Advances in the Pathogenesis of Oral Submucosal Fibrosis and Animal Models

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Luo et al.: Pathogenesis of Oral Submucosal Fibrosis

Oral submucosal fibrosis is a common precancerous state, which is characterized by abnormal collagen deposition, epithelial atrophy and microvascular disease. There are many factors that may contribute to the development of the pathogenesis, including environmental factors, genetic predisposition, and immune system abnormalities. About 7%-13% of oral submucosal fibrosis cases can transform into malignant tumors. Therefore, how to establish an animal model with the same pathological mechanism as human oral submucosal fibrosis is an urgent problem to be solved. For animal models, researchers have successfully established various animal models of oral submucosal fibrosis by transplanting human oral mucosal tissue or inducing the condition with specific drugs. These models provide a powerful tool for studying the pathogenesis and treatment of oral submucosal fibrosis. Consequently, this paper summarized the pathogenesis of oral submucosal fibrosis and six methods of constructing the oral submucosal fibrosis animal models, so as to identify new drug targets for the development of anti-oral submucosal fibrosis drugs, lay a theoretical foundation and provide new treatment thinking for clinical work.

Key words: Oral submucosal fibrosis, pathogeny, arecoline, animal models, collagen

Oral Submucous Fibrosis (OSF) was first described by Schwartz in 1952, and the term was first proposed by Jens J. Pindborg and Satyavati M. Sirsat in 1966, which is still in use today1,2. OSF is a chronic, progressive and scarred precancerous state, which is characterized by excessive and abnormal collagen deposition in laminae propria layer and submucous layer, leading to tissue fibrosis, glassy degeneration and muscular degeneration2-6.

OSF was originally restricted in India, but it has not spread to Asian populations in America, England, China and Southeast Asia, which has gradually become a severe health issue in the world7-11. According to the statistics of the World Health Organization (WHO), there are over 5 million patients with OSF worldwide4,11,12. For the time being, animal model with identical pathogenesis to human OSF has not been built, and how to completely mimic an animal model similar to human OSF is an urgent problem which needs to be solved. Therefore, this paper summarized the etiology, pathogenesis and animal model construction methods of OSF, with an aim to provide new drug targets for the development of anti-OSF drugs, and to provide theoretical foundation and a new treatment thinking for clinical practice.

ETIOLOGY OF OSF

The etiology of OSF remains unclear, and it is currently considered that it is an oral mucosal disease might be caused by multiple factors. As suggested by epidemiologic studies, betel nut chewing is one of the most important risk factors for inducing OSF13-15. Betel nut contains a large amount of arecoline, which plays an important role in the pathogenesis of OSF as mentioned in current study16-18. It is demonstrated in some other studies that the risk of OSF in the betel nut chewing population increases by 109-287 folds compared with the non-chewing population, besides, the betel nut chewing frequency is positively correlated with the duration19,20.

Other risk factors include smoking, eating spicy...
foods, vitamins B and C, iron deficiency, gene mutations, autoimmunity and human papilloma virus infection\textsuperscript{[4,20-22]}. The morbidity of OSF varies depending on race and region, and it is also closely correlated with factors such as diet, habit and culture\textsuperscript{[23-25]}. As reported in some study, betel nut chewing, smoking, drinking and other habits will increase the risk of OSF\textsuperscript{[20,26,27]}.

**PATHOGENESIS OF OSF**

The pathogenesis of OSF is still unclear, which may be related to factors including altered collagen homeostasis, hypoxia, together with increased production of inflammatory cytokines and growth factors\textsuperscript{[28-32]} (fig. 1).

**Altered collagen homeostasis:**

At the moment, a majority of studies have indicated that, OSF is the consequence of altered collagen homeostasis, namely, the reduced collagen clearance and the increased collagen synthesis\textsuperscript{[19,30]}.

**Decreased collagen clearance:**

Collagen stabilization, defective Extracellular Matrix (ECM) dynamics and suppression of collagen phagocytosis will reduce the collagenase activity, which subsequently decreases collagen degradation, and finally leads to the decreased collagen clearance rate\textsuperscript{[28,30]}.

Arecoline promotes cross-linking between collagen peptide chains, thereby endowing collagen the ability to resist collagenase-mediated degradation. Besides, arecoline dose-dependently up-regulates cystatin C in buccal mucosal fibroblasts\textsuperscript{[30,33-35]}. The up-regulation of cystatin C in turn inhibits the cysteine protease, thus decreasing collagen degradation and further stabilizing collagen in OSF\textsuperscript{[30]}. Flavonoids such as tannin and catechin are other important components in betel nut, which exert synergistic effects with alkaloids to decrease collagen degradation by suppressing collagenase and stabilizing collagenous fiber, finally inducing the occurrence of OSF\textsuperscript{[36,37]}.

Fibroblast phagocytosis exerts a critical function in regulating ECM remodeling through collagen degradation\textsuperscript{[30]}. However, fibroblasts derived from OSF patients lack the effective collagen phagocytic function, which may be related to the ECM deposition and eventually fibrosis\textsuperscript{[13,38]}.

**Increased collagen synthesis:**

Betel nut-induced local mucosal inflammation will recruit activated T cells and macrophages, and increase the levels of cytokines and TGF-\(\beta\)\textsuperscript{[44,45]}. TGF-\(\beta\) can activate the procollagen genes Collagen Alpha (\(\alpha\)) 2(I) (COL1A2), COL3A1, COL6A1, COL6A3 and COL7A1 chain to remarkably increase collagen production\textsuperscript{[30,46,47]}. At the same time, it promotes the production of procollagenase and up-regulates the secretion of Lysyl Oxidase (LOX) that initiates collagenous fiber cross-linking\textsuperscript{[48]}.

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**Fig. 1: Schematic diagram of the pathogenesis hypothesis of OSF**
The active components in betel nut can induce the massive synthesis of collagen in oral mucosal cells. The alkaloids in betel nut will stimulate fibroblasts to synthesize collagen by activating fibroblasts and increasing collagen production. Copper further increases collagen production and oral fibroblast cross-linking. Meanwhile, catechin and flavonoid induce the occurrence of fibrosis through accelerating collagen production and cross-linking.

Further, the addition of slaked lime (calcium hydroxide) into betel nut will not only create a kind of alkaline environment, promote the oxidation of polyphenols, and increase the release of Reactive Oxygen Species (ROS), but it also accelerates the hydrolysis of betel nut into arecoline, which can induce change in fibroblast phenotype, thereby increasing the formation of collagen with altered molecular structure.

**Hypoxia:**

Hypoxia-Inducible Factor-1 (HIF-1) is a kind of nuclear transcription factor existing in hypoxic cell nuclei, which can bind to Deoxy Ribonucleic Acid (DNA) and activates the expression of numerous hypoxia genes in the hypoxic environment. In the early stage of OSF, the high expression of HIF-1 induces the transcription of downstream cytokines like TGF-β, promotes fibroblast proliferation and collagen synthesis, suppresses collagen degradation, and ultimately results in OSF.

**Increased production of inflammatory cytokines and growth factors:**

When chewing betel nut, the crude fiber will damage the oral mucosa, then induce inflammatory response in epidermal cells and activate the macrophages to secrete cytokines. TGF-β is the major cytokine participating in OSF progression, which regulates the expression of type I collagen and α-Smooth Muscle Actin (SMA) in myofibroblasts.

As discovered in a study, in vitro arecoline up-regulates the inflammatory cytokines and growth factors such as Interleukin-1 (IL-1), IL-6, IL-8, TGF-β, tumor necrosis factor-α, fibroblast cytokines, and platelet-derived growth factor, and down-regulates the interferon-Gamma (γ) level to promote collagen synthesis, leading to tissue fibrosis. Moreover, arecoline can stimulate the biosynthesis of Connective Tissue Growth Factor (CTGF) in oral mucosal fibrosis through the ROS, nuclear factor-kappa B, Jun N-terminal Kinase (JNK) and p38 Mitogen-Activated Protein Kinases (p38 MAPK) pathways in a dose- and time-dependent manner.

Changes in cytokines and growth factors will result in fibroblast proliferation and collagen synthesis near the injury site, thereby causing OSF.

**PREPARATION OF OSF ANIMAL MODEL**

There are numerous methods to prepare OSF animal model, among them, six methods together with their merits and demerits have been elaborated below.

**Application of bleomycin in preparing the OSF rat model:**

Bleomycin is one of the most extensively applied drugs in the preparation of fibrosis model. Some scholars have injected bleomycin into the buccal mucosa of Sprague-Dawley (SD) rats to prepare the OSF animal model. Gao et al. selected 20 Specific-Pathogen-Free (SPF) grade adult male SD rats weighing (220±20) g in their experiment. After isoflurane anesthesia, 100 μl of the 1 mg/ml bleomycin dilution was injected into bilateral buccal mucosae, respectively, once a day for 8 d consecutively. The results suggested that, after injection of bleomycin for 8 w, the OSF model was successfully prepared.

Zhang et al. prepared bleomycin at 0.1 mg/ml, 0.5 mg/ml, 1 mg/ml and 2 mg/ml, injected 100 μl of bleomycin at the above 4 concentrations into bilateral buccal mucosae of rats every day for 8 d consecutively, and the rat model of OSF was prepared.

This method can successfully mimic the clinical and pathological manifestations of human OSF at the animal level, and has been extensively applied, with the merits of rapid preparation and good reproducibility. However, it is still associated with certain limitations; for instance, due to the different pathological process from betel nut chewing, pathological changes show atypical hyperplasia and pulmonary fibrosis, and repeated injections will cause unwanted inflammation, edema and congestion.

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Construction of OSF mouse model based on the arecoline drinking water method:

As indicated in some research, arecoline is a key factor inducing OSF, and some scholars have adopted the arecoline drinking water method to construct the OSF mouse model. Wen et al.\textsuperscript{[69]} utilized 27 SPF grade male Balb/c mice and fed them with 500 mg/l arecoline dissolved in water for 12 w. According to their observations, early OSF-like lesions were observed on the buccal mucosa. Wen et al.\textsuperscript{[70]} added a high concentration of arecoline (1000 mg/l) in the drinking water, and early OSF-like changes occurred in lingual mucosa in the 8\textsuperscript{th} w, while OSF-like changes were also observed in oral and buccal mucosae in the 12\textsuperscript{th} w, and transparent changes were seen in the lingual and palatal mucosae in the 20\textsuperscript{th} w, indicating the advanced OSF.

In this method, mice have a free access to the drinking water, the induction method is simple and reproducible after excluding the interference of foreign substances and age, the modeling time is shortened, and the drug effect on the entire oral mucosa is controlled through single dose administration. However, this method greatly affects the whole body, and is slightly different from the local human OSF process induced by betel nut chewing.

Construction of the OSF mouse model through injection of betel nut extract:

It is suggested by numerous studies \textit{in vitro} that, Areca Nut Extract (ANE) can increase collagen formation through the trans-differentiation of myofibroblasts that express the intracellular marker protein α-SMA, and regulate MMP activity to further decrease collagen degradation. Moreover, ANE can induce fibrosis in different cell lines via signaling molecules such as TGF-β, CTGF, IL-6 and prostaglandin E2\textsuperscript{[71,72]}. To verify the inhibitory effect of photobiomodulation therapy on mouse OSF, Chiang et al.\textsuperscript{[72]} injected ANE into the buccal mucosa and induced fibrosis within 1 mo. Besides, they observed the OSF-like pathological features. Shekatkar et al.\textsuperscript{[73]} injected 50 μl ANE (50 mg/ml) into the right buccal mucosa in Swiss albino mice at 2 d intervals within 12 w and induced OSF.

This model is stable and rapid, but repeated injections will induce unnecessary inflammation, edema and congestion, and interfere with the common OSF formation pathways. In addition, the pathological process is slightly different from betel nut chewing, and the composition of ANE remains unclear.

Preparation of OSF mouse model through applying betel nut extract:

The OSF mouse model is prepared through applying ANE onto the buccal mucosa. Perera \textit{et al.}\textsuperscript{[74]} randomly selected 20 adult male albino Balb/c mice with the age of 10-12 w (weight, 28-30 g), applied ANE onto the buccal mucosal surface for twice a day, 6 d a w for 300-600 consecutive d, and successfully prepared the OSF model. Yang \textit{et al.}\textsuperscript{[75]} applied ANE at four concentrations (0, 0.5 mg/ml, 2 mg/ml, and 8 mg/ml) in buccal mucosa of SD rats for 16 w, and the SD rat model of OSF was successfully constructed under the stimulation of 2 mg/ml and 8 mg/ml ANE. However, the model was not constructed by stimulation of 0 and 0.5 mg/ml ANE, revealing that this method requires a certain concentration of ANE when constructing the OSF model.

Compared with other modeling methods, this method can mimic the human OSF pathological process, but it is time-consuming, almost the whole life cycle of mice is needed to induce OSF, and the ANE composition remains unclear. Consequently, numerous confounding factors (especially aging) may affect the pathological process of OSF in the model.

Construction of the OSF mouse model through drip of betel nut extract:

The OSF mouse model is prepared through intravenous drip of ANE in the entire oral cavity. Shekatkar \textit{et al.}\textsuperscript{[73]} given drip of 50 μl ANE (50 mg/ml) into the whole oral cavity in the Swiss albino mice every day, mice were not allowed to drink water within 3 h after administration, and the OSF model was constructed after 12 w. Meanwhile, as revealed by the study by Shekatkar \textit{et al.} the method of oral drip of ANE was superior to oral local injection of ANE. This method is easy in operation, brings fewer discomforts to the mice, and mimics the natural development of the disease, but the ANE composition remains to be further determined.

Construction of the OSF rabbit model through injection of phenol solution:

The rabbit OSF model is constructed through
injection of phenol solution into the rabbit buccal mucosa. Lin et al. injected 4% phenol solution in the buccal mucosa twice a week, and they successfully induced the OSF-like lesions in rabbits 4 w later. Lin et al. used the model to determine the relation between OSF pathological changes and endogenous collagenase activity, and whether exogenous collagenase treatment improved the food intake function. This model is rarely used, and its merits and demerits remain unclear.

CONCLUSION

OSF is prevalent in Asian countries and has spread in North America and Europe. At present, the etiology of OSF may be related to factors like betel nut chewing, smoking, eating spicy foods, vitamin B, vitamin C and iron deficiency, gene mutations, autoimmunity and human papillomavirus infection. The pathogenesis of OSF is complex and still not fully understood. The pathogenesis of OSF may be associated with factors such as altered collagen homeostasis, hypoxia, together with increased inflammatory cytokine and growth factor production. For the time being, there are mainly six methods for constructing the OSF animal models, which have their own advantages and disadvantages. The premise of constructing an animal model with identical pathogenic mechanism to human OSF is to determine the etiology and pathogenesis of OSF. Consequently, more studies are needed to further investigate the etiology and pathogenesis of OSF, so as to provide theoretical foundation and new treatment thinking for the prevention and treatment of OSF.

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Wen Luo, Jie Mei and Kaiyue Zheng contributed equally to this work and shared the first authorship. Xi Xie and Hai Bin Luo are the corresponding authors.

Conflict of interests:

The authors declared no conflict of interest.

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