table 2). These genes were then categorised into KEGG pathways using GENECODIS analysis and 83 pathways were identified as significantly enriched pathways (Supplementary Table 3). Among these pathways, we focussed on the focal adhesion pathway because this pathway was implicated in cancer cell migration and invasion. A total of 58 genes were identified in this pathway (Supplementary Table 4). The gene set was then analysed with a publicly available gene expression data set in GEO (accession number: GSE9844) and genes upregulated (log₂ ratio > 0.5) in HNSCC were chosen. To gain further insight into which genes were affected by *miR-29s*, we performed genome-wide gene expression analysis using SAS, and genes downregulated (log₂ ratio < -0.5) by *miR-29a* transfection were selected. Entries from the gene expression data were approved by GEO, and were assigned GEO accession number GSE47657. As a result, 16 candidate genes were identified as *miR-29s* targets in the focal adhesion pathway (Table 1). Among those candidates, we focussed on *LAMC2* (a component of laminin-332), which is a specific ligand of *ITGA6* (a component of $\alpha \beta \beta 4$ integrin). Laminin 332- $\alpha \beta \beta 4$ integrin interaction is known to promote tumour invasion and cell survival through PI3K and RAC1 activation. Therefore, we focussed on *LAMC2* and *ITGA6* in further analyses.

LAMC2 and *ITGA6* were directly regulated by *miR-29s*. We performed qRT–PCR and western blotting in SAS to investigate whether *LAMC2* and *ITGA6* expression was downregulated by restoration of *miR-29s*. The mRNA and protein expression levels of *LAMC2*/LAMC2 were significantly repressed in *miR-29s* transfectants in comparison with mock or miR-control transfectants (P<0.005, Figure 3A and B). Similarly, the mRNA and protein expression levels of *ITGA6*/ITGA6 were significantly repressed in *miR-29s* transfectants (P<0.005, Figure 4A and B).

We performed a luciferase reporter assay in SAS to determine whether *LAMC2* and *ITGA6* mRNA had target sites for *miR-29s*. The TargetScan database predicted that two putative *miR-29s*binding sites existed in *LAMC2* 3'-UTR (positions 611–627 and

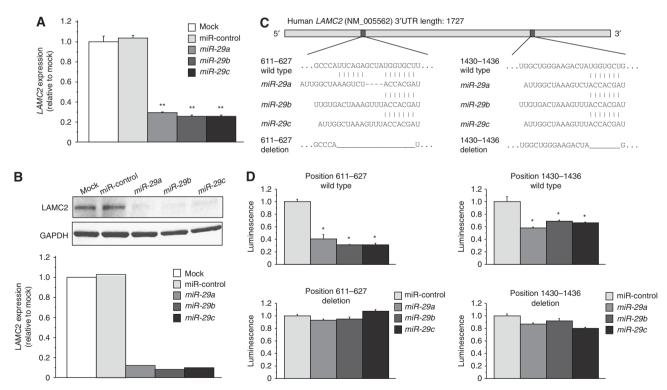


Figure 3. Direct regulation of LAMC2 by miR-29s. (A) The LAMC2 mRNA expression 72 h after transfection with 10 nm miR-29s in SAS was evaluated with qRT–PCR. As an internal control, GUSB was used. **P<0.005. (B) LAMC2 protein expression 72 h after transfection with miR-29s in SAS was evaluated with western blot analysis. As a loading control, GAPDH was used. The expression ratio of LAMC2/GAPDH was calculated with ImageJ software. (C) TargetScan database analysis indicated two putative miR-29s-binding sites existed in the LAMC2 3'-UTR (positions 611–627 and 1430–1436). With regard to these two sites, partial wild-type sequences of LAMC2 3'-UTR and those with the deleted miR-29s target sites were constructed and inserted between the Xhol and Pmel restriction sites in the 3'-UTR of the hRluc gene in the psiCHECK-2 vector. (D) The vectors were co-transfected with miR-29s or miR-control into SAS cells. Firefly luciferase activity was normalised to Renilla luciferase activity. Relative luciferase activity in miR-29s transfectants was compared with that in miR-control cultures, which was set at 1, in cells transfected with wild type 3'-UTR or deletion 3'-UTR. *P<0.0083.

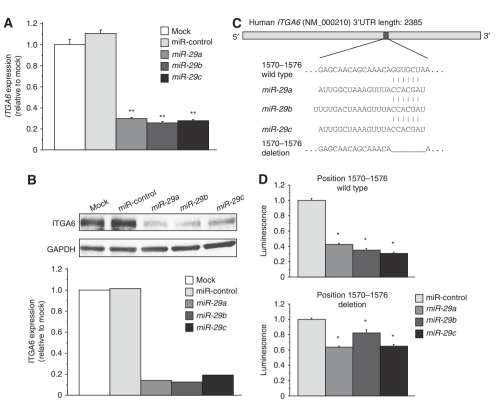


Figure 4. Direct regulation of *ITGA6* by *miR-29s*. (A) The *ITGA6* mRNA expression 72 h after transfection with 10 nm miR-29s in SAS was evaluated with qRT–PCR. As an internal control, *GUSB* was used. **P<0.005. (B) ITGA6 protein expression 72 h after transfection with *miR-29s* in SAS was evaluated with western blot analysis. As a loading control, GAPDH was used. The expression ratio of ITGA6/GAPDH was calculated with ImageJ software. (C) TargetScan database analysis indicated a putative *miR-29s*-binding site existed in the *ITGA6* 3'-UTR (positions 1570–1576). With regard to this site, partial wild-type sequences of *ITGA6* 3'-UTR and those with the deleted *miR-29s* target site were constructed and inserted between the *Xhol* and *Pmel* restriction sites in the 3'-UTR of the h*Rluc* gene in psiCHECK-2 vector. (D) The vectors were co-transfected with *miR-29s* or miR-control into SAS cells. Firefly luciferase activity was normalised to *Renilla* luciferase activity. Relative luciferase activity in *miR-29s* transfectants was compared with that in miR-control cultures, which was set at 1, in cells transfected with wild-type 3'UTR or deletion 3'-UTR. *P<0.0083.

1430–1436, Figure 3C). We used vectors encoding either the partial wild-type sequence of the 3'-UTR of LAMC2 mRNA, including the predicted miR-29s target sites, or 'deletion' vectors, that is, those lacking the miR-29s target sites. We found that the luminescence intensity was significantly reduced by transfection of miR-29s with the vector carrying the wild-type 3'-UTR of LAMC2, whereas transfection with deletion vectors (positions 611-627 or 1430-1436 had been removed) blocked the decrease in luminescence (P < 0.0083, Figure 3D). As for ITGA6, the TargetScan database predicted a putative miR-29s-binding site in ITGA6 3'-UTR (positions 1570-1576). We constructed a vector encoding either the partial wild-type sequences of 3'-UTR of ITGA6 or those with a deleted *miR-29s* target site (Figure 4C). The luminescence intensity was significantly reduced by transfection of miR-29s with the vector of wild-type 3'-UTR of LAMC2 (P<0.0083, Figure 4D). Although the luminescence intensity was also reduced by transfection of miR-29s with the deletion vector, the reduction was smaller than that with wild type (Figure 4D).

Effects of silencing *LAMC2* and *ITGA6* on cell proliferation, migration and invasion in SAS. To investigate the functional role of *LAMC2* and *ITGA6*, we performed loss-of-function studies using si-*LAMC2* and si-*ITGA6* transfectants. First, we evaluated the knockdown efficiency of si-*LAMC2* and si-*ITGA6* treatments in SAS. Western blotting and qRT–PCR indicated that the two siRNAs effectively downregulated *ITGA6* and *LAMC2* expression in SAS (Supplementary Figure 3).

The XTT assays demonstrated that cell proliferation was significantly inhibited in si-LAMC2 transfectants in comparison

with the mock or si-control transfectant in SAS (percentage of the mock: 64.7 ± 2.2 , with P < 0.0167; Figure 5A). On the other hand, proliferation was not inhibited in si-*ITGA6* transfectants in comparison with the mock or si-control transfectants in SAS (Figure 5B).

The migration assays demonstrated that cell migration activity was significantly inhibited in both si-*LAMC2* and si-*ITGA6* transfectants in comparison with the mock or si-control transfectants in SAS (percentage of the mock: si-*LAMC2*, 14.3 ± 7.2 ; si-*ITGA6*, 15.8 ± 12.5 , with *P* < 0.0167 for both; Figure 5C and D).

Matrigel invasion assays demonstrated that cell invasion activity was significantly inhibited in both si-*LAMC2* and si-*ITGA6* transfectants in comparison with the mock or si-control transfectants in SAS (percentage of the mock: si-*LAMC2*, 23.4 ± 14.0; si-*ITGA6*, 19.6 ± 14.3, with P < 0.0167 for both; Figure 5E and F).

DISCUSSION

In the postgenome sequencing era, improved understanding of noncoding RNA is necessary for continued progress in cancer research. For elucidation of the molecular mechanisms underlying HNSCC, we have examined tumour-suppressive miRNAs, focussing on their regulated molecular targets and novel cancer pathways based on HNSCC expression signatures (Kikkawa *et al*, 2010; Nohata *et al*, 2011). Our previous miRNA expression signatures demonstrated that the *miR-29s* family (*miR-29a/b/c*) was reduced in several types of cancers (Kikkawa *et al*, 2010; Nohata *et al*, 2011). Decreased expression of *miR-29s* was

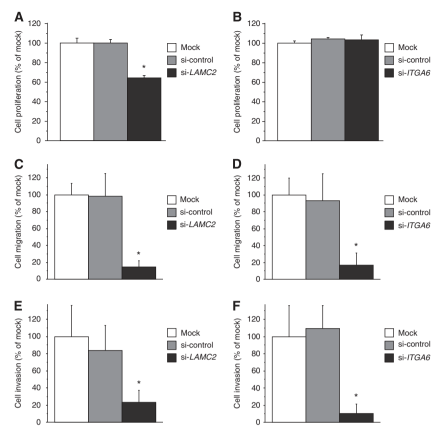


Figure 5. Effect of si-LAMC2 or si-ITGA6 transfection on cell proliferation, migration and invasion in SAS. Cell proliferation 72 h after transfection with 10 nm si-LAMC2 (**A**) or si-ITGA6 (**B**) in SAS was determined with the XTT assay. Cell migration activity 48 h after transfection with 10 nm si-LAMC2 (**C**) or si-ITGA6 (**D**) in SAS was determined with the migration assay. Cell invasion activity 48 h after transfection with 10 nm si-LAMC2 (**E**) or si-ITGA6 (**D**) in SAS was determined with the migration assay. Cell invasion activity 48 h after transfection with 10 nm si-LAMC2 (**E**) or si-ITGA6 (**F**) in SAS was determined with the Matrigel invasion assay. *P<0.0167.

described in other types of cancers such as cholangiocarcinoma, nasopharyngeal cancer, non-small cell lung cancer, cervical cancer, hepatocellular carcinoma, malignant peripheral nerve sheath tumour and mantle cell lymphoma (Fabbri et al, 2007; Mott et al, 2010; Xiong et al, 2010; Zhao et al, 2010; Li et al, 2011; Presneau et al, 2013), all of which are consistent with our results. In contrast, upregulation of miR-29s was reported in breast cancer, colon cancer and acute myeloid leukaemia (Gebeshuber et al, 2009; Han et al, 2010; Fu et al, 2012). Therefore, the molecular mechanisms by which miR-29s expression is deregulated vary according to cancer types. In this study, our data showed that all members of the miR-29s family were significantly reduced in cancer tissues. Moreover, restoration of miR-29s strongly inhibited cancer cell migration and invasion. Thus, the miR-29s family functions as a group of tumour suppressors, and they might contribute to metastasis in HNSCC.

To elucidate the silencing mechanisms of *miR-29s* in HNSCC cells, we performed preliminary analysis of 5-Aza-dC treatment as a demethylation agent on HNSCC cells. Our data demonstrated that *miR-29s* were not recovered by the treatment (data not shown), suggesting that silencing of *miR-29s* were not caused by methylation of CpG islands on their genomic regions. Three annotated genes (*CR1L, CD46* and *C1orf32*) are located around the *miR-29b-2/miR-29c* cluster region on human chromosome 1q32.2 (Supplementary Figure 1). Our microarray expression data of HNSCC (GEO accession number: GSE 36951) showed that expression changes of the three genes were not observed in cancer tissues. According to the UniGene database, expression levels of other genes shown in Supplementary Figure 1 are extremely low.

Although the molecular mechanism by which *miR-29s* is silenced in HNSCC is still unknown, some important studies have

examined the promoter region of *miR-29s* in the human genome. The human *miR-29b-1/miR-29a* promoter region contains two putative E-box sites (MYC-binding site), a Gli-binding site and four NF- κ B-binding sites. Increased expression of MYC silenced *miR-29b-1/miR-29a* expression in cholangiocarcinoma cells (Mott *et al*, 2010). Furthermore, NF- κ B signalling, which is known to be activated in inflammation-related cancers, directly repressed *miR-29b-1/miR-29a* promoter activity (Wang *et al*, 2008; Mott *et al*, 2010). Recent data showed that GATA3, a transcription factor, promotes differentiation, suppresses metastasis and alters the tumour microenvironment in breast cancer by inducing *miR-29b* expression (Chou *et al*, 2013). Thus, it will be necessary to identify the transcription factors contributing to HNSCC. Further detailed analysis is needed to understand the molecular mechanisms of *miR-29s* silencing in HNSCC cells.

Metastasis is the final stage of cancer progression and is the cause of most cancer deaths. The epithelial-to-mesenchymal transition (EMT) is considered to be an important step in cancer progression and metastasis, and TGF- β signalling contributes to EMT processes (Heldin *et al*, 2012). Recent data have suggested that TGF- β 1 inhibits the expression of *miR-29s* and promotes the expression of extracellular matrix (ECM) components (Maurer *et al*, 2010; Roderburg *et al*, 2011). These data suggest that *miR-29s* could play an important role in modulating TGF- β signalling during cancer cell migration and invasion. It will be of interest to more fully identify the molecular targets and pathways of *miR-29s* in HNSCC, particularly those contributing to cancer cell migration and invasion.

The miRNAs are unique in their ability to regulate many protein-coding genes. Bioinformatic predictions indicate that miRNAs regulate > 30% of protein-coding genes (Lewis *et al*,

2005). We have taken the position that the identification of novel cancer pathways and responsible genes regulated by tumoursuppressive *miR-29s* is an important first step in understanding HNSCC oncogenesis. Based on this view, we categorised *miR-29s* target genes into known pathways using the KEGG pathway database. These data have facilitated the understanding of tumour-suppressive miRNA-regulated molecular pathways in human cancer. We devised this method of analysis and found that tumour-suppressive miRNA could efficiently regulate cancer-associated pathways (Kinoshita *et al*, 2012; Nohata *et al*, 2013; Yoshino *et al*, 2013).

In this study, we focussed on the 'focal adhesion' pathway because restoration of miR-29s inhibited cancer cell migration and invasion in HNSCC cell lines. Cell adhesion is the binding of a cell to a surface, such as an ECM or another cell, and a fundamental determinant of a wide range of cell biology responses and signalling (Parsons *et al*, 2010). Recent evidences suggested that several miRNAs including miR-29s regulated multiple molecular components of the cell adhesion machinery and contributed to functional key regulators of adhesion-associated processes (Valastyan and Weinberg, 2011). Our hypothesis is that

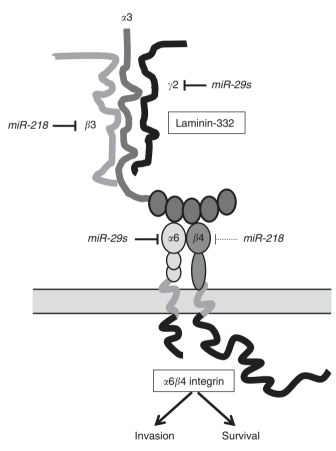


Figure 6. Illustration of tumour-suppressive *miR-29s* and *miR-218* regulation of laminin-332 and $\alpha 6\beta 4$ integrin. Regulation of laminin-332 and $\alpha 6\beta 4$ integrin by *miR-29s* and *miR-218* is shown (modified a Figure in Guess and Quaranta, 2009). Laminin-332, which consists of laminin $\alpha 3$, $\beta 3$ and $\gamma 2$ chains, is a major component of basement membranes, but is also an important autocrine ligand produced by cancer cells to promote tumourigenesis. Laminin-332– $\alpha 6\beta 4$ integrin interaction promotes tumour invasion and survival by activation of PI3K and RAC1. Both LAMC2 and ITGA6 were directly regulated by tumour-suppressive *miR-29s*, whereas LAMB3 was directly regulated by tumour-suppressive *miR-218* (our previous study). In addition, *ITGB4* has a putative *miR-218*-binding site in the 3'-UTR region.

downregulation of tumour-suppressive miRNAs in cancer cells leads to upregulation of oncogenes. Therefore, we combined the gene expression data of miR-29a transfectants and the GEO expression data of upregulated genes in HNSCC, generating 16 candidate target genes for miR-29s in focal adhesion pathways. We focussed on laminin $\gamma 2$ (LAMC2), a member of laminin-332, and $\alpha 6$ integrin (*ITGA6*) as responsible genes that contribute to cancer cell migration and invasion in HNSCC. Numerous clinical studies indicated that overexpression of LAMC2 was positively correlated with invasiveness and poor survival in several types of cancer (Guess and Quaranta, 2009), suggesting that LAMC2 functions as an oncogene and deeply contributes to cancer metastasis. Because of the small number of the HNSCC samples, no positive data were obtained between clinical significance of the HNSCC patients and the LAMC2 and ITGA6 expression levels in the present study. Further examination should be necessary to elucidate the expression levels of LAMC2 and ITGA6 and the clinicopathological parameters and patient survival of HNSCC.

Integrins are a large family of cell surface receptors composed of two subunits (α and β) that bind to ECM components. Most types of cells require integrin-mediated signal pathways for proliferation, migration, invasion and survival. To date, a total of 18 different α -subunits and 8 different β -subunits have been identified, accounting for at least 24 distinct integrin heterodimers (Gilcrease, 2007). Among these integrins, laminin-332 interacts with two major integrin receptors, $\alpha 3\beta 1$ and $\alpha 6\beta 4$, promoting the formation of focal adhesions and stable anchoring contacts (Marinkovich, 2007). The $\alpha 6\beta 4$ integrin is reportedly involved as a link between the cytoskeleton and ECM as well as in the activation of a variety of intercellular signalling processes in cooperation with growth factor receptors (Falcioni *et al*, 1997; Mariotti *et al*, 2001; Trusolino *et al*, 2001).

Of particular interest, laminin-332 and $\alpha 6\beta 4$ integrin interaction triggers a number of signalling cascades in cancer cells, promoting both cell migration and cancer cell survival (Marinkovich, 2007). Furthermore, our previous data demonstrated that tumoursuppressive miR-218, which was frequently reduced in HNSCC, directly regulated laminin-332, suggesting that miR-218 contributed to cancer cell migration and invasion through regulating focal adhesion pathways (Kinoshita et al, 2012). Thus, the available data suggest that tumour-suppressive miR-29s and miR-218 contribute to cancer cell migration and invasion through their regulation of focal adhesion pathways, especially targeting laminin-332 and $\alpha 6\beta 4$ integrin signalling (Figure 6). Evidence indicates that integrins crosstalk with receptor tyrosine kinases in an integrin type-dependent manner through a variety of specific mechanisms (Falcioni et al, 1997; Mariotti et al, 2001; Trusolino et al, 2001). Thus, the understanding of tumour-suppressive miRNAs (miR-29s and miR-218) and their regulation of laminin-integrin signalling should shed light on HNSCC metastasis as well as delineate more effective strategies for future therapeutic interventions for this disease.

CONCLUSIONS

Our data showed that all members of the *miR-29s* group were frequently downregulated in HNSCC. They function as tumour suppressors and normally inhibit cancer cell migration and invasion through their regulation of focal adhesion pathways, especially via laminin-332 and $\alpha 6\beta 4$ integrin. Elucidation of cancer pathways regulated by tumour-suppressive *miR-29s* should shed light on HNSCC metastasis as well as delineate more effective strategies for future therapeutic interventions for this disease.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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