Development of a Decellularized Hypopharynx with Vascular Pedicle Scaffold for use in Reconstructing Hypopharynx

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Development of a decellularized hypopharynx with vascular pedicle scaffold for use in reconstructing hypopharynx

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Abstract: Background: Hypopharynx reconstruction after hypopharyngectomy is still a great challenge. Perfusion decellularization is for extracellular matrix (ECM) scaffolding and had been used in organ reconstruction. Our study aimed to prepare an acellular, natural, three-dimensional (3D) biological hypopharynx with vascular pedicle scaffold as the substitute materials to reconstruct hypopharynx. Result: Scanning electron microscope (SEM) and immuno showed that the decellularized hypopharynx with vascular pedicle scaffold retained intact native anatomical ECM structure. Myoblasts were observed on the recellularized scaffolds with bone marrow mesenchymal stem cells (BMSCs) induced by 5-azacytidine implanted in the rabbit greater omentum by immunohistochemical analysis. Conclusion: The decellularized hypopharynx with vascular pedicle scaffold prepared by detergent perfusion in our study has an potential to be an alternative material to pharynx reconstruction.

Keywords: decellularization, extracellular matrix, hypopharynx, vascular pedicle, hypopharyngeal squamous cell carcinoma

1. Introduction

Hypopharyngeal squamous cell carcinoma (HSCC) accounts for approximately 3% to 5% of all head and neck squamous cell carcinomas (HNSCC), and is less prevalent than most other major HNSCC, such as the laryngeal cancer and oral squamous cell carcinomas (1, 2). However, these patients with HSCC have worse prognosis than other types of HNSCC since the tumor in these patients has developed into stage III or IV when diagnosed (1, 3, 4). The current treatment methods of HSCC mainly include surgical resection and concurrent chemoradiotherapy (CRT). Hypopharynx refers to the muscular canal from the upper margin of hyoid bone to the lower margin of cricoid cartilage. The tumor may invade larynx, epiglottis or tongue, etc. The surgical resection of advanced hypopharyngeal cancer is so extensive that primary pharyngeal closure cannot complete directly or may lead to a high risk for pharyngocutaneous fistula postoperatively (5-8). Furthermore, the wide range of resection will lead to the loss of voice and swallowing function, which reduce the patients’ life quality. Reconstruction is critical to maintain hypopharynx functions like voice and swallowing, as well as improve patient’s life quality.

Hypopharynx reconstruction has been a great challenge. Recently, the common materials used to reconstruct hypopharynx are autogenous flaps, including pedicled flaps and free flaps. The
pedicled flaps include pectoralis major pedicled flaps, supraclavicular artery flaps, and free flaps include radial forearm free flap, anterolateral thigh free flap, jejunal free flaps, temporomandibular free flaps, serratus anterior free flap, gracilis muscle free flap, etc (5, 9-19). It causes secondary injury to sick patients and is difficult to complete second repair when the autogenous flaps necrosed.

With the development of tissue engineering, a substantial number of biological materials were used in organ or tissue reconstruction including artificial and native biomaterials. Common artificial biomaterials included collagen, fibrin, silk, alginate and so on. However, these materials are lack of the structural, chemical, biological, and mechanical complexity of native ECM (20). The simple native biomaterials such as allogeneic acellular dermal matrix patch are only used to repair a small range of defects (21-23). In our experiment, the complex full-thickness ECM with vascular pedicle scaffold were prepared to solve the large scale of hypopharyngeal defects.

Decellularization techniques basically preserves all the characteristics of natural ECM and decellularized scaffold can provide a good micro-ecological environment for the growth, proliferation and differentiation of cells (24, 25). Ott et al. prepared a acellular, natural, three-dimensional (3D) biological cardiac scaffold retaining the natural structure by perfusion and decellularization, and successfully simulated cardiac tissue with a certain contraction function (26). They also decellularized rat, porcine, and human kidneys by perfusion, yielding acellular scaffolds with vascular, cortical and medullary architecture, collecting system and ureters and transplanted in orthotopic position in rat successfully (27). Moreover, it was reported that decellularized scaffolds for liver, pancreas, lung, heart, and mitral valve were prepared by decellularization techniques (28-36). The antegrade or retrograde perfusion can completely retain the vascular bed (26, 37, 38). Zhang, Jian, et al prepared decellularized rectus abdominis scaffold by decellularization techniques for repairing the defect of abdominal wall and the decellularized scaffold showed good neovascularization and functional regeneration ability (38). Previously we have prepared decellularized laryngeal scaffolds with preserved cartilage by the same method and induced myocytes successfully (39, 40). Repairing hypopharyngeal defects with vascularized flaps can reduce the incidence of postoperative pharyngocutaneous fistula, infection and other complications(41). However, there is no report using decellularized hypopharynx with vascular pedicle to solve the defect after hypopharyngectomy.

Therefore, a complex full-thickness native decellularized hypopharynx with vascular pedicle scaffold was prepared by perfusing the ascending pharyngeal artery with 1%SDS and triton X-100 in our experiment. And the results indicated that the composite scaffold retained intact native anatomical ECM structure and component by HE staining and SEM and low immunogenicity by immunofluorescence. In addition, we examined the regenerative myoblasts growing in the ECM by immunohistochemical analysis. We aimed to prepared a new alternative material for hypopharynx reconstruction.

2. Materials and methods

2.1 Animals

All rabbits used in the present study were obtained from the Chengdu Dashuo Biotechnology Co. LTD (Chengdu, China) and the research complied with all committee regulations approved by the Ethics Committee for Animal Experimentation of Clinical Medical College and The First Affiliated Hospital of Chengdu Medical College. Adult New Zealand White Rabbits (n = 36),
weighing 2.5-3 kg, were used as donor animals and randomly divided into 2 groups (n = 12 each): (1) the perfusion group (decellularized hypopharynx with vascular pedicle), and (2) the control groups (fresh hypopharynx with vascular pedicle with no treatment). Two scaffolds were extracted per rabbit. The scaffolds (n = 3/time/group) were observed by macroscopic visualization, histological examination, SEM, molecular analysis, and immunological assessment. The remaining animals (n = 3/time/group) were implanted in the greater omenta of recipient rabbits who were anesthetized by intraperitoneal sodium pentobarbital administration (40 mg/kg). The implantation procedure and tests are described in detail below.

2.2 Purfuson decellularizion of hypopharynx with vascular pedicle
The adult New Zealand white rabbits were euthanized by intraperitoneal sodium pentobarbital administration (100 mg/kg). After systematic heparinization through the ear vein, a median incision of the neck exposed hypopharynx and trachea fully. The two common carotid arteries were dissected. Their superior and inferior extremities, and other branches were ligated, but their branches (superior thyroid arteries, superior laryngeal arteries and ascending pharyngeal artery) were preserved. One transfusion needle was inserted into the left common carotid arteries. The hypopharynx and common carotid arteries were removed and put on a super-clean bench. The perfusion sequence was in accordance with the Ott’s method [26]. The hypopharynx was infused at a speed of 99 ml/h by a micropump (Guangxi Veryark Technology Co., Ltd., Guangxi, China) under aseptic conditions. Briefly, heparinized phosphate buffered saline (PBS, Sigma) containing adenosine was perfused into the left carotid for 15 min and then 1% sodium dodecyl sulfate (SDS; Sigma) in deionized water was perfused for 16 h. This was followed by perfusion with deionized water for 15 min and 1% Triton X-100 (Sigma) in deionized water for 30 min. In the end, the hypopharynx was perfused with PBS containing 100 U/ml of penicillin-G (Sigma) and 100 U/ml of streptomycin (Sigma) for 48 h. During the purfusion, we maintained the entire hypopharynx in RPMI-1640 culture media without serum in an aseptic culture flask.

2.3 Characterization of decellularized scaffolds

2.3.1 Histological analysis
The samples were fixed in 10% formalin in PBS for 24 h at room temperature and then washed in distilled water, dehydrated in graded alcohol, embedded in paraffin, and sectioned. The sections were stained with hematoxylin and eosin (HE).

2.3.2 Scanning Electron Microscope (SEM)
The decellularized scaffolds were fixed with 2.5% glutaraldehyde in PBS for 24 h at 4°C. The samples were washed in PBS, dehydrated in graded alcohol, critical point-dried, and coated with gold. The samples were observed via SEM (Hitachi S-2500 Scanning Electron Microscope) to analyse the ultrastructure of the surface of pharyngeal constrictor and vessel.

2.3.3 PCR determination of DNA content
Three types of tissue samples were examined: fresh hypopharynx, decellularized hypopharynx, and decellularized vessel (about 25 mg, n = 3 for each sample). DNA was isolated with a commercially available extraction kit (Tiangen, Beijing, China). The total DNA content was measured by absorption at 260 nm on a UV eVIS spectrophotometer (Biorad, US). The total DNA was used as a template for polymerase chain reaction (PCR) analysis with primers specific for β-actin(F: 5’-CGAGATCGTGGGACAT-3’ R: 5’-CAGGAAGGAGGCTGGGAAC-3’). PCR products were separated by electrophoresis in 1.5% agarose gels.

2.3.4 Immunohistochemical analysis
The decellularized and fresh scaffold were fixed with 4% paraformaldehyde in 0.1 M PBS containing 30% sucrose at 4 ℃. Then the samples were cut to 10 mm sections on a freezing microtome (Leica), mounted on poly-L-lysine-coated slides, and washed in 0.1 M sodium PBS with 0.25% Triton X-100 (PBS/Triton, pH 7.2). Subsequently, the pieces were incubated with rabbit anti-MHC class II Polyclonal antibody (Bioss, Beijing, China) (1: 100) at 4℃ for overnight. The pieces were then incubated with CoraLite488-Conjugated Affinipure goat anti-rabbit IgG (H+L) (Bioss, Beijing, China) (1: 300) for 30 min and washed twice in PBS. Finally, the samples were stained with DAPI for 3 min and washed with 0.1 M PBS. All pieces were observed under a fluorescence microscope (Olympus).

2.3.5 Cytotoxicity of decellularized hypopharynx with vascular pedicle

5-mm diameter decellularized composite scaffold samples were cultured in L-DMEM supplemented with 15% fetal bovine serum, 100 U/ml penicillin, 100 U/ml streptomycin for 1 day and 2 days. Bone marrow mesenchymal stem cells (BMSCs) were cultured in 96-well plate (Sigma). The supernatant was removed and replaced with the medial soaked with decellularized composite scaffold (the experimental group, n = 5) for culturing 1, 2, 3 days. Correspondingly, the control groups were cultured in the normal medial without decellularized hypopharynx with vascular pedicle for culturing 1, 2, 3 days. 10μL CCK-8 reagent (Sigma) were added to culture for 4h. The OD value at 450 nm was determined by a microplate absorbance reader (HBS-1096C Pro, Nanjing, China) to determine the cell viability in each well.

2.4 Surgical transfer to the greater omentum

The abdominal skin of the anesthetized rabbits was sterilized. A midline incision in the abdominal all exposed the greater omentum. The fresh samples and decellularized samples were explanted and placed on the greater omentum. Then the greater omentum was sutured as a capsule with 6-0 nylon silk sutures and abdominal was anastomosed. All animals were returned to their cages with no immunosuppressive agents administered and treated with penicillin for 3 days. Transplanted scaffolds were acquired from the greater omentum of receptor rabbits after 3, 8 and 12 weeks. For histological analysis, the sections were stained with HE and the operation referenced to the process above.

2.5 Recellularization of decellularized hypopharynx

2.5.1 Isolation, culture and myogenic differentiation of BMSCs

Twelve receptor rabbits (4-week-old rabbits) were were euthanized by intraperitoneal sodium pentobarbital administration (100 mg/kg). Three of them were used for immunocytochemistry and the remaining rabbits for recellularization. The legs were soaked in 70% ethanol for 5 min. Then bone marrow was aspirated from the femurs in a sterile fashion. Bone marrow cells mixed with heparin were suspended and washed in PBS. Mononuclear cells were isolated by density gradient centrifugation (Histopaque 1.077 g/ml; Sigma). Cells were washed and finally resuspended in L-DMEM (1500 rpm, 10 min; Sigma) and seeded at a density of 10^6 cells/ml in L-DMEM supplemented with 15% fetal bovine serum (FBS, Hyclone, USA), 100 U/ml penicillin, 100 U/ml streptomycin, and 2 mol/L L-glutamine (Sigma) in 100 cm² petri dish. Cultures were maintained in a humidified atmosphere with 5% carbon dioxide at 37 ℃. The media was changed every 3 days. At 80% confluence, primary BMSCs were washed twice in Sterile Saline Solution and cultured with 5-azacytidine (10 mmol/L; Sigma) for 24 h for differentiation into myocytes. Cells were digested with 0.25% trypsin-EDTA (Sigma) and passaged at the ratio of 1: 3. Cells between passage 3 and passage 5 were used. Cells were examined by immunocytochemistry with antibody
to α-sarcomeric actin. The receptor rabbits were provided with penicillin for 1 week and kept in a sterile environment postoperatively.

2.5.2 Recellularized scaffolds

The decellularized hypopharynx scaffold was placed in L-DMEM with 15% FBS after perfusion. Cells were digested and cells (80-100×10^6) were suspended in 1 ml L-DMEM with 15% FBS. They were injected multipointly in the decellularized matrix with a 20ul microinjection needle. 30 min later, cellular numbers were examined and quantified in the effluent from the decellularized matrix after injection. This recellularized scaffolds was immersed in L-DMEM with 15% FBS in a humidified atmosphere with 5% carbon dioxide at 37°C for 24 h.

2.5.3 Histological analysis of recellularized scaffolds in vitro

For histological analysis, SEM was used to examine the cells adhered to the decellularized hypopharynx scaffold and the operation referenced to the process above.

2.5.4 Surgical transfer to the greater omentum

The operation of recellularized hypopharyngeal explantation referenced to the process above and the recellularized scaffold was implanted on the greater omentum for 1 week.

2.5.5 Histology and immunohistochemistry of recellularized scaffolds in vivo

Histological examination was used to examine the cells on the scaffold. For histological analysis, the sections were stained with HE and the operation referenced to the process above. Immunohistochemistry was used to examine the signs of myogenic differentiation on the decellularized scaffolds. For immunohistochemistry analysis, sections were dewaxed and rehydrated, antigen repair, perforated and incubated with a series of blocking solution. Following by, sections were exposed to the primary antibody to α-sarcomeric actin (1: 200 dilution, mouse monoclonal antibody, skeletal muscle Ab-2; Thermo, USA) for 1 h at 37°C. Sections were washed in PBS and then exposed to the secondary antibody (1: 1000 dilution, goat antimouse IgG; China). Subsequently, sections were incubated with avidin/ horseradish peroxidase (HRP) for 30 min at 37°C. Subsequently, the sections were stained with chromogenic substrate diaminobenzidine (DAB) for 5 minutes at room temperature. Sections were then counterstained with hematoxylin.

3. Results

3.1 Characterization of the decellularized scaffold

The images of the decellularized tissue indicated that it is efficient to remove the majority of the cellular element by perfusing detergent. After 16 hours of 1% SDS and triton X-100 perfusion, the decellularized scaffolds (Fig. 1B) were transparent and intact and there was no significant change in size compared to the fresh scaffolds macroscopically (Fig. 1A). HE staining indicated that there was only extracellular matrix and no cellular components in the decellularized hypopharynx (Fig. 1D) and ascending pharyngeal artery (Fig. 1F), while cellular components were found in the fresh hypopharynx (Fig. 1C) and ascending pharyngeal artery under light microscope (Fig. 1E).

SEM indicated that all cellular components were removed, but the ECMs were preserved. Collagen bundles can be observed on the surface of decellularized hypopharynx (Fig. 2B) and ascending pharyngeal artery scaffold (Fig. 2D), while collagen bundles and ECM cannot be seen in the native hypopharynx (Fig. 2A) and ascending pharyngeal artery with cellular components (Fig. 2C).

The average DNA content of the native hypopharynx and ascending pharyngeal artery
samples was 1650 ng/mg dry weight versus 50 ng/mmg of decellularized hypopharynx dry weight and 42 ng/mg of decellularized vessel dry weight. The average DNA content of the decellularized hypopharynx and vessel samples were less than that in the fresh hypopharynx with vessel samples (P < 0.05; Fig. 3). DNA analysis showed that the housekeeping gene (β-actin) in the decellularized hypopharynx and ascending pharyngeal artery sample were not amplified.

3.2 Immunogenicity of the decellularized scaffold

Immunofluorescence staining indicated that the presence of the markers of MHC-II in fresh tissue can be observed (fluorescence on hypopharynx and ascending pharyngeal artery) (Fig. 4A). However, the markers of MHC-II were not present on the decellularized scaffold. Meanwhile, DAPI staining indicated that there were a mass of nuclei in the fresh hypopharynx and ascending pharyngeal artery. But no nuclei were found in the decellularized hypopharynx and ascending pharyngeal artery. The cells were eliminated completely by perfusion.

All the implanted fresh hypopharynx samples showed strong immunological rejection (Fig. 4B). The native scaffolds and decellularized samples were encapsulated in the greater omentum. HE staining indicated that there were extensive acute inflammatory cells infiltrating in the fresh samples, while slight acute inflammatory cells infiltrating in the decellularized samples implanted in the greater omentum after 3 weeks. Some lymphocytes and necrotic tissue were observed in the fresh samples after 8 weeks. However, fewer lymphocytes and neovascularization were observed in the decellularized samples after 8 weeks. When the samples were implanted after 12 weeks, the decellularized samples degraded completely and greater omentum became thicker and the fresh samples were not discovered and purulent secretion were observed.

3.3 Cytotoxicity of decellularized hypopharynx with vascular pedicle

CCK8 assay was examined the cytotoxicity of decellularized scaffold. As examined by the CCK8 assay, no significant differences were observed between the experimental and control groups for extracts of decellularized scaffold in terms of the proliferation of rabbit BMSCs (Figure 5) (n = 6, P > .05).

3.4 Characterization of BMSCs and BMSCs induced by 5-azacytidine

BMSCs had adhered to the bottom of the petri dish and the shape of cells was fusiform after 3 days of cell culture (Figure. 6A). After 3 weeks of induced culturing by 5-azacytidine, cells still kept their fusiform shape, while the cell morphology were enlarged (Figure. 6B). Immunocytochemistry indicated that the expression of α-sarcomeric was positive in the cells induced by 5-azacytidine (Figure. 6C), while the expression of α-sarcomeric was negative in the cells without 5-azacytidine (Figure. 6D).

3.5 Characterization of recellularized scaffolds

SEM indicated that cells were polygonal in shape and adhered to the surface of recellularized scaffolds in vivo (Fig. 7A). HE staining showed cycloidal and oversized cells like regenerated myocytes grewed in the recellularized scaffold after implanted on the greater omentum for 1 week (Fig. 7B). And immunocytochemistry confirmed that the BMSCs differentiated into myocytes for positive expression of α-sarcomeric in the recellularized scaffold (Fig. 7C).

4. Discussion

Hypopharyngeal squamous cell carcinoma (HSCC) is a common malignant tumor in head and neck and the main therapy was operation, companied by chemoradiotherapy (CRT). The range
of hypopharyngotomy is too extensive to close pharyngeal defect primarily. The autogenous flaps and artificial biomaterials have been considered as an alternative materials nowadays to repair the hypopharyngeal defects. However, the native hypopharynx tissue structure contained mucosal layer, submucosa, muscle layer and serosal layer. The autogenous flaps and artificial biomaterials cannot imitate the native anatomicals tructure of hypopharynx. Meanwhile, the autogenous flaps to reconstruction will cause second injury to sick and frail patient and artificial biomaterials are lack of vascellum nourishing the tissue. We aimed to prepare an acellular, natural, three-dimensional (3D) biological hypopharynx with vascular pedicle scaffold by perfusion techniques to reconstruct laryngopharynx.

ECM is composed of around 300 different proteins and carbohydrates, which is mainly composed of collagens, proteoglycans, and glycoproteins (42-44). So ECM serves as a building support for cell growth in tissue and affect cell migration, lineage commitment, cellular behavior during disease progression, stem cell proliferation and differentiation (24, 45-47). SDS and triton X-100 can eliminate cellular component and preserve ECM efficiently (48, 49). In our study, an acellular, natural, 3D biological hypopharynx with vascular pedicle was prepared by perfusing 1% SDS for 16 h and triton X-100 for 30 min. The decellularized composite scaffold was transparent macroscopically after perfusion. Histological staining and SEM showed that the hypopharyngeal native ECM structure was preserved including mucosal layer, submucosa, muscle layer and no residual cell component was found. The decellularization criteria are as follows: 1) < 50 ng of dsDNA per mg dry weight, 2) DNA fragment length <200 bp, and 3) lack of visible nuclear material in tissue sections (28). PCR and agarose electrophoresis indicated that little nucleic acid was residual. The slighter immune rejection of decellularized composite scaffold in vivo compared to the fresh samples further verified that none of cells exist in decellularized composite scaffold. CCK-8 suggested that the medial soaked with decellularized composite scaffold had no toxic effects on cell proliferation. In conclusion, the decellularized hypopharynx with vascular pedicle scaffold in our study retained the native, full-thickness, complex structure and component of ECM and microvasculature, which has the potential to repair extensive and complex defect in clinic. However, the allogeneic acellular dermal matrix patches are too simplex to repair complex defects. The artificial tissue-engineered complex scaffold like 3D print cannot imitate the native structure of ECM and lack native elaborate microvasculature nursing tissue for the method of overlay printing. Meanwhile, the artificial biomaterials are lack of native component of ECM. Moreover, the economic cost of artificial scaffold is too high to be used extensively. By contrast, the native tissue-engineered biomaterials are low-cost and easy available.

The immunogenicity of the decellularized hypopharynx with vascular pedicle scaffold was removed by infusing 1% SDS and triton X-100 and it has a great potential to be a native muscle by recellularization. The main histocompatibility complex (MHC) resulting in immunological rejection mainly present on the cellular surface, while the ECM has no immunogenicity (50). Immunofluorescence staining indicated that the presence of markers of MHC-II was negative on the decellularized scaffolds in the study. It suggested that no cells were found in the decellularized composite scaffold by detergent perfusion. And the immunological rejection of decellularized composite scaffold in the allogeneic rabbit greater omentum was weaker visibly than the native tissues. The immunocytes were fewer in the decellularized composite scaffold and greater omentum encapsulating the decellularized scaffolds compared with the fresh samples. And the
decellularized scaffolds in the greater omentum degrade completely after 3 months. It indicated that decellularized composite scaffold prepared in the experiment had little immunogenicity and favorable biodegradability. The cartilage-preserving decellularized laryngeal scaffolds by the same method had no significant immune rejection after implanted in an allogeneic rat (39). The decellularized kidney yielding acellular scaffolds with vascular, cortical and medullary architecture, collecting system and ureters was prepared by perfusion and it was transplanted orthotopically in xenogeneic rat successfully by Ott, etc (27). The regenerated kidney produced urine and parenchymal bleeding or microvascular thrombus formation were not observed in bloodperfused vasculature of explanted regenerated kidneys (27). It suggested that acellular materials with little immunogenicity could be prepared by detergent perfusion for reconstructing organs or tissues.

Marrow mesenchymal stem cells (MSCs) are pluripotent progenitor cells that possess self-renewal and multi-directional differentiation (51). MSCs can differentiate into different kinds of cells and be easily amplified in vitro (52). Recent years, some studies showed that MSCs have immunoregulatory bioactivity and the main immunological characteristics are low immunogenicity and high immunosuppressive ability (53-55). MSCs produced a large variety of paracrine to exert its immunoregulatory effects, rather than contact-dependent mediators (56). MSCs have attracted great attention for their regenerative activity and strong immunosuppressive property of MSCs over the decades. In our experiment, BMSCs adhered to and proliferated on the decellularized hypopharynx with vascular pedicle scaffold in vitro and vivo. Moreover, BMSCs are successfully induced into myocytes by cultured with 5-azacytidine in the decellularized scaffolds. It suggested that the decellularized composite scaffold we prepared may have an positive effect on inducing BMSCs into myocytes. Ott et al. perfused the aorta with SDS and triton X-100 to prepare decellularized hearts with an acellular, perfusable vascular architecture, competent acellular valves and intact chamber geometry and under physiological load and electrical stimulation, constructs could generate a degree of pump function in recellularized heart (26). Vascularized islet organs were prepared by endothelializing acellular lung matrixes and the bio-fabricate islet released physiologic insulin and reduced hyperglycemia successfully when they were i supra-physiologic arterial glucose levels in vivo and ex vivo (57).

The hypopharyngeal reconstruction after hypopharyngectomy has been a challenge all the time. The common materials have inevitable defects. Autogenous flaps will cause second injury to patients and cannot repair the defect repeatedly when the flaps necrosed. The artificial scaffolds are lack of the component and structure of native ECM. The allogeneic acellular dermal matrix patches are of no full-thickness, complex, microvascular ECM. However, in anatomical structure, the decellularized hypopharynx with vascular pedicle scaffold we prepared retained the hypopharyngeal full-thickness native structure of ECM. On a biochemical level, the decellularized composite scaffold retained the native component of ECM and little immunogenicity. In conclusion, the decellularized composite scaffold has an potential to be an alternative material to hypopharyngeal reconstruction. In future, we plan to epithelialize the mucosa and prepare the functional hypopharynx.

5. Conclusions

The defect of hypopharyngectomy of HSCC is extensive and there is not perfect patching materials at present. An acellular, natural, 3D biological decellularized hypopharynx with vascular pedicle scaffold as the substitute materials for hypopharyngeal reconstruction could be prepared
by perfusing 1% SDS and triton X-100. This composite decellularized scaffold preserved the structure and components of the natural ECM and have little immunogenicity. The decellularized composite we prepared had good prospect in hypopharyngeal reconstruction as an allogeneic material.

**Author Contributions:** Nan Hou, Die Lv: ideals, design of methodology, creation and writing—original draft preparation, Nan Hou: acquisition of the financial support for the project leading to this publication. Die Lv, Xiao Li Xu, Yan Qing Lu, Jing Zhi Li, Ying Tang: provision of study materials, specifically performing the experiments, data presentation, Rui Na Ma, Yun Zheng: verification, reproducibility of results, supervision.

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**Declarations**

**Ethics approval and consent to participate:** This study was approved by Ethics Committee of Chengdu Medical College.

**Consent for publication:** Not applicable.

**Conflicts of Interest:** The author declares no conflict of interest.

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Figures

Figure 1

Please See image above for figure legend.

Fig. 2 Scanning electron micrographs (SEM) showed the fresh hypopharynx (A), fresh ascending pharyngeal artery (C), decellularized hypopharynx (B) and decellularized ascending pharyngeal artery (D). The collagen bundles was observed in the decellularized tissue (B, D).

Figure 2

Please See image above for figure legend.
Fig. 3 PCR analysis of β-actin in extracts of the decellularized hypopharynx with vascular pedicle. Gel electrophoresis of PCR products for β-actin in ECM of the decellularized hypopharynx, decellularized ascending pharyngeal artery, and native hypopharynx with vascular pedicle (A). Bands represented respectively the residual of DNA content in ECM of decellularized hypopharynx, decellularized ascending pharyngeal artery, and native hypopharynx with vascular pedicle (B).

Figure 3

Please See image above for figure legend.
Fig. 4 Immunogenicity testing. Immunofluorescent staining showing that the presence of MHC-II antigens in the fresh tissue, low immunoreactivity in the decellularized tissue, and DAPI staining indicated that there were a mass of nuclei in the fresh scaffold, none of nuclei in the decellularized scaffold (A). Characterization of the implanted scaffolds in vivo (B). HE staining showed lots of immunocytes infiltrating in the fresh samples, and few immunocytes infiltrating in the decellularized samples after 3 weeks. The structure of native hypopharynx was destroyed and some necrotic tissue (☆) were observed after 8 weeks. However, the structure of ECM in the decellularized hypopharynx was unbroken and neovascularization (→) were observed. Thicker greater omentum was seen compared with the native in the fresh samples and purulent secretion in the fresh samples after 12 weeks.

Figure 4

Please See image above for figure legend.
Fig. 5 Cytotoxicity assay. CCK8 assay for cytotoxicity examination of the scaffolds: the extract of ECM in decellularized hypopharynx with vascular pedicle scaffold (the medial soaked with decellularized composite scaffold for culturing 1, 2 days) and well plate alone with OD values, \( n = 5, \ P > 0.05 \). The cells were cultured with the extract for 24, 48, and 72 h.

Figure 5

Please See image above for figure legend.
Fig. 6 Characterization of cells. BMSCs were cultured for 3 days (A). Cells were induced by 5-azacytidine for 3 weeks (B). Immunocytochemistry showed the positive expression of α-sarcomeric in the cells induced by 5-azacytidine for 3 weeks (C) and the negative expression in the cells without 5-azacytidine for 3 weeks (D).

Figure 6

Please See image above for figure legend.