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AFM nanoindentation detect elastic modulus of tongue squamous carcinoma cells with different metastatic potential

**Running title:** AFM nanoindentation detect elastic modulus of TSCC

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Abstract

Although significant advances have been made in the molecular mechanisms that influence tongue squamous cell carcinoma (TSCC) metastasis, less is known about the association between the biomechanical behavior of the TSCC cells and their metastasis. To this goal, atomic force microscope (AFM) nanoindentation via the rate-jump method was used to detect the elastic modulus of TSCC cells from patients and cell lines with different metastatic potential. We found that TSCC cells with higher metastatic potential showed a reduction in elastic modulus as compared to TSCC cells with lower metastatic potential; moreover, the decrease in elastic modulus was accompanied by changes in epithelial–mesenchymal transition (EMT) and cytoskeleton (F-actin and β-tubulin), a smaller nucleus size, and a large nucleus/cytoplasm ratio. The present findings demonstrate a close relationship between cellular elastic modulus and metastasis of TSCC. Also, the detection of elastic modulus by AFM nanoindentation via the rate-jump method can potentially be used as a method to grade the metastatic potential of TSCC.

Keywords

Tongue squamous cell carcinoma; elastic modulus; metastasis; atomic force microscopy; nanoindentation
Background

Tongue squamous cell carcinoma (TSCC) is significantly more aggressive than other forms of oral squamous cell carcinomas (OSCC), and a propensity for rapid local lymph node (LN) metastasis. Improvement in patient survival requires a better understanding of tumor metastasis, which would allow aggressive tumors to be detected earlier in the disease process and targeted therapeutic interventions to be developed. Our previous studies revealed that the metastasis of TSCC may be related to many deregulated genes, such as SOD2, miR-138 and miR-222/221 [1-8]. In these studies we found that reductions in miR-138 are correlated with an enhanced metastatic potential and induce EMT (epithelial–mesenchymal transition) in TSCC cells. Similar to our studies, most of the current research regarding cancer metastasis has focused on biological capabilities such as molecular genetics and gene signaling, whereas the mechanical nature of the invasion process of the cancer cells has been ignored so far [9]. Indeed, the mechanical properties of living cells are strongly linked to the molecular alterations in the progression of cancer. Cumulative studies have revealed that the mechanical properties of individual cells (such as elastic and viscous) are important in the cell growth, motility and metastasis of cancer [10]. Malignant cancer cells are either less elastic (softer) or less viscous (less resistant to flow) than their normal counterparts, and metastatic cancer cells exhibit an even lower resistance to deformation than nonmetastatic cancer cells [11-13]. However, very few studies have focused on the effect of the elasticity of TSCC cells on their metastasis.

On the other hand, advances in experimental nano-biomechanics during the past two decades have enabled direct, real-time mechanical probing and the manipulation of single cells and molecules [10]. Commonly used experimental techniques include nanoindentation [14], micropipette aspiration [15] and microfluidic techniques [16]. Among these methods, nanoindentation is the most popular because quantitative measurement of the mechanical properties can be made directly without the need of carrying out time consuming post-experimental analyses. When nanoindentation is carried out on atomic force microscopes (AFM), the Hertzian model is commonly used to analyze the data to generate an elastic modulus of the sample [14, 17]. However, since the Hertzian model assumes the sample to be purely linear elastic, while the cytoplasm-membrane-cytoskeleton structure of a cell
should not behave in a purely linear elastic manner, the Hertzian model is almost definitely a wrong
description of the cell’s nanoindentation response. In fact in such analyses of experimental data by the
Hertzian model, the resultant elastic modulus estimates are often found to increase with the loading
rate [18, 19]. To solve this problem, our group has developed a rate-jump protocol for AFM
nanoindentation which can produce much more reliable measurement of the elastic modulus of soft
samples than the Hertzian model. This protocol has been found to yield elastic modulus that is
independent of the rate-jump magnitude from soft samples [18-20].

It is well known that the mechanical properties of human cells are largely governed by their
cytoskeleton [10]. During the descent of a cell from a mature, differentiated state to a highly mitotic
and motile cancerous state, the cytoskeleton regresses from an ordered and fairly stiff structure to a
more irregular and compliant state [21,22]. Furthermore, some other factors, such as EMT [23], the
overall cell size, the nucleus size [24], and chromatin organization [25], are all found to affect the
cell’s mechanical behavior. However, exactly how these cellular structural factors influence the
mechanical properties of cell remains quite ambiguous.

To investigate the relationship between the elastic modulus and metastatic potential of TSCC, the
everastic modulus of cancer cells from cell lines with different metastatic potentials and primary cell
samples from TSCC patients were examined using AFM nanoindentation via the above-mentioned
rate-jump method [18, 19]. Moreover, various cellular structural factors such as the cytoskeleton
structure, EMT, nucleus size, etc. were also investigated with various imagining techniques, so as to
explore the structure-mechanical property relationship of different types of tongue cancer cells with
different metastatic potential.
Methods

Primary cell culture

 Archived tissue samples for primary cell cultures from four TSCC cases were used in the present study. Clinical characterizations are summarized in Table S1. All of the patients underwent curative surgery and did not receive any adjuvant therapy prior to surgery, and all of the experiments with primary samples were performed within a few days after surgery. The present study was approved by the Ethical Committee of the First Affiliated Hospital, Sun Yat-Sen University. Primary cell cultures were maintained as previously described [22]. Briefly, TSCC tissues were disinfected with Betaisodona (Mundipharma, Limburg, Germany), rinsed twice in PBS, minced and placed in 2.5 mg/ml Dispase II (Roche, Mannheim, Germany) in DMEM for 18 to 24 h at 4°C. Subsequently, tissues were incubated in 0.25% trypsin/EDTA for 5 min. Trypsin activity was stopped with DMEM containing 10% fetal bovine serum (FBS). The suspensions were centrifuged, resuspended in DMEM/F12 (containing 10% FBS, 1,000 units/ml penicillin and 500 μg/ml streptomycin), and cultured at 37°C with 5% CO₂.

Cell culture

 UM1 and UM2 are paired TSCC cell lines with different metastatic potentials that were previously established from a single patient [26]. Another two paired cell lines with different metastatic potentials (UM1/Control and UM1/miR-138, 1386Ln/Control and 1386Ln/miR-138) were also used in this study [6]. These cell lines were stable transfected cells with control mimic or miR-138, respectively. 1386Ln is a cell line generated from lymph node metastatic disease from hypopharynx squamous cell carcinoma [6]. All the cells were maintained in DMEM/F12 containing 10% FBS, 1,000 units/ml of penicillin and 500 μg/ml of streptomycin in a 37°C incubator with 5% CO₂.

Wound healing assay

 Wound healing assay experiments were performed as previously described [27]. Images were taken at time points 0 h and 24 h post-wounding using a Nikon Diaphot TMD inverted microscope (4×). The relative distance traveled by the leading edge from 0 h to 24 h was assessed using Photoshop 7.0 software (n = 6).

Transwell invasion assay
Transwell invasion assay experiments were performed as previously described [27]. Briefly, Biocoat Matrigel invasion chamber inserts (BD Biosciences, NJ, USA) were equilibrated for 2 h at 37 °C in serum-free medium. Cells were seeded in serum-free medium in the upper chamber and allowed to invade through the Matrigel to the lower chamber for 24 h. Cells on the bottom surface of the filter were rinsed and permeabilized with 1% Triton X-100 in PBS for 20 min, stained with DAPI in the dark and visualized under a fluorescent microscope. Three random fields were captured at a magnification of 10× (n = 3). The number of cells on the bottom surface was compared between groups.

**Western Blot Analysis**

Western blot analysis was performed as previously described [28] using antibodies specific to E-cadherin, vimentin (Cell Signaling Technology, Beverly, MA, USA) and GAPDH (Sigma-Aldrich, MO, USA).

**Immunohistochemical analysis**

Immunohistochemistry was performed according to previous described [29] to detect the EMT markers (E-cadherin and vimentin) using a commercially available kit (Invitrogen, Carlsbad, CA). The relative intensities of the completed immunohistochemical reactions were evaluated by 3 independently trained observers who were blinded to the clinical data. Image-Pro Plus v6.0 software (Media Cybernetics, USA) was used to score the relative intensities.

**Scanning electron microscopy**

SEM was performed to investigate alterations in the cellular morphology. Briefly, cells were plated onto 14 mm diameter coverslips and placed in a 24-well plate for 24 h. Cells were then washed with PBS buffer and fixed with 2.5% glutaraldehyde in 0.1 M cacodylate buffer for 2 h at 4°C. After washing with cacodylate buffer with 0.1 M sucrose, cells were dehydrated for 5 min in an increasing series of ethanol solutions (30%, 50%, 75%, 90% and 100%). The samples were dried in a Critical Point Dryer and coated with gold prior to SEM scanning (Hitachi S4800 FEG SEM, Japan).

**Nucleus size and nucleus/cytoplasm (N/C) ratio measurement**

The nucleus size and N/C ratio measurements were performed using immunofluorescence imaging
Briefly, cells were fixed with 2.5% glutaraldehyde solution and permeabilized with 0.1% Triton X-100 in PBS. After a PBS wash, the cells were stained with anti-β-tubulin-Cy3 (Sigma-Aldrich, MO, USA) for cytoplasmic observation and Hoechst 33342 (1 μg/mL in PBS, Sigma-Aldrich, MO, USA) for nuclear observation. The cells were then washed with PBS and observed under a fluorescent microscope. The emission wavelengths for the Hoechst 33342 and Cy3 β-tubulin are 330-380 nm and 510-560 nm, respectively. The ImageJ (v. 1.45s) software by the National Institute of Health (NIH) was used to measure the areas of the cells and their nuclei, and the N/C ratio data were calculated with the following equation [31],

$$\frac{\text{Area}_{\text{nucleus}}}{\text{Area}_{\text{cell}} - \text{Area}_{\text{nucleus}}}$$ (1)

**Immunofluorescence analysis**

Cytoskeletal organization (F-actin and β-tubulin) was performed using the direct immunofluorescence method [30]. The filamentous actin (F-actin) was stained with a phalloidin-fluorescein isothiocyanate (phalloidin-FITC) conjugate solution (Sigma-Aldrich, MO, USA), and the β-tubulin was stained with anti-β-tubulin-Cy3. The slides were examined using a fluorescence microscope (Leica, Germany). Representative images of the cells were captured at a magnification of 400× and in the same exposure times of 200 milliseconds. The emission wavelength for the FITC-phalloidin and Cy3 β-tubulin are 450-490 nm and 510-560 nm, respectively. The mean intensity of F-actin and β-tubulin inside each entire cell was measured using ImageJ. In this method, the image was first converted into 8-bit format, and was then duplicated and used to determine the corresponding measured regions. A threshold function was applied to highlight the structure in the copy image and the resultant image was then converted into a binary image using the “binary” function of ImageJ. This image was then converted back to the original 8-bit format, and the mean intensity of the filaments was obtained by the function “analyze particles” in the software. In each group, the mean intensity of the filaments was obtained from 10 cells, and the corresponding mean intensities of different groups were compared, and in the Results section below, a typical image from each group is shown.
Atomic force microscopy nanoindentation

Nanoindentation experiments were performed in an AFM (JPK NanoWizard II, Germany) to measure the cellular elastic modulus as previously described [19, 20]. A silicon nitride cantilever-tip (CSG01, supplied by NT-MDT Company, Russia) was used, and the end of the tip was cut off using focused ion-beam (FIB) milling (FEI Quanta 200 3D FIB/SEM) to form a flat-ended cylindrical tip with diameter 1.01μm as shown in our previous study [20]. The choice of the flat-ended tip geometry with a cylindrical shape here ensures that the tip-cell contact area is a constant during the nanoindentation test, and this greatly simplifies the analysis [32]. The cantilever sensitivity was calibrated by indenting on the glass substrate in the presence of DMEM cell culture medium, and the spring constant of the tip was calibrated to be 0.037 N/m using the thermal fluctuation method prior to AFM nanoindentation. The cells were cultured on glass coverslips and placed on a Biocell™ holder (JPK, Germany), which ensured a constant 37°C environment for the cells during the entire AFM nanoindentation assay. AFM nanoindentation was performed by controlling the movement of the clamping base of the AFM cantilever-tip via a PZT scanner while the sample sat on a stationary base, and the deflection of the AFM cantilever at the tip’s end was recorded by a position-sensitive photo-diode detector.

The indentation schedule employed in this study is shown in Figure 1A and 1B. Each cell was mechanically probed by AFM at its most bulging part, where the nucleus was likely to be, as shown in Figure S1. The cell is not homogeneous, and therefore the values of elastic modulus may change significantly at different positions of the cell. In this case, indenting on the most bulging part of cell with the help of microscopic observation should largely reduce measurement errors [33, 34]. Moreover, indenting on this location will ensure that the membrane-cytoskeleton-nucleus response of the cell to be measured which is more pertinent to metastasis through narrow openings. The indentation depth was chosen to be at least 1 μm in order to better simulate deformations which occur physiologically as described by Xu et al [35]. Instead of running the test at a constant displacement rate as in the Hertzian method, in the rate jump method, a step change in the displacement rate is imposed, which results in a step change in the load rate. Theoretical considerations as previous described have shown that the relationship between these two step changes yields an elastic modulus
which is intrinsic of any power-law viscoelastic material [36]. As is shown in our previous study, the 
Hertzian method gives an apparent elastic modulus which increases with the loading rate, but the 
rate-jump method gives elastic modulus values which are independent of the rate-jump magnitude 
[19]. In the present study, each cell was just indented once, and 30 cells were randomly selected to be 
indented in each sample group for further statistical analysis.

In the rate-jump protocol, the elastic modulus of the sample is given by [19]

\[ \Delta \dot{P} = 2a \left( \frac{E}{1-v^2} \right) \Delta \dot{h} \]  

(2)

where \( h \) and \( P \) are the indentation depth and indentation force, and \( a, E \) and \( v \) are the radius of the 
cylindrical tip’s end (0.505 micron in the present study), the sample’s intrinsic Young’s modulus and 
Poisson’s ratio, respectively. In Eq. (2), \( \Delta \dot{P} \) and \( \Delta \dot{h} \) are jumps in the rates of \( P \) and \( h \) across the 
unloading point, and in the AFM used, the indentation force \( P \) is evaluated as  \( P = k(\delta - h) \) where \( k \) is 
the spring constant of the AFM cantilever, \( \delta \) is the programmed movement of the clamp base of the 
cantilever, and \( h \) is the deflection of the AFM tip as detected by the photo-diode detector (see ref. [19] 
for details). As explained in detail elsewhere [18, 20, 36], the rate-jump method involving eqn. (2) 
assumes the tested sample to be viscoelastic, and the elastic modulus \( E/(1-v^2) \) returned from eqn. 
(2) is the resultant of all the elastic spring elements in the viscoelastic constitutive law of the sample 
while the viscous dashpot elements, which can be nonlinear in general, do not play any role. Eqn. (2) 
is the result of a subtraction between the effects of the constitutive law just before and just after the 
rate-jump point [36], and the viscous dashpots are immaterial here because their deformations are 
continuous across the rate jump and are therefore subtracted out, i.e. they do not react to the rate jump.
Only the elastic springs react to the rate jump and this is why eqn. (2) contains only the net elastic 
modulus although the sample is viscoelastic. In the event if the spring and dashpot elements are time 
or load-history dependent, as would be the case for large deformations when the cytoskeleton, for 
example, becomes significantly modified, then the \( E/(1-v^2) \) returned from eqn. (2) will be the 
instantaneous value at the moment when the rate jump is applied [36]. The validity of this rate-jump
method in general has been verified in a wide range of soft materials on different mechanical testing platforms, including depth-sensing indentation on bone tissues, polymers and soft metals [18,37,38], micro-plate compression on stem-cell inserted collagen micro-masses for tissue engineering purposes [39], AFM nanoindentation of cells and soft polymers [19,20, 39], as well as macroscopic uniaxial tensile testing on polymers [40]. In the results below, the values of the Young modulus $E$ are reported and these were calculated from the measured reduced moduli $E/(1-\nu^2)$ by assuming the Poisson ratio $\nu$ of the cells to be 0.5.

**Statistics**

The data were analyzed with Student’s $t$ test to determine the significance between two variables or one-way analysis of variance (ANOVA) to calculate the significance when there were more than two variables. Associations between elastic modulus and cell migration and invasion were tested using Pearson’s product-moment correlation and Spearman’s rank correlation.
Results

Elastic modulus of TSCCs with different metastatic potentials

Among the TSCC cases (Table S1), Case 3 had LN metastasis and recurrence after one month and died six months postsurgery; however, Cases 1, 2 and 4 did not have LN metastasis, and no recurrence occurred during the 10-18 months of follow-up. Using the wound healing and transwell assays, we found that UM1 cells and UM1/Control cells had a higher migratory and invasive ability (i.e. higher metastatic potential) than UM2 cells and UM1/miR-138 cells, respectively (Figure 2). Similarly, primary cells obtained from Case 3 had the highest metastatic potential than those obtained from the other three cases (Figure 2). In another paired OSCC cell line, 1386Ln/control cells exhibited significantly elevated metastatic potential when compared to 386Ln/miR-138 cells (Figure S2).

Atomic force microscope nanoindentation via the rate-jump method was used to detect the elastic modulus of these TSCC cells with different metastatic potentials. As shown in Figure 1C, Table 1 and Table S2, UM1 cells, UM1/Control cells and 1386Ln/Control cells (with higher metastatic potential) had significantly lower elastic modulus values when compared to UM2, UM1/miR-138 and 1386Ln/miR-138 cells (with lower metastatic potential), respectively. Primary cells obtained from Case 3 also exhibited significantly lower elastic modulus when compared to primary cells obtained from the other three cases. The histograms for each sample were provided in supplemental Figure S3, in which the distribution of elastic modulus values for the less invasive cells showed a normal distribution. The relationship between cellular elastic modulus and metastatic potential was further examined and indicated a strong association between elastic modulus and metastatic potential as shown in Table S3. These results revealed that cells with higher metastatic potential had lower elastic modulus values than cells with lower metastatic potential, which means that they were softer or more deformable.

The relationship between EMT and elastic modulus of TSCC

To investigate the relationship between elastic modulus and EMT, TSCCs with different metastatic potential were observed under SEM and the expression of EMT markers (E-cadherin and vimentin) was also assessed. Figure 3 and Figure S4A show that the primary cells obtained from Case 3, UM1
cells, UM1/Control cells and 1386Ln/Control cells (all with lower elastic modulus and higher metastatic potential) were all spread out and had a fibroblastic morphology. On the other hand, primary cells obtained from the other three cases, UM2 cells, UM1/miR-138 cells and 1386Ln/miR-138 cells (all with higher elastic modulus and lower metastatic potential) were more adherent to one another showing promoted formation of cell clusters. More surface protrusions, microvilli and filopodia-like processes were found in cells with lower elastic modulus and higher metastatic potential.

Moreover, as shown in Figure 4, primary cells obtained from Case 3, UM1 cells and UM1/Control cells (with lower elastic modulus and higher metastatic potential) had higher expression levels of vimentin and lower expression levels of E-cadherin as compared with primary cells obtained from the other three cases, UM2 cells and UM1/miR-138 cells (with higher elastic modulus and lower metastatic potential), respectively. These observations are in agreement with the hypothesis that EMT may be related to the elastic modulus and metastatic potential.

The relationship between Cytoskeleton and elastic modulus of TSCC

As shown in Figures 5, 6, and Figure S4B, the F-actin cytoskeleton in TSCCs features a dense cortical layer underneath the plasma membrane and an isotropic network throughout the cell body. β-tubulin was localized in the space between the nucleus and the cell membrane. F-actin and β-tubulin intensity profiles in the primary cells obtained from Cases 1, 2 and 4, UM2 cells, UM1/miR-138 cells and 1386Ln/miR-138 cells (with higher elastic modulus and lower metastatic potential) were significantly higher than those in primary cells obtained from Case 3, UM1 cells, UM1/Control cells and 1386Ln/Control cells (with lower elastic modulus and higher metastatic potential), respectively. Moreover, UM2 cells, UM1/miR-138 cells and 1386Ln/miR-138 cells had a thick cortical layer of F-actin and a thick nuclear layer of β-tubulin. These results imply that lower elastic modulus in TSCC cells with higher metastatic potential are related to less organization and decreased intensity of F-actin and β-tubulin.

The relationship between nucleus size and elastic modulus of TSCC

As shown in Table 2 and Table S2, primary cells obtained from Case 3, UM1 cells, UM1/Control cells
and 1386Ln/Control cells (with lower elastic modulus and higher metastatic potential) had significantly smaller nucleus sizes and higher N/C ratios than primary cells obtained from the other three cases, UM2 cells, UM1/miR-138 cells and 1386Ln/miR-138 cells (with higher elastic modulus and lower metastatic potential), respectively. Thus, the nucleus size and the N/C ratio may lead to a reduction in the elastic modulus and an increase in metastatic potential.
**Discussion**

The elastic modulus is an inverse measure of the deformability or compliance of individual cells and is therefore important in the initiation of cancer [10]. Many studies have shown that malignant tumor cells display either lower elastic modulus (softer) and/or lower viscosity (less resistant to flow) [13, 21, 41, 42]. Although Remmerbach et al. recently showed that both oral epithelial cell lines and primary cancer cells have an increased deformability compared with their normal counterparts [22], not many studies have focused on oral cancer. Recently researchers also found that mechanical properties are a critical determinant for the efficiency of cancer cell invasion [11, 21, 30, 41, 42]. Swaminathan et al. found that cancer cells with the highest migratory and invasive potential are five times less stiff (with lower elastic modulus) than cells with the lowest migration and invasive potential [11]. In the present study on tongue cancer cells, we revealed that, just like other types of cancer, cells with lower metastatic potential from TSCC cell lines and patients had significantly higher elastic modulus (i.e., less deformable) than those of cells with higher metastatic potential. The elastic modulus of tongue cancer cells with lower metastatic potential is only around twice of that of cells with higher metastatic potential, which means that the difference between them is less significant than those reported in previous studies [11]. It is believed that this is due to the individual differences among different types of cancer cells. It should also be noted that, based on the elastic modulus measured in this work, the difference between elastic modulus of tongue cancer cells with high metastatic and low metastatic potential is less significant than those of other types of cancers, and therefore a protocol for accurate measurement of the elastic modulus of cancer cells is necessary for evaluating the metastatic potential. Thus, these findings suggest that cellular elastic modulus may be related to the metastasis of TSCC and can be used as a cell marker and a diagnostic parameter for TSCC metastasis. Moreover, the present study also shows that the rate-jump method for AFM nanoindentation can be used as a standard protocol for measuring the elastic modulus of living cells, since the measured values are intrinsic properties of the cells.

Tumor metastasis is the dominant cause of death in patients with TSCC. EMT is a critical step in the process of metastasis, which leads to cancer spread and treatment failure. Recently, studies from our
group and other groups have demonstrated that EMT is important in determining the biological behaviors and clinical prognosis of tongue cancer [5, 43]. For example, Sakamoto found that the development of delayed neck metastasis in stage I/II TSCC was closely related to the induction of EMT in primary tumor cells [43]. Our previous studies have suggested that downregulation of miR-138 is associated with mesenchymal-like cell morphology and enhanced cell migration and invasion [3-8]. Besides, EMT-associated changes were also found to influence cellular mechanics [44]. Buckley et al, found that alveolar epithelial cells (A549) with TGF-β1 stimulation gave rise to a significant increase in stiffness and induced EMT changes [44]. In the present study, we also found that TSCC cells with lower elastic modulus and higher metastatic potential had EMT changes. These observations revealed that EMT might lead to a decrease in elastic modulus and an increase in the metastatic potential of TSCC.

The cytoskeleton is the structural framework that predominantly shapes a cell and provides its mechanical rigidity. According to the tensegrity theory, a cell can be viewed as a tensional structure in which actin filaments bear tensile loads and microtubules bear compression loads [30]. Among the different cytoskeletal filaments, F-actin provides the highest resistance to deformation [10]. Changes in the cytoskeleton would therefore reflect the overall mechanical properties of the cell [21, 24, 45-47]. Compared with nonmetastatic cancer cells, Guck et al. found that the increased deformability (decrease in elastic modulus) of cancer cells with metastatic competence appears to be accompanied by a reduction in the amount of F-actin [21]. In the present study, we observed differences in the cell cytoskeleton (F-actin and β-tubulin) that were accompanied with changes in cell metastatic potential and elastic modulus. In the cells with lower elastic modulus and higher metastatic potential, the cytoskeleton was reduced and less organized, with lower levels of F-actin gathered underneath the plasma membrane and decreased levels of β-tubulin organized around the nucleus. These observations confirmed that changes in the cytoskeleton may regulate the cellular elastic modulus and induce cell motility, invasion and metastasis.

Directed-cell migration, which often requires nuclear reshaping to allow passage of cells through narrow openings, is dependent not only on changes in cytoskeletal elements but also on global
chromatin condensation [25, 48]. Rolli et al. found that the diameters of the nuclei of pancreatic epithelial cancer cells (Panc-1) are reduced by 12% after sphingosylphosphorylcholine treatment, which can enhance the invasive behavior and deformability of Panc-1 [24]. Stroka et al. also found that increased cell stiffness correlates with an increase in cell area [49]. Similar to these studies, we demonstrated that TSCC cells with higher metastatic potential had smaller nuclei and higher N/C ratios than cells with lower metastatic potential. Thus the small nuclei may decrease the overall elastic stiffness of the cells and allowed them to pass through narrow openings, and this corresponds to an increased invasion of TSCC cells.

From the above, the present findings revealed a close relationship between cellular elastic modulus and the metastatic potential of TSCC cells. The EMT, cytoskeleton, nucleus size and nucleus/cytoplasm ratio are all factors contributing to a change of the cellular elastic modulus. Combined with our previous studies [3-8], we hypothesize that TSCC cells with higher metastatic potential undergo a series of molecular changes, such as deregulation of miR-138, then induce EMT and cytoskeleton disorganization, modulate the nucleus size and nucleus/cytoplasm ratio, and finally have their cellular elastic modulus reduced and metastasis mediated. Thus, the cellular elastic modulus may be used to grade the metastatic potential of TSCC and as a diagnostic parameter for TSCC metastasis. In addition, since the present rate-jump method of AFM nanoindentation takes only minutes rather than hours to complete as in invasion assays, this is an efficient and reliable approach for classifying cancer cells to enable broad screening of suspicious lesions with metastasis of TSCC.

To conclude, the significance of this work can be summarized as follows. 1) We used a new method (rate-jump AFM nanoindentation) to measure the elastic modulus of cancer cells in which the viscous effects can be eliminated giving rise to reliable and intrinsic measurements of the elastic modulus. 2) Although tongue squamous cell carcinoma (TSCC) is significantly more aggressive than other forms of oral squamous cell carcinomas, how metastasis is linked to cellular biomechanics is basically unknown for this type of carcinoma. Using both TSCC cell lines and TSCC cells from patients with different metastatic potential, we found a close relationship between the elastic modulus and metastatic potential of TSCC cells. 3) With the efficient rate-jump AFM nanoindentation method, we
demonstrated that the cellular elastic modulus can be used as a diagnostic parameter for grading the metastatic potential of TSCC. 4) By indenting on the nucleus location of the cells, we showed that the cellular elastic modulus (and hence metastatic potential) of TSCC cells are related to the filament organization in the cytoskeleton, the nucleus size and also the nucleus/cytoplasm ratio.
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Figure legends

Figure 1 The elastic modulus values of TSCCs with different metastatic potentials
A: Loading schedule for the rate jump method. When doing experiment, the tip was first retracted for 2 μm within 1s, and then was made to approach the cells with a constant speed 0.55μm/s for 10s and then held for 30s. After that, the tip was retracted with a constant speed 0.1μm/s for 10 s. The final unload point at 41s was treated as the rate-jump point where eqn. (2) was applied.
B: The indentation force vs time curve in rate jump tests using the loading profiles in Fig. 1A.
C: The elastic modulus of TSCC cells. Inside each box chart, the body denotes the values of 90% tested samples in the region, the number on top of each box chart is the mean value for each group, which is also the line inside the body box. The two ends of each box chart are the maximum and minimum value. Significant differences were found between Primary cells obtained from Case 3 and primary cells obtained from the other three cases; UM1 cells and UM2 cells; UM1/Control cells and UM1/miR-138 cells (P<0.05).

Figure 2 The metastatic potential of TSCC cells
The migration ability of TSCC cells was assessed using a wound healing assay (A, B, C). The invasion ability of TSCC cells was assessed using a transwell invasion assay (D, E, and F). UM1 cells, UM1/Control cells and primary cells obtained from Case 3 had a higher migratory and invasive ability than UM2 cells, UM1/miR-138 cells and primary cells obtained from the other three cases, respectively. *: P< 0.05.

Figure 3 The morphology of TSCCs with different elastic modulus
The cell morphology of TSCC primary cells and cell lines were detected under SEM as described in the Materials and Methods section. Primary cells obtained from Case 3, UM1 cells and UM1/Control cells had a fibroblastic morphology. Primary cells obtained from the other three cases, UM2 cells and UM1/miR-138 cells had a more adherent phenotype, which promoted the formation of cell clusters. ➝: microvilli; ➞: filopodia-like processes.
**Figure 4. The expression of EMT markers in TSCCs with different elastic modulus**

The EMT markers E-cadherin and vimentin were detected as described in the Materials and Methods section. Using immunohistochemistry (A, B), Primary cells obtained from Case 3 (in both cancer and LN tissue) had a higher expression level of vimentin and a lower expression level of E-cadherin compared with primary cells obtained from the other three cases, \( P < 0.05 \). Semi-quantitative analysis of the immunohistochemistry is shown in B. Relative staining intensities were analyzed by using Image Pro Plus 6 software. Western blot analysis (C, D) demonstrated a higher expression level of vimentin and a lower expression level of E-cadherin in UM1 cells compared to UM2 cells (C), or UM1/Control cells compared to UM1/miR-138 cells (D).

**Figure 5 F-actin in TSCCs with different elastic modulus**

F-actin was detected with immunofluorescence as described in the Materials and Methods section. The F-actin cytoskeleton features a dense cortical layer underneath the plasma membrane and an isotropic network throughout the cell body. Nuclear staining showed that the shadows within the cells coincide with the nucleus (not included in the image). F-actin intensity was significantly decreased in the cells with lower elastic modulus [i.e., primary cells obtained from Case 3, UM1 cells and UM1/Control cells] compared with cells with higher elastic modulus [i.e., primary cells obtained from the other three cases, UM2 cells and UM1/miR-138 cells], respectively. \( P < 0.05 \).

**Figure 6 \( \beta \)-tubulin in TSCCs with different elastic modulus**

\( \beta \)-tubulin was detected with immunofluorescence as described in the Materials and Methods section. The \( \beta \)-tubulin was localized in the space between the nucleus and the cell membrane. Nuclear staining showed that the shadows within the cells coincide with the nucleus (not included in the image). \( \beta \)-tubulin intensity were significantly increased in primary cells obtained from the other three cases, UM2 cells and UM1/miR-138 cells compared with primary cells obtained from Case 3, UM1 cells and UM1/Control cells, respectively. \( P < 0.05 \).
Table S1 Clinical characteristics of four TSCC cases

<table>
<thead>
<tr>
<th>Case</th>
<th>Gender</th>
<th>Age (ys)</th>
<th>TNM</th>
<th>Pathology</th>
<th>LN metastasis</th>
<th>Recurrence</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>male</td>
<td>53</td>
<td>T3N0M0</td>
<td>Middle-D</td>
<td>N</td>
<td>N</td>
<td>18m</td>
</tr>
<tr>
<td>2</td>
<td>female</td>
<td>37</td>
<td>T3N0M0</td>
<td>High-D</td>
<td>N</td>
<td>N</td>
<td>12m</td>
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<tr>
<td>3</td>
<td>male</td>
<td>59</td>
<td>T2N1bM0</td>
<td>Middle-D</td>
<td>Y</td>
<td>1m ps</td>
<td>die 6m ps</td>
</tr>
<tr>
<td>4</td>
<td>male</td>
<td>59</td>
<td>T1N0M0</td>
<td>Middle-D</td>
<td>N</td>
<td>N</td>
<td>10m</td>
</tr>
</tbody>
</table>

*: ys: years; ps: post-surgery; D: differentiation; Y: yes; N: no; m: months
Table S2 Elastic modulus and nucleus size of 1386Ln/Control cells and 1386Ln/miR-138 cells

<table>
<thead>
<tr>
<th></th>
<th>nucleus size (µm²)</th>
<th>N/C ratio</th>
<th>$E_{\text{elastic}}$ (kPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1386Ln/Control (n=38)</td>
<td>178.19±48.17*</td>
<td>0.39±0.05*</td>
<td>4.51±1.14*</td>
</tr>
<tr>
<td>1386Ln/miR-138 (n=59)</td>
<td>207.30±78.34</td>
<td>0.25±0.03</td>
<td>6.77±1.50</td>
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</tbody>
</table>

The data are presented as mean±standard deviation. *: compared between 1386Ln/Control cells and 1386Ln/miR-138 cells, $P<0.05$. 
Table S3 The association between elastic modulus/migratory and elastic modulus/invasion

<table>
<thead>
<tr>
<th></th>
<th>Pearson</th>
<th></th>
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<th>Spearman</th>
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<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>r</td>
<td>p value</td>
<td>r</td>
<td>p value</td>
<td></td>
</tr>
<tr>
<td>Elastic modulus/relative migration</td>
<td>TSCC cell lines</td>
<td>-0.688</td>
<td>0.000</td>
<td>-0.677</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TSCC cases</td>
<td>-0.682</td>
<td>0.000</td>
<td>-0.782</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>Elastic modulus/relative Invasion</td>
<td>TSCC cell lines</td>
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<td>-0.632</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TSCC cases</td>
<td>-0.720</td>
<td>0.000</td>
<td>-0.782*</td>
<td>0.000</td>
<td></td>
</tr>
</tbody>
</table>

TSCC cell lines include UM1, UM2, UM1/Control and UM1/miR-138; TSCC cases include case 1, case 2, case 3 and case 4. The TSCC cell lines and cases were classified by their migration and invasion (shown in Figure 2 and Figure S2), with the migration and invasion value of UM2 cells or case 1 falling into 1, and the migration and invasion value of other TSCC cell lines and cases was described in comparison with UM2 cells or case 1, respectively.
Supplementary figure legends:

Figure S1 TSCC cells were mechanically probed by AFM at their most bulging part

Figure S2 The migration and invasion ability of another paired OSCC cell lines (1386LN/Control and 1386LN/miR-138)

The migration ability was assessed using a wound healing assay (A). The invasion ability was assessed using a transwell invasion (B). 1386LN/Control cells had a higher migratory and invasive ability than 1386LN/miR-138 cells. *: $P < 0.05$.

Figure S3 Histograms of elastic modulus for each sample from TSCC cell lines and patients

Figure S4 Morphology and cytoskeleton of another paired OSCC cell lines (1386LN/Control and 1386LN/miR-138)

A: The morphology of 1386Ln/Control and 1386Ln/miR-138 cells under SEM. 1386Ln/Control cells appeared spread out assuming a fibroblastic morphology. 1386Ln/miR-138 cells appeared more adherent to one another and promoted formation of cell clusters.

B: Cell cytoskeleton F-actin and β-tubulin in 1386Ln/Control and 1386Ln/miR-138. The F-actin cytoskeleton features a dense cortical layer underneath the plasma membrane and an isotropic network throughout the cell body. The F-actin stress fiber was found to reduce and less organize in higher metastatic potential 1386Ln/Control as compared to 1386Ln/miR-138. *: $P < 0.05$. The β-tubulin was localized in the space between nucleus and cell membrane, the β-tubulin was found to increase and more organize (around nucleus) in 1386Ln/miR-138 as compared to 1386Ln/Control. *: $P < 0.05$. 

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Fig. 3

Case 1  Case 2  UM1  UM2
Case 3  Case 4  UM1/control mimic  UM1/miR-138 mimic