Chitosan films promote formation of olfactory neurospheres and differentiation of olfactory receptor neurons*

Sheng-Tien Li¹, Tai-Horng Young¹, Chi-Te Wang³, Tsung-Wei Huang²,³,⁴

¹ Institute of Biomedical Engineering, College of Medicine and College of Engineering, National Taiwan University, No. 1, Sec. 1 Jen-Ai Road, Taipei 100, Taiwan
² Department of Electrical Engineering, College of Electrical and Communication Engineering, Yuan Ze University, Taoyuan, Taiwan
³ Department of Otolaryngology, Far Eastern Memorial Hospital, Taipei, Taiwan
⁴ Department of Health Care Administration, Oriental Institute of Technology, Taipei, Taiwan

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Background: Olfactory dysfunction significantly impairs the life quality of patients. Therefore, a model needs to be developed for anosmia. Chitosan is a biodegradable natural polysaccharide that has been widely studied for regenerative purposes in the nervous system. However, whether chitosan promotes differentiation of olfactory receptor neurons or regulates formation of neurospheres in the olfactory system remains unexplored.

Methodology: Olfactory neuroepithelial cells were isolated from embryonic wistar rats on day 17, and cultured on controls and chitosan films for 12 days. The effects of treatment were assessed using immunocytochemistry, quantitative polymerase chain reaction and western blots following culturing. The substrate of poly-L-lysine-co-laminin was adopted as a control.

Results: In contrast to the flat layer on controls, olfactory neuroepithelial cells form olfactory neurospheres on chitosan films with steadily increasing diameter. The olfactory neurospheres contain basal cells, as well as immature and mature olfactory receptor neurons. The expression level of olfactory marker protein is higher on chitosan films than those on controls in gene and protein levels, and the olfactory transduction elements also express a similar trend. Mature olfactory receptor neurons are found predominantly at the periphery of the olfactory neurospheres.

Conclusions: Chitosan films not only facilitate formation of olfactory neurospheres, but also promote differentiation of olfactory receptor neurons. Chitosan is a potential biomaterial to establish an in vitro culture model to treat olfactory dysfunction in future.

Key words: chitosan; anosmia; olfactory receptor neurons; olfactory mucosa

Introduction

Olfactory dysfunction significantly affects quality of life by altering appetite, reducing the ability to identify nearby dangerous situations, and even resulting in depression (1,2). Hyposmia varies in the general population from 13% to 18%, and anosmia from 4% to 6% (3). In the olfactory system, the olfactory neuroepithelium (ON) operates the first relay point of odor sensation. The mammalian ON, which harbors olfactory receptor neurons (ORNs), is located in the nasal cavity, while this neuroepithelium is split into three compartments, basal, middle and apical. The basal compartment is close to the basal lamina of the epithelium, and consists of basal cells that express the achaete-scute family bHLH transcription factor 1 (Ascl1); the middle compartment contains immature and mature ORNs that individually express βIII tubulin and olfactory marker protein (OMP), and the apical layer of the ON comprises sustentacular cells (4). Particularly, unlike other cranial neurons, ORNs can be continuously replaced with new ones during normal neuronal turnover and environmental insults (5). However, this unique ability deteriorates with age and in response to various toxic factors, including smoking, viruses, inflammation, medication and environmental factors, leading to olfactory dysfunction (6,7). Although many approaches, including vitamin A, zinc and various growth factors, have promising findings, they have not been applied as clinical practice methods for olfactory dysfunction (8-10). Topical or systemic application of glucocorticoids is a common clinical treatment, but its therapeutic effect is controversial. Oral steroids can improve olfaction in sinonasal disease, but the reduced sense
of smell tends to recur shortly after the treatment is stopped (13).
Topical steroids are well tolerated and are effective treatments of olfactory loss associated with severe nasal and sinus disease (12). However, it appears to have a minor effect on olfactory dysfunction in some reports, especially when considering long-term changes (13,14).

A therapeutic trend is transplantation and engraftment of basal cells in the damaged ON (15). The ON can be regenerated using autologous olfactory basal cells to repair damage, and to alleviate the consequent dysfunction (16). Hence, an in vitro culture system that can encourage olfactory neuronal differentiation and expansion of basal cells is essential for the regeneration of ON. Among culture conditions, biomaterials play a crucial role in mediating cellular morphology, proliferation and differentiation. Chitosan is a cationic polysaccharide with a variable number of randomly located D-glucosamine and N-acetyl-glucosamine groups (17). This biopolymer has been demonstrated to bridge large gaps in the peripheral nerves with chitosan-coated silicone tubes. Chitosan is also reported to regulate formation of neurospheres (18-20). Compared with the adherent culture, this sphere formation in vitro culture has numerous multipotent progenitors to facilitate incorporation into host epithelium, which is a requirement for eventual translational use (21). Therefore, neurospheres can serve as biomarkers for engraftment capacity. However, whether chitosan can promote differentiation of olfactory neuroepithelial cells (ONCs) or regulate formation of olfactory neurospheres remains unexplored. The objective of this study is to evaluate the effect of chitosan films on development and differentiation of ONCs in an in vitro culture system.

**Materials and methods**

**Cell preparation and culture**

All experimental procedures and animals used were approved by the National Taiwan University Animal Care and Use Committee (agreement number: 20140357). Briefly, the ON was obtained from embryonic wistar rats on day 17 (E17) using the protocol in our previous study (22). Briefly, the ON was cut into small pieces in cold Hank’s balanced salt solution and then digested with 0.125% Trypsin/EDTA for 30 minutes at 37°C. After digestion, the dissociated cells were collected by centrifugation and resuspended in DMEM-F12 (serum-free) containing 20 ng/mL bFGF, 20 ng/mL EGF, 2% B27 and 1% penicillin/streptomycin. The half medium was added on alternate days. For the experiment, harvested cells were seeded on tissue culture plates pre-coated with poly-L-lysine-co-laminin (PLL-La) or on chitosan films at 37°C in a humidified 5% CO₂ and 95% air atmosphere.

**Preparation of substrate-coated culture plates**

Substrate-coated was coated on tissue culture plates according to a protocol detailed previously (23). Briefly, culture plates were pretreated with PLL (P2636, Sigma-Aldrich, St. Louis, MO, USA) pretreated with PLL (P2636, Sigma-Aldrich, St. Louis, MO, USA) to a protocol detailed previously. Substrate-coated was coated on tissue culture plates according to a protocol specified previously. Preparation of substrate-coated culture plates was conducted as follows:

1. The nascent films were neutralized using 0.5 N NaOH. Finally, these films were sterilized in silicone tubes. Chitosan is also reported to regulate formation of olfactory neuroepithelial cells (ONCs) or on chitosan films at 37°C in a humidified 5% CO₂ and 95% air atmosphere.

2. Table 1. List of gene-specific primers used for qPCR assay.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Sequence</th>
<th>Product size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ascl1</td>
<td>ACTTGACTCTATGGCAGGTT</td>
<td>151bp</td>
</tr>
<tr>
<td>βIII tubulin</td>
<td>ATGAGGGAGGTGTGAGCAG</td>
<td>102bp</td>
</tr>
<tr>
<td>OMP</td>
<td>CTTGATTTCCAGACGACG</td>
<td>122bp</td>
</tr>
<tr>
<td>Gαs</td>
<td>TTGACACGAGAAGACGCTG</td>
<td>148bp</td>
</tr>
<tr>
<td>ADcy3</td>
<td>CGTGGTCTCTGTTGTTGTCG</td>
<td>124bp</td>
</tr>
<tr>
<td>GAPDH</td>
<td>GCGCTGCTTCCTGTTGACAGC</td>
<td>101bp</td>
</tr>
</tbody>
</table>

at a concentration of 52 μg cm⁻². Then, 2 μg cm⁻² laminin was coated on top of the PLL. The culture plates were then put in the incubator at 37°C until their use at the same day. The excessive laminin was removed immediately before seeding ONCs. This group was adopted as controls.

**Preparation of chitosan films culture plates**

Chitosan was coated on tissue culture plates using methods described in a previous study, with some modifications (20). Briefly, 0.5 mL 1% (w/w) chitosan solution (C-3646, Sigma-Aldrich) dissolved in 0.5 N acetic acid was added into each well of six-well plates and evaporated in a convection oven at 60°C until the casting solution became solid. Then, the nascent films were neutralized using 0.5 N NaOH. Finally, these films were sterilized in 70% alcohol under ultraviolet light, and rinsed extensively with distilled water, before being used for cell culture.

**Morphological examination**

Cell morphology was observed under an inverse phase contrast microscope (TS-100, Nikon, Tokyo, Japan) and scanning electron microscope (S-4800, Hitachi, Tokyo, Japan). The diameter and projected area of spheroids were calculated using the software Image J in ten randomly selected fields for each sample. Only those with the diameter of more than 30μm were considered spheroids. Ratio of olfactory neurosphere projected area was defined by a percentage of projected area of neurosphere in a view.

**Reverse transcription- quantitation polymerase chain reaction (RT-qPCR)**

Total RNA was extracted from the cultured cells using a Trizol reagent (Invitrogen, CA, USA). The synthesis of first-strand cDNA was performed by the protocol from High-Capacity cDNA Reverse Transcription Kits (Applied Biosystems, Foster City, CA, USA). The cDNA was applied as the template for the qPCR assay.
of specific gene expression performed with the gene-specific primers listed in Table 1. The gene expression level was analyzed and normalized to GAPDH for each cDNA sample. Relative quantity of gene expression was calculated with the expression levels each group divided by those in controls at day 6.

Immunocytochemical staining

The samples were fixed with 4% paraformaldehyde and permeabilized with 0.1% Triton X-100 (X100; Sigma-Aldrich) for 10 minutes at room temperature. After fixing, the samples were blocked in 3% bovine serum albumin for 20 minutes, and incubated with primary antibodies at 4°C overnight. The primary antibodies and their dilution utilized were used in this study: anti-Ascl1 (Ab74065; 1:1000; Abcam, Cambridge, UK), anti-βIII tubulin (Ab118627; 1:1000; Abcam), anti-OMP (ab62144, 1:100; Abcam), anti-OMP (NB110–74751, 1:100; Novus, CO, USA), anti-olfactory neuron specific-G protein (Golf) (GTX110520; 1:100; GeneTex, CA, USA), anti-adenylate cyclase type 3 (ADCY3) (Ab125093; 1:1000; Abcam) and anti-5-bromo-2′-deoxyuridine (BrdU) (MAB3424; 1:1000; Millipore, Billerica, MA, USA). The samples were then washed and incubated with Alexa Fluor 488 and/or Alexa Fluor 555-conjugated secondary antibodies, and counterstained with DAPI. Particularly, before day 6, cells were incubated with 10μM BrdU for 24 hours to identify S-phase cells since BrdU, a thymidine analog, would incorporate into the newly synthesized DNA in replicating cells. Images were taken with Leica DMi6000 microscope and a confocal microscope (LSM510, Carl Zeiss, Germany). Further, the BrdU-positive cells within olfactory neurospheres were counted and divided by the total number using ImageJ in at least ten randomly selected fields.

Western blot analyses

Each lysate was obtained from individual culture wells using radioimmunoprecipitation assay lysis buffer containing protease inhibitor cocktail (Roche Diagnostics, Indiana, IN, USA) to avoid protein degradation. The protein samples were then denatured and separated by 12% SDS-PAGE gels, and blotted onto PVDF membranes (Millipore, Billerica, MA, USA). After blocking in a CIS blocking buffer (CIS-Biotechnology, Taiwan) at room temperature for 60 seconds, the membranes were probed with the specific primary antibody at 4°C overnight, washed, incubated in HRP-conjugated secondary antibodies, and finally visualized using enhanced chemiluminescence (ECL; Millipore). The antibodies used in this study were anti-Ascl1 (Ab74065; 1:1000), anti-βIII tubulin (Ab118627; 1:1000), anti-OMP (Ab98124; 1:1000), anti-olfactory neuron specific-G protein (Golf) (Ab74049; 1:1000) and anti-adenylate cyclase type 3 (ADCY3) (Ab125093; 1:5000) (all from Abcam, Cambridge, MA, USA). After detachment of previous primary antibodies, the membranes were also probed with GAPDH antibodies.
which acted as an internal control. Immunoreactive bands were quantified using Image J.

Measurement of calcium transients

Olfactory neurospheres cultured on chitosan films at day 12 were plated on gelatin-coated 35mm coverslips for 1 day prior to testing. Briefly, cells were loaded with 1μM fura-2 AM (F-1225, Invitrogen) and 20μg/ml Pluronic® F-127 (P3000MP, Invitrogen) at room temperature for 30 minutes in a culture medium, followed by 2 rinses with Hank’s Balanced Salt Solution containing HEPES for 30 minutes to cleave the excess fura-2 AM. A fura2-loaded cell was then respectively exposed to the alternating excitation wavelengths of 340nm and 380nm for Ca^{2+}-bound and Ca^{2+}-free dyes by IonOptix system (IonOptix, Milton, MA, USA). A solution of 300μM 3-isobutyl-1-methylxanthine (IBMX, Invitrogen) plus 10μM forskolin (FSK) acted as a stimulator to induce intracellular calcium transients, according to a previous protocol (23). The calcium waves were described using the ratio 340/380. All parameters were analyzed off-line using IonWizard (IonOptix, Milton, MA, USA).

Statistical analysis

The results were observed at least three independent experiments, with the data expressed as mean ± standard deviation (SD). Statistical significance was evaluated using paired t-test or one way ANOVA followed by Tukey’s post-hoc test with p < 0.05 being considered significant.

Results

Morphology of ONCs on chitosan films vs. controls (PLL-La)

Cells undergo morphological changes after contact with bio-materials to stabilize the cell-substrate interface, which plays a significant role in regulating subsequent cell behaviors such as survival, proliferation and differentiation. In controls, cells with fusiform-like morphology attached to the substrate and extended neurites at day 3 (Figure 1A (a)). Fusiform cells in this group were variable in size and shape, and lengthened neurites over time. The ONCs reached approximately 80% confluence until day 9, and retained flat and bipolar-like morphology. Lots of round cells appeared at day 12 (Figure 1A (d)). Conversely, ONCs cultured on chitosan films exhibited small neurospheres by day 3 (Figure 1A (e)). Small colony-like structures were sometimes formed. The number and size of those neurospheres gradually rose after day 6, while maintaining semi-attachment to chitosan films. The diameter of olfactory neurospheres steadily increased from 46μm ± 13.6μm at day 3 to 132μm ± 24.5μm at day 12, and BrdU was also expressed within olfactory neurospheres (7.5% ± 2.7%, Figures 1B–C). BrdU, an analog of thymidine, labels newly synthesized DNA in mitotic cells, and is commonly used in the detection of proliferating cells. Additionally, this projected area covered by neurospheres was larger on chitosan films than on controls. The percentages of the projected area on chitosan films were 25% ± 3.9%, compared to 4% ± 4.1% on controls at day 12, showing a significant difference (p < 0.05) (Figure 1D).

The ORN differentiation of ONCs on chitosan films vs. controls

ONCs have been well characterized for stage-specific markers (4). In the ONC development, a marker of basal cells, Ascl1, plays a crucial choice point at an early stage in the ORN lineage to initiate a differentiation program. OMP is a prototypical form, and is
Li et al. expressed almost exclusively in mature ORNs, while βIII tubulin is a neuron-specific isoform of tubulin, and is widely accepted as a marker of immature ORNs. At day 6, immunofluorescence revealed that ONCs on both groups expressed one of these stage-specific markers, Ascl1, βIII tubulin and OMP (Figure 2). βIII tubulin was clearly present throughout the neuron, in soma, dendrites and axons (Figure 2E). At day 12, the expression level of βIII tubulin significantly fell on both groups (Figs. 2H & 2K), but the expression level of OMP was higher on chitosan films than on controls (Figures 2L & 2I). The distribution of mature ORNs with positive OMP gathered at spheroid cells was also noteworthy (Figure 2L). Further evaluation of gene expression with RT-qPCR revealed 2.0 ± 0.1 fold upregulated Ascl1 and 1.3 ± 0.1 fold upregulated βIII tubulin genes on chitosan films comparing to those on controls at day 6 (Figures 3A & 3B, p < 0.05). Notably, the mRNA ratio of OMP relative to the internal gene GAPDH was significantly higher on chitosan films than on controls (5.7 ± 1.0 fold) at day 12 (Figure 3C, p < 0.05). Western blot analyses of Ascl1, βIII tubulin and OMP further confirmed the results (Figure 4).

Figure 5. Immunocytochemical detection of Golf and ADCY3 on controls (a–b) and chitosan films (c–d) (A). ONCs are immunos toned at day 12, and identified with Dylight488 (green). Nuclei are labeled with DAPI (blue). The micrographs show that Golf and ADCY3 are respectively located in cell bodies and membranes. RT-qPCR measurements for functional elements on controls and chitosan films. ONCs on chitosan films have significantly higher levels than controls of Golf mRNA (B) and ADCY3 mRNA (C). Western blot analyses of Golf, ADCY3 and GAPDH proteins in ONCs at day 12 (D). Golf/GAPDH ratios (E). ADCY3/GAPDH ratios (F). The expressions of Golf and ADCY3 protein in ONCs are significantly higher on chitosan films than on controls. Fura-2-loaded olfactory neurospheres are stimulated with FSK + IBMX, and the perfusion time of stimulators is indicated by the horizontal bars above the traces (G). Scale bar = 100μm. Results are expressed as means ± SD. Asterisk indicates p < 0.05.

Olfactory transduction elements of ORNs on chitosan films vs. controls
To further determine whether ORNs have functional characteristics in these cultures, the levels of Golf and ADCY3 were measured, as these are essential elements and hallmarks of functionally mature ORNs (24). Immunofluorescence revealed at day 12 that ORNs on chitosan films expressed more Golf and ADCY3 than those on controls (Figure 5A). The Golf and ADCY3 were respectively found throughout their cell bodies and membranes, in accordance with previous reports (25) (Figure 5A, micrographs). The chitosan films and controls had significantly different immunofluorescence patterns. The distribution of Golf- and ADCY3-positive cells also gathered at spheroid cells. In comparison to controls, the gene expression level of ONCs on chitosan films exhibited significant upregulation of Golf and ADCY3 genes at day 12, at 4.6-fold and 3.2-fold respectively (Figures 5B & 5C, p < 0.05). Western blot analyses of Golf and ADCY3 further confirmed these findings (Figure 5D-F). Particularly, the olfactory neurospheres showed a typical response peak of intracellular calcium in response to IBMX+FSK, suggesting that olfactory neurospheres cultured on chitosan films would remain functional ORNs (Figure 5G).
Analysis of olfactory neurospheres

Apparent neurospheres appeared at day 6, with an average diameter of 71.3μm ± 13.7μm. The small population of multicellular spheroids on chitosan films had fine filamentous extensions from their terminal knobs. They are cilia-like structures (B, arrow). (Magnification ×6000). Representative confocal images show cellular composition of chitosan-expanded olfactory neurospheres. Photographs of sectioned olfactory neurospheres show βIII Tubulin (green, C) and double staining for OMP (green) with Ascl1 (red, D), Golf (red, E) or ADCY3 (red, F). Nuclei are labeled with DAPI (blue). Diagram of the cellular distribution within olfactory neurospheres (G). Scale bar = 20μm.

Discussion

Several groups of researchers have developed primary culture systems derived from ON in embryonic, neonatal or adult murine to study neurogenesis in the past two decades (22,23). In particular, the ON still contains many immature ORNs that are able to further mature in vitro at the E17 stage (27). According to our previous study, these primary ONCs simultaneously expressed markers of horizontal basal cells and proliferation, i.e. cytokeratin 5 and BrdU, at day 3 (22). Among substrates adopted as controls, PLL has a significantly positive effect on proliferation of ONCs, and laminin is an extracellular matrix within the ON. Both substrates not only improve axon outgrowth of ORNs and its pathfinding in the olfactory system, but are also widely used as substrates in ONC cultures (22,23). Therefore, this investigation harvested rat ONCs at E17, and utilized PLL-La as a substrate for comparison (23). As depicted in Figure 1A (a–d), ONCs can proliferate on controls with variable morphology to reach confluency after 12 days. However, in comparison to the two-dimensional monolayer culture, providing more physiologically relevant information and creating an in vivo similarity in three-dimension culture systems have many advantages, which may benefit treatment of olfactory dysfunction in the future.

Chitosan has been widely utilized in tissue engineering of neurological and nasal systems, as it bridges large gaps in the peripheral nerves with chitosan-coated silicone tubes, and exhibits permeation-enhancing properties for mucosal drug delivery (18,29,30). Although several reports have demonstrated that chitosan films can promote formation of spheres (19,20), no reports have considered whether chitosan promotes formation of ONCs. This work found that ONCs cultured on chitosan films form olfactory neurospheres, in contrast to the almost flat layer on controls (Figure 1A). Cells within the olfactory neurospheres express the markers of basal cells and proliferation, i.e. Ascl1 and BrdU (Figure 6D & Figure 1C). The expression of BrdU within olfactory neurospheres explains why the diameter of spheres steadily rises during culture periods (Figure 1B–C). These findings indicate that chitosan can promote formation of olfactory neurospheres with numerous basal cells, which may be valuable for translational purpose in the future.

Another important issue is whether chitosan promotes differentiation and maturation of ORNs. This work indicates that ONCs express more terminal differentiation markers of ORNs, i.e. OMP, Golf and ADCY3 on chitosan films in gene and protein levels than on controls, implying a significantly greater number of mature ORNs on chitosan films. Additionally, many olfactory neurons develop cilia-like structures extending from the process of an ORN (Figure 6B). The confocal microscope analyses of olfactory neurospheres reveal that mature ORNs expressing OMP, Golf and ADCY3 are co-localized within the out-layer of the spheroids (Figure 6D–F). This finding indicates that spheroids may start to develop polarity and behave like their in vivo counterparts (31) (Figure 6G).

Conclusion

This investigation is the first to show that chitosan promotes...
formation of olfactory neurospheres that not only maintain proliferation of basal cells, but also enhance ORN maturation. However, whether these olfactory neurospheres can be applied to facilitate ON regeneration in vivo needs further investigation.

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Authorship contribution
STL: study design, acquisition and interpretation of data, drafting of the manuscript, final approval of the version, accountability for all aspects of the work. THC, CTW, and TWH: study design, acquisition and interpretation of data, revision of the manuscript, final approval of the version, accountability for all aspects of the work.

Conflict of interest
The authors declared that no conflict of interest exists.

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