Overexpression of the oncostatin-M receptor in cervical squamous cell carcinoma is associated with epithelial–mesenchymal transition and poor overall survival

Justyna A Kucia-Tran¹,6, Valtteri Tulkki¹,6, Stephen Smith¹, Cinzia G Scarpini¹, Katherine Hughes², Angela M Araujo¹, Ka Yin Matthew Yan¹, Jan Botthof¹, Eduardo Pérez-Gómez³, Miguel Quintanilla⁴, Kate Cuschieri⁵, Maria M Caffarel *,¹,7 and Nicholas Coleman *,¹

¹Department of Pathology, University of Cambridge, Cambridge CB2 1QP, UK; ²Department of Veterinary Medicine, University of Cambridge, Cambridge CB3 0ES, UK; ³Department of Biochemistry and Molecular Biology I, School of Biology, Complutense University, Madrid 28040, Spain; ⁴Department of Cancer Biology, Instituto de Investigaciones Biomédicas Alberto Sols, Consejo Superior de Investigaciones Científicas (CSIC), Universidad Autónoma de Madrid (UAM), Madrid 28029, Spain and ⁵Scottish Human Papilloma Virus Reference Laboratory, Specialist Virology Centre, Royal Infirmary of Edinburgh, Edinburgh EH16 4SA, UK

Background: Copy-number gain of the oncostatin-M receptor (OSMR) occurs frequently in cervical squamous cell carcinoma (SCC) and is associated with adverse clinical outcome. We previously showed that OSMR overexpression renders cervical SCC cells more sensitive to the major ligand oncostatin-M (OSM), which increases migration and invasion in vitro. We hypothesised that a major contribution to this phenotype would come from epithelial–mesenchymal transition (EMT).

Methods: We performed a comprehensive integrated study, involving in vitro cell line studies, in vivo animal models and numerous clinical samples from a variety of anatomical sites.

Results: In independent sets of cervical, head/neck and lung SCC tissues, OSMR expression levels correlated with multiple EMT-associated phenotypic markers and transcription factors. OSM treatment of OSMR overexpressing cervical SCC cells produced consistent EMT changes and increased tumour sphere formation in suspension culture. In a mouse model, OSMR overexpressing SCC cells treated with OSM showed significant increases in lung colonisation. The biological effects of exogenous OSM were mirrored by highly significant adverse overall survival in cervical SCCs with OSMR overexpression (N = 251).

Conclusions: OSM:OSMR interactions are able to induce EMT, increased cancer stem cell-like properties and enhanced lung colonisation in SCC cells. These changes are likely to contribute to the highly significant adverse outcome associated with OSMR overexpression in cervical SCCs.

There are ~300 000 deaths from cervical carcinoma worldwide each year (Forman et al, 2012). Most cases are squamous cell carcinomas (SCC), which develop from precursor squamous intraepithelial lesions (SILs; Groves and Coleman, 2015). Vaccination against high-risk human papillomavirus (HPV), the necessary cause of cervical SCC, is of no benefit to women who are already
persistently infected with the virus (Stanley et al., 2014). Improved understanding of cancer biology can be translated into new treatments for cervical SCC, where novel targeted therapies would be an important advance.

Early studies identified oncogenes of potential significance in cervical SCC, by screening for genes located in regions of frequent copy-number gain (CNG) that showed significant associations between expression levels and copy number (Ng et al., 2007). One of the most common regions of CNG in advanced cervical SCC is on chromosome 5p, where a candidate driver gene is the oncostatin-M receptor (OSMR), a cell surface cytokine receptor that is suitable for antibody-mediated therapeutic inhibition (Caffarel and Coleman, 2014). In our previous work, we observed gain of the genomic locus containing OSMR in 61% of 36 cervical SCC samples using array CGH and also found OSMR gain in 61% of 110 independent cervical SCCs using fluorescence in situ hybridisation (Ng et al., 2007). SCCs with OSMR CNG showed significantly higher OSMR expression than those without CNG (P < 0.001; Ng et al., 2007). OSMR overexpression (which we define as levels at least twofold greater than the mean of normal cervix samples) was seen in 50% of 18 non-selected cervical SCCs (Ng et al., 2007). Other groups independently showed chromosome 5p CNG in >60% of advanced cervical carcinomas (Narayan et al., 2007; Scotto et al., 2008), while OSMR was overexpressed at 3.1-fold greater levels in 30 cervical carcinomas compared with 20 cases of normal cervical squamous epithelium (Scotto et al., 2008). Importantly, we found that OSMR CNG was associated with significantly adverse clinical outcome in cervical SCC, with a relative risk of death vs cases without CNG of 3.6 (Ng et al., 2007).

In cervical SCC cells in vitro, OSMR overexpression conferred greater sensitivity to the major ligand oncostatin-M (OSM), which induced increased cell migration and invasion (without affecting cell proliferation), plus an ability to induce angiogenesis (Winder et al., 2011). Expression profiling of OSM-treated cell lines and cervical SCC tissue identified consistent patterns of gene induction following OSM:OSMR interactions, which led to functional analyses of OSM-induced promalignant effects in cervical SCC cells (Winder et al., 2011). We identified that one mechanism by which OSMR confers increased motility and invasiveness to cervical SCC cells is by inducing transglutaminase-2 (tissue transglutaminase), which, among other functions, interacts with cell surface integrin-α5β1 and fibronectin (Caffarel et al., 2013). In the present study, we hypothesised that a further mechanism mediating the promalignant effects of OSMR overexpression in cervical SCC cells is induction of epithelial–mesenchymal transition (EMT), a process linked with invasion and metastasis (Pollyak and Weinberg, 2009).

EMT is a dynamic and plastic mechanism that involves complex transcriptional and post-transcriptional cellular changes (Craene and Berx, 2013). These lead to downregulation of epithelial genes and upregulation of mesenchymal markers, orchestrated through EMT-associated transcription factors (EMT-TFs), including Snail, Slug (Snail2), Zeb1 and Zeb2 (Lamouille et al., 2014). EMT is characterised by changes in cell–cell contact, shape, motility and invasiveness and may also be associated with the acquisition of stem cell-like properties (Mani et al., 2008), including the ability to grow as cell spheres in suspension culture. In carcinomas, EMT and cancer stem cell (CSC)-like features are thought to arise through common mechanisms of induction, including those originating in the tumour microenvironment (TME; Korkaya et al., 2011; Jung et al., 2015). We previously identified that OSMR levels in SCCs correlated positively with those of fibronectin and integrin-α5 (Caffarel et al., 2013), both regarded as mesenchymal markers (Ocaña et al., 2012).

We investigated the functional associations between OSMR overexpression and EMT by examining cervical SCC cells in vitro and in preclinical metastasis models, and by analysing multiple clinical sample data sets. Our evidence further supports therapeutic targeting of OSMR in SCCs of the cervix and potentially other anatomical sites.

**MATERIALS AND METHODS**

**Tissue samples and gene expression profiles.** All clinical tissue sample sets studied are summarised in Supplementary Table S1.

We measured OSMR expression levels by qRT–PCR in: 13 cervical SCCs and 3 samples of normal ectocervical squamous epithelium (set 1; Scottish HPV Reference Laboratory, Edinburgh); 27 SCCs of the larynx and 9 samples of normal larynx (set 2; CNIO Tumor Bank Core Unit, Madrid, Spain); 18 skin SCCs and 7 samples of normal skin (set 3; CNIO Tumor Bank Core Unit, Madrid, Spain). All three tissue sample sets were studied with Local Research Ethics Committee approval. In addition, we reanalysed our previous microarray gene expression profiling of 32 samples of lesional epithelium microdissected from cervical high-grade SILs (HSILs) and low-grade SILs (LSILs) and 12 samples of normal ectocervical squamous epithelium (set 4; Gene Expression Omnibus Accession Number GSE27678; Ng et al., 2007; Winder et al., 2011; Caffarel et al., 2013).

We also reanalysed published gene expression profiles from: 33 cervical SCCs plus 21 normal cervix (GSE9750, set 5; Scotto et al., 2008); 35 oral SCCs plus 6 normal oral mucosa samples (GSE10121, set 6; Sticht et al., 2008); 31 tongue SCCs plus 26 normal tongue, including 20 matched sample pairs (GSE13601, set 7; Estilo et al., 2009); 13 vulvar SCCs plus 5 normal vulva (GSE38230, set 8; Micci et al., 2013); 15 skin SCCs, 14 actinic keratosis and 13 normal skin, including 13 matched sample pairs (GSE32628, set 9; Hameetman et al., 2013); 5 skin SCCs, 4 actinic keratosis and 6 normal skin (GSE2503, set 10; Nindl et al., 2006); 12 matched sample pairs of skin SCC and normal skin (set 11; Hudson et al., 2010).

**Analysis of next-generation sequencing data.** We analysed expression levels of OSMR and other EMT genes in the publically available RNA sequencing results of The Cancer Genome Atlas (TCGA). Computational analysis and statistical testing of the next-generation sequencing (NGS) data were conducted using the R statistical programming language. Filtered and log 2 normalised RNA expression data, along with all available clinical data, were downloaded from the GDAC firehose database (run: stddata__2015_06_01) for each gene of interest from the CESC (cervical cancer), HNSCC (head and neck squamous cell carcinoma) and LUSC (lung squamous cell carcinoma) collections (sets 12-14; Supplementary Table S1). Data downloads were performed using the FirebrowseR database access R package.

Correlation testing for associations between expressed genes was performed using the cor.test function in R, to calculate the coefficient of variation (r) and test for significant deviation from no correlation. A list of 16 EMT-related genes (Supplementary Table S2) was derived from published literature (Pollyak and Weinberg, 2009; Nieto, 2011; Craene and Berx, 2013; Lamouille et al., 2014). Plotting of the correlation data was performed using the ggplot2 R package. Survival analysis was performed using the survival R package. Kaplan–Meier estimated survival curves were constructed using the clinical data. Statistical testing of differences between survival curves used the G-rho family of tests, as implemented in the survdiff function of the survival package.

In addition, we analysed published NGS data from 24 cervical SCC samples, in which the carcinoma and tumour-infiltrating cells had been separated by flow sorting, followed by RNA sequencing of the two cell populations (set 15; Supplementary Table S1; Punt et al., 2015).

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Cells and cell culture. We used the representative OSMR overexpressing cervical SCC cell lines CaSki (HPV16-positive) and SW756 (HPV18-positive), in which OSMR was expressed at 3.4- and 4.5-fold greater levels than in a pool of four independent cultures of basal-type normal cervical squamous cells (Ng et al., 2007). There was no overexpression of OSM in either cell. The cells were obtained from the American Type Culture Collection (ATCC-LGC, Middlesex, UK) and verified by short tandem repeat profiling (Muralidhar et al., 2011). Cell culture and OSM treatment were performed as described (Winder et al., 2011). OSM (R&D Systems, Abingdon, UK) was added at 10 ng ml$^{-1}$, in accordance with previous studies examining long-term effects of OSM in tissue culture (Winder et al., 2011). OSM secretion into culture medium was measured by sandwich ELISA (human OSM DuoSet ELISA, R&D Systems), according to the manufacturer’s instructions.

Sphere forming assay. Cells in monolayer cultures were trypsinised, filtered through a 30-μm filter (Systex, Milton Keynes, UK) and resuspended in DMEM medium containing 1% methy cellulose (R&D systems), 20 ng ml$^{-1}$ EGF and 20 ng ml$^{-1}$ b-FGF (both PeproTech, London, UK). Cells were then added to non-adhesive 96-well plates (Greiner bio-one Cellstar, Monroe, NC, USA), at 1000 cells per well, and treated every three days with medium containing 10 ng ml$^{-1}$ OSM, or control medium. The total number of spheres in each well was counted after 10 days and numbers in OSM-treated wells compared with those of control wells. Spheres were then isolated by centrifugation and the RNA extracted.

Real time quantitative PCR and western blotting. Relative mRNA levels were measured by QuantiTect One-Step SYBR-Green qRT–PCR (Qiagen, Crawley, UK) as described (Caffarel et al., 2013), using the primers and conditions listed in Supplementary Table S3. Expression ratios were calculated using the comparative threshold cycle (Ct) method (Pfaffl, 2001) and normalised using three housekeeping genes; ACTB, TBP and HMBS (Vandesompele et al., 2002). Western blotting and densitometry were performed as described (Caffarel et al., 2013), using the primary antibodies listed in Supplementary Table S4. In keeping with our previous work (Muralidhar et al., 2007; Muralidhar et al., 2011; Winder et al., 2011), we defined OSMR overexpression in SCCs as levels twofold or more greater than the mean for corresponding normal cervical squamous epithelium samples. The latter represented primary cultures for cell lines and microdissected epithelium for tissues.

Generation of OSM overexpressing SW756 cells and bioluminescent SW756 cells. SW756 cells were stably transfected with 2 μg of pUNO1-hOSM expression construct (InvivoGen, Toulouse, France), using Lipofectamine LTX with PLUS reagent (Thermo Fisher Scientific, Loughborough, UK) followed by Blasticidin (Thermo Scientific, France), using Lipofectamine LTX with PLUS reagent (Thermo Scientific, France), using Lipofectamine LTX with PLUS reagent. SW756 cells were stably transfected with 2 μg of OSMR expression construct (InvivoGen, Toulouse, France), using Lipofectamine LTX with PLUS reagent (Thermo Scientific, France), using Lipofectamine LTX with PLUS reagent. SW756 cells were stably transfected with 2 μg of OSMR expression construct (InvivoGen, Toulouse, France), using Lipofectamine LTX with PLUS reagent.

Small-molecule inhibition and RNA interference. Cells were treated for 2 h with medium supplemented with TGI01348 (Axon Medchem BV, Groningen, The Netherlands), a small-molecule inhibitor of JAK2. This was used at 1 μM, which was the lowest concentration that showed inhibition of STAT3 phosphorylation on western blotting. Negative control cells were treated with equivalent volumes of DMSO vehicle (vol/vol) only.

STAT3 or OSMR were each depleted using four pooled siRNAs (ON-TARGETplus SMARTpool L-003544 or L-008050 respectively, both Dharmacon, Lafayette, CO, USA), compared with cells treated with non-targeting control (NTC) pooled siRNAs (siCONTROL non-targeting siRNA-1, Dharmacon). Individual siRNAs were used at 10 nM, producing 40 nM pools. Cells were transfected at 20–30% confluence using Lipofectamine RNAiMAX (Invitrogen, Paisley, UK), as described (Hanning et al., 2013a; Hanning et al., 2013b). OSM or PBS was added 24 h after transfection and cells treated for 48 h.

Tail vein injection and subcutaneous xenografts. For experimental metastasis experiments, bioluminescent SW756 cells (1 × 10$^6$ in 100 μl) were injected into the lateral tail vein of 5–6-week-old female NOD SCID (NOD.CB17–Prkd$c^{+/+}$/NcrCrl) mice (Charles River, Oxford, UK), using 29-gauge insulin needles. Growth of lung tumours was monitored non-invasively using optical whole-body imaging at weekly intervals. Mice were injected with d-luciferin intraperitoneally and imaged 15 min later using the IVIS Spectrum preclinical imaging system (Perkin Elmer, Waltham, MA, USA). Mice were treated three times a week with intraperitoneal 100 μl injections containing 1 μg OSM or PBS only. Animals were killed 11 weeks after tail vein injection and the lungs examined histologically. The tissue samples were fixed in 4% formaldehyde in PBS and embedded in paraffin. Five-micron sections were stained with H&E, observed with a DM LB light microscope (Leica, Wetzlar, Germany) and photographed using a DC500 digital camera (Leica).

To establish subcutaneous xenografts, 5–6-week old female NOD SCID (NOD.CB17–Prkd$c^{+/+}$/NcrCrl) mice were implanted subcutaneously in the flank with 5 × 10$^6$ bioluminescent SW756 cells. Mice were treated three times a week with peri-tumoural injections of 100 μl containing 1 μg OSM or PBS only. All mice were palpated three times a week until tumours were detected and thereafter tumours were measured with callipers. Tumour volumes were calculated as (4 × width/2) × (height/2) × (length/2). Animals were killed nine weeks after cell transplantation and the subcutaneous tumours collected. Metastases in visceral organs and brain were evaluated by quantification of bioluminescent signal and by histopathology. To quantify skeletal metastases, visceral organs were removed and the mice assessed by optical imaging.

All mice were maintained in conventional cages within a specific pathogen-free animal facility. They were sacrificed by neck dislocation. All animals were treated in strict accordance with the local ethical committee (University of Cambridge Licence Review Committee) and the UK Home Office guidelines.

Statistics. For comparisons between groups we used Student’s T test and ANOVA, with post-hoc analysis by the Student–Newman–Keuls’ test. Unless otherwise stated, results are expressed as mean ± s.e.m. Timecourses for the development of lung metastases in vivo were compared using linear mixed effects modelling, implemented in R via the lme4 package.

RESULTS

OSMR expression associates with EMT markers in cervical and other SCCs. We used TCGA gene expression data for multiple SCC sample sets to test for correlations between OSMR levels and those of 16 genes associated with EMT (Supplementary Table S2). In 251 cervical SCCs (sample set 12), we observed strong and significant ($P<0.05$) correlations between OSMR and eight of the 16 genes, namely fibronectin (FN1), FOX22, HMG2A, integrin–α5 (ITGAV3), MMP10, Snail (SNAI1), Slug (SNAI2) and ZEB1 (Figure 1). Importantly, similar correlations were also seen in SCCs from other anatomical sites. OSMR levels correlated with 14 of the 16 EMT genes in 514 head/neck SCCs (sample set 13) and with all 16 genes in 492 lung SCCs (sample set 14) (Supplementary Table S2).
OSM induces EMT and CSC-like features in OSMR-overexpressing cervical SCC cells in vitro. As SCC tissue samples include both epithelial and stromal components, we used cervical SCC cells in vitro (Ng et al, 2007) to investigate the direct associations between OSMR overexpression and EMT. In both CaSki and SW756, treatment with exogenous OSM at 10 ng ml$^{-1}$ for 72 h induced a more mesenchymal morphology, with cell elongation, reduced cohesion and a more refractile appearance on bright-field microscopy (Figure 2A). Consistent with these changes, the OSM-treated cells showed reduced mRNA and protein expression of the epithelial marker E-cadherin (Figure 2B), together with increased expression of fibronectin (Figure 2C), the EMT-TFs Snail and ZEB2 (Figure 2D), and the matrix metalloproteinases MMP9 and MMP10 (Figure 2E). The mesenchymal marker vimentin was also induced in CaSki cells (Supplementary Figure S1). The effects of OSM could be reduced or abrogated following depletion of OSMR using pooled siRNAs (Supplementary Figure S1). SW756 cells that constitutively overexpressed OSM following stable transfection also showed a more mesenchymal morphology, identical to that of cells treated with exogenous OSM (Supplementary Figure S2A), and also showed increased levels of fibronectin, Snail, ZEB2, MMP9 and MMP10, compared with control-transfected cells (Supplementary Figure S2B).

In addition, treatment with exogenous OSM increased the formation of tumour spheres by SW756 cells grown in suspension culture for 10 days (Figure 3A), consistent with increased CSC-like properties (Wang et al, 2013). When compared with untreated control spheres, the OSM-treated spheres showed increased levels of Snail and ZEB2, as well as the pluripotency marker SOX2 and OSMR itself (Figure 3B). The untreated spheres showed increased levels of Snail and OSMR compared with SW756 cells in monolayer culture. Untreated CaSki cells formed spheres very efficiently and there were no additional effects of OSM treatment.

EMT induction requires JAK2/STAT3 signalling. As OSM:OSMR interactions are known to induce STAT3 (Ng et al, 2007), we tested the importance of JAK/STAT3 signalling in the induction of EMT-related genes. To investigate the role of JAK2, cells were treated with OSM for 15 min, followed by assessment of STAT3 phosphorylation by western blotting after 30 min and measurement of Snail transcript levels after 24 h. In this experiment, cells pretreated with the small-molecule JAK2 inhibitor TG101348 for 2 h prior to the addition of OSM showed reduced STAT3 phosphorylation and Snail induction (Figure 4A). In addition, STAT3 depletion using pooled siRNAs (Supplementary Figure S3) inhibited the induction of fibronectin and Snail proteins following OSM treatment for 48 h (both Figure 4B) and also inhibited expression of vimentin, ZEB2, MMP9 and MMP10 in one or both of the OSMR overexpressing cell lines (Figure 4C).
Figure 2. OSM induces EMT in OSMR overexpressing cervical SCC cells. (A) Effects of OSM treatment (72 h) on the morphology of OSMR overexpressing cells in monolayer culture. (B–E) Levels of EMT-associated markers in OSMR overexpressing cervical SCC cells treated with OSM for the indicated time points, compared with control treated cells (C). Graphs show transcript levels and blots show protein levels. B, CDH1 (E-cadherin), C, fibronectin, D, EMT-TFs Snail and ZEB2, E, MMP9 and 10. Transcript quantification values are from three independent experiments, whereas the western blots show the data from one representative experiment. The protein loading control was β-actin (lower row of the blots). As the blots in C and D were done on the same membrane, the same loading controls are shown for each. *P<0.05, **P<0.01, ***P<0.001.
OSM increases metastasis of OSMR-overexpressing SW756 cervical SCC cells in vivo. Mice injected intravenously with luciferase-expressing SW756 cells and treated three times per week with intraperitoneal OSM injections showed significantly increased lung metastasis, compared with PBS-treated control animals (Figure 5A and B). There were no histological differences between the lung metastases seen in the two experimental groups (n = 5 mice per group; Figure 5C).

Subcutaneous xenografts of SW756 cells treated three times per week with peri-tumoural OSM injections showed no difference in growth rates compared with PBS-treated control xenografts (n = 4 per group; Figure 5D). There were no macroscopic or histological differences between the primary xenografts from the two experimental groups. However, the OSM-treated group showed more skeletal metastases (n = 2 per group; Figure 5D). There was no evidence of metastasis to other sites, including liver and lung.

OSMR is overexpressed in multiple SCCs and associated with worse overall survival. In view of the effects of OSM:OSMR interactions on metastasis in vivo, we investigated the frequency of OSMR overexpression in SCCs, as well as the associations with overall survival. When compared with equivalent normal tissues, OSMR was overexpressed in SCCs from numerous sites, including cervix, head/neck, vulva and skin (sets 1–3, 5–11) (Figure 6A–C; Supplementary Figure S4). In cells flow-sorted from cervical SCCs (set 15; Punt et al., 2015), OSMR was overexpressed by the carcinoma cell fraction only, with very little expression in the CD45 + tumour-infiltrating immune cells (Figure 6D). In contrast, OSM was expressed strongly by the immune cells, with minimal expression in the carcinoma cells (Figure 6D).

Lower-level increases in OSMR expression were also seen in premalignant squamous lesions. In the cervix, there were gradual increases in expression levels from normal cervix, through LSILs to HSILs (set 4; Supplementary Figure S4B). OSMR overexpression (levels at least twofold greater than the mean of the 12 normal cervix samples) was seen in 3/21 HSILs, 0/11 LSILs and 0/12 normal cervix. In skin tissue, OSMR levels in actinic keratosis, a potentially premalignant condition (Fu and Cockerell, 2003), were intermediate between those of normal skin and cutaneous SCC (sets 9 and 10; Supplementary Figure S4F and S4G).

We next examined the associations between OSMR expression levels and clinical outcome, using cervical SCCs from TCGA. In this analysis, we were unable to define OSMR overexpression by reference to normal cervix samples, which were not included in the data set. Accordingly, we divided the SCCs into two groups, those with OSMR expression levels above the median for the data set (n = 126) and those with expression levels at or below the median (n = 125). Importantly, the SCCs with high OSMR expression showed a highly significant adverse overall survival, compared with those with low expression (P = 0.006; Figure 6E).
This finding was confirmed in a small independent set of 13 cervical SCCs (set 1). Of these, seven (54%) overexpressed OSMR, defined as levels at least twofold greater than the mean of the three normal cervix samples in the data set. These seven SCCs showed significantly worse overall survival than the six SCCs without OSMR overexpression (Figure 6A). None of the three normal cervix samples in this data set showed OSMR overexpression.

**DISCUSSION**

This is the first study to show that OSMR overexpression is associated with EMT features and adverse clinical outcome in SCC samples and that OSM can enhance EMT, CSC-like properties and metastasis in SCC cells. We identified that OSMR is frequently overexpressed in SCCs from multiple anatomical sites and provides the malignant cells with increased responsiveness to the major ligand OSM. The frequency of OSMR overexpression observed in the cervical SCCs of sample set 1 (54%) was similar to our previous observations in an independent data set (50%) (Ng et al, 2007). In total, OSMR overexpression was present in 52% of the cervical SCCs we examined (16/31), consistent with the observed frequency of OSMR CNG (61%) and our evidence of significant associations between gene copy number and expression (Ng et al, 2007; Winder et al, 2011).

In cervical SCC samples, OSMR was almost exclusively expressed by the carcinoma cells, while OSM was almost exclusively expressed by the infiltrating immune cells. The latter observation is consistent with evidence that OSM may be produced by a wide range of cells in the TME, including macrophages, dendritic cells and T-lymphocytes (Tanaka and Miyajima, 2003). We previously demonstrated that OSM stimulation of OSMR overexpressing cervical SCC cells leads to increased invasion and migration in vitro (Winder et al, 2011). We now show that such changes are accompanied by increased metastatic ability in vivo and that EMT is an important association of these malignant features.

Our present and previous (Caffarel et al, 2013) in vitro data demonstrate that exogenous OSM rapidly induces a consistent pattern of EMT-like changes in OSMR overexpressing cervical SCC cells, with downregulation of E-cadherin and upregulation of several mesenchymal markers. The latter include MMP9 and MMP10, which are key mediators of ECM destruction during carcinoma cell invasion (Nistico et al, 2012). All seven of the genes that we tested and showed to be upregulated in the SCC cells (fibronectin, MMP9, MMP10, Snail, vimentin, Zeb2 and integrin-α5 (Caffarel et al, 2013)), also showed significant positive associations with OSMR levels in one or more of the TCGA SCC tissue data sets (Figure 1; Supplementary Table S2). We concluded that the correlations between OSMR and multiple mesenchymal markers in clinical samples are likely to include direct effects on the carcinoma cells. However, contributions to these observations from cells of the TME cannot be excluded.

The effects of OSM treatment of OSMR overexpressing SCC cells also included increased CSC-like features, as measured using suspension cultures. The spheres formed by OSM-treated cells showed increased expression of mesenchymal genes, as well as the pluripotency marker Sox2, similar to observations in breast adenocarcinoma cells (West et al, 2014). Interestingly, OSMR was overexpressed in untreated cell spheres, compared with cells in monolayer culture, and was further induced by OSM. Altogether, these data indicate that OSMR is directly linked with CSC-like properties in cervical SCC cells. As EMT is a dynamic, reversible process (Thiery et al, 2009), reversion of OSM-exposed SCC cells to a more stem-like state would support clonal expansion and the development of tumour heterogeneity, within both primary lesions
and metastatic deposits. It might also be expected that high levels of OSMR and/or OSM in SCCs would confer resistance to chemotherapy and radiotherapy (Nieto, 2011). Indeed, this effect may have contributed to the association between OSMR levels and adverse clinical outcome that we observed in multiple cervical SCC data sets, including TCGA samples.

Figure 5. OSM increases lung colonisation by OSMR overexpressing SW756 cervical SCC cells. (A) Time course quantification of the thoracic bioluminescent signal from mice following tail vein injection of SW756 cells and intraperitoneal injection of OSM or PBS (n=5 per group). (B) Luminescence images from the end point of the experiment shown in A. (C) Representative images of lung tissue from the experiment shown in A. Images a–c are from PBS-treated animals, while images d and e are from OSM-treated animals. Image c is a high-magnification view of a region of image b. (D) The line graph (upper row, left) shows growth of primary subcutaneous xenografts of SW756 cells treated with peri-tumoral injections of OSM or PBS (n=4 per group). The images (lower row) were generated from bioluminescence analysis of mice at the end of the experiment, after removal of the xenografts and visceral organs. The images were overlapped with cartoon outlines to indicate the approximate locations of the skeletal luminescence signals. However, they are not precisely localised, due to variations in the orientation of the mice during optical imaging. The dot plot (upper row, right) shows the relatively greater levels of skeletal luminescence in mice bearing xenografts treated with OSM versus those treated with PBS (n=2 per group).
The enhanced responsiveness to OSM in OSMR overexpressing cervical SCC cells is associated with induction of multiple signalling pathways, including phosphorylation of STAT3 (Ng et al., 2007). Our present data indicate that STAT3 is important in inducing mesenchymal markers and the major EMT-TF Snail. One mediator of STAT3 phosphorylation is JAK2, as indicated by specific small-molecule inhibition. It is likely that other signalling pathways induced by OSM in OSMR overexpressing SCC cells are also of functional significance. In breast adenocarcinoma, MAPK signalling is associated with downregulation of oestrogen receptor (an epithelial marker; West et al., 2012) and PI3K signalling with increased CSC-like properties (West et al., 2014). Detailed dissection of the roles of signalling pathways in the multiple promalignant effects of OSM in cervical SCC cells is now required.

Figure 6. OSMR is overexpressed in SCCs from multiple sites and is associated with adverse overall survival in cervical SCC. (A–C) Levels of OSMR expression in SCC versus normal tissue from cervix (A), larynx (B) and skin (C). The line graphs in A show overall survival in the 13 cervical SCCs in sample set 1, according to OSMR expression levels. (D) Levels of OSMR and OSM in flow-sorted carcinoma and CD45+ infiltrating immune cells from cervical SCCs (sample set 15). (E) Kaplan–Meier curves showing overall survival for cervical SCCs with high vs low expression of OSMR, as determined from TCGA data (sample set 12).
There are likely to be multiple effectors of the transcriptional changes characteristic of EMT. Our current data, from in vitro experiments and clinical samples, indicate a role for the EMT-TFs Snail, Slug, ZEB1 and ZEB2. It will now be important to investigate the mechanisms responsible for the induction of these EMT-TFs, including similarities and differences with other cell types. For example, the OSMR target transglutaminase-2 (Caffarel et al, 2013) has been linked to EMT in ovarian malignancy (Shao et al, 2009) and may also have a role in SCCs. In breast adenocarcinoma cells OSM induces Lin28, which represses Let7 microRNAs and so upregulates protein-coding gene targets including HMGAA2 (Guo et al, 2013). The latter is a non-histone chromatin protein that acts as an architectural transcription factor and is known to bind the SNAI1 promoter (Thuault et al, 2008). Whether similar mechanisms apply in tumours of squamous histogenesis remains to be determined, particularly given the lineage specificity of microRNA profiles (Lu et al, 2005). However, we note the significant correlations between levels of OSMR and HMGA2 across the TCGA SCCs (Figure 1 and Supplementary Table S2). There may also be differences in mechanisms of EMT induction in SCCs at different sites. For example, OSMR levels correlated with some EMT markers (e.g., Twist2 and MMP2) in head/neck and lung SCCs but not cervical SCCs, while the high-risk HPV oncopgenes E6 and E7 that drive cervical and some head/neck SCCs have been linked with EMT induction (Qureshi et al, 2015).

Our in vivo data indicate that the metastatic capacity of OSMR overexpressing cervical SCC cells increases following exposure to OSM, as assessed by a lung colonisation assay. This demonstrates an important role for OSMR early in the metastatic process, for example via increased extravasation and infiltration of secondary sites (Nguyen et al, 2009). We observed no histological differences between the lung metastases at the end of the experiment, consistent with the reversibility of EMT, via the reciprocal process of mesenchymal–epithelial transition (Brablzet, 2012). In mice bearing cervical SCC xenografts, the increased skeletal metastases may have been due to several consequences of OSM:OSMR interactions, including enhanced cell motility/detachment, invasion, blood vessel intravasation and extravasation/colonization (Nguyen et al, 2009; Caffarel and Coleman, 2014). Our observations support the clinical evidence that OSMR overexpression in cervical SCC is associated with significantly worse overall survival, as well as with gene expression profiles indicating EMT. These data resonate with those in breast adenocarcinoma, where high OSMR expression correlates with shorter overall survival and recurrence free survival, together with features of EMT (West et al, 2014). Recently, evidence for an association between OSMR overexpression and adverse outcome has also been identified in a non-epithelial malignancy, glioblastoma multiforme (GBM), again with evidence of an association with EMT (Natesh et al, 2015).

Our data have indicated an important mechanism by which OSMR overexpression provides a selective advantage in cervical SCCs. Cells capable of greater migration/invasion are more likely to access conditions of improved oxygenation and nutrient availability and would therefore be provided with a competitive advantage in mixed cell populations. Induction of EMT/CSC-like properties increases metastasis and therapeutic resistance (Poljak and Weinberg, 2009), both of which are likely to contribute to the adverse clinical outcome associated with OSMR overexpression in cervical SCC. Together, our findings further strengthen the case for inhibiting OSM:OSMR interactions as a targeted anti-cancer therapy (Caffarel and Coleman, 2014). The observation that OSMR is commonly overexpressed in a range of SCCs, as well as other carcinomas and GBMs, indicate that effective targeting of this promalignant pathway may produce clinical benefits in a wide range of common cancers.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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