Expression of the stem cell factor in fibroblasts, endothelial cells, and macrophages in periapical tissues in human chronic periapical diseases

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ABSTRACT. Stem cell factor (SCF), an important stem cell cytokine, has multiple functions. Fibroblasts (FBs), mature mast cells, endothelial cells (ECs), and eosinophil granulocytes can produce SCF in the inflammatory process. Therefore, we aimed to observe SCF expression in FBs, ECs, and macrophages (MPs) in periapical tissues in human chronic periapical disease and investigate the effects of cells expressing SCF in pathogenesis of the disease. Healthy (N = 20), periapical cyst (N = 15), and periapical granuloma (N = 15) tissues were fixed in 10% formalin for 48 h, embedded in paraffin, and stained with hematoxylin and eosin to observe histological changes. SCF expression was observed in FBs, ECs, and MPs in periapical tissues by double immunofluorescence. CD334, CD31, and CD14 are specific markers of FBs, ECs, and MPs, respectively. Results showed that densities of CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs were significantly increased in periapical tissue groups (P < 0.01). There were no significant differences in CD334-SCF double-positive FB and CD31-SCF double-positive EC levels between the two periapical tissue groups (P > 0.05). CD14-
SCF double-positive MP density was considerably higher in periapical granulomas than in cysts (P < 0.01). FB, EC, and MP levels were significantly high and densities of CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs improved considerably in chronic periapical tissues, suggesting that the cells might be related to occurrence, development, and pathogenesis of chronic periapical disease.

Key words: Chronic periapical disease; Stem cell factor; Fibroblasts; Endothelial cells; Macrophages

INTRODUCTION

Stem cell factor (SCF), also known as steel factor or c-Kit ligand, has two protein forms, namely, membrane-bound stem cell factor and soluble stem cell factor, which mainly act on proliferation, migration, survival, and differentiation of hematopoietic progenitor cells, melanocytes, and generative cells (Lennartsson and Rönnstrand, 2012). In the inflammatory process, fibroblasts (FBs) and endothelial cells (ECs) can produce SCF. Since mature mast cells (MCs) and eosinophil granulocytes have tropism for SCF, they can adhere to fibronectin or ECs and exhibit degranulation (Pradier et al., 2014). In this study, double immunofluorescence (DIF) was used to observe the expression of CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs in human chronic periapical disease tissues and investigate their involvement in the disease mechanism.

MATERIAL AND METHODS

Subjects

Study subjects were recruited from the Affiliated Stomatological Hospital of Jinan University between July 2013 to July 2015. The patient group consisted of 15 patients with periapical cysts and 15 patients with periapical granuloma (ages ranging from 24 to 65 years). The 20 control samples were obtained from healthy premolar teeth extracted from individuals (ages ranging from 15 to 23 years) for orthodontic purposes. Pregnant women or individuals with systemic diseases were excluded. Individuals who were not on antibiotics three months prior to study commencement were selected. The study met specified standards of the Institutional Review Board (IRB) and received its approval. In addition, informed written consent was obtained from all study participants.

Groups and materials

Samples were divided into the three groups: the control, periapical granuloma, and radical cyst groups. The control group consisted of 20 samples obtained from healthy premolar teeth extracted for orthodontic purposes. The periapical granuloma group consisted of 15 samples showing round or oval transmission shadows with low density in the apical area (diameter ≤1 cm) detected by X-rays imaging. Lesion tissues were pathologically examined to be granulated, accompanied with capillary proliferation and high inflammatory...
cell infiltration and without non-keratinized stratified squamous epithelium. The radicular
cyst group consisted of 15 samples showing round or oval transmission shadows with low-
density in the apical area (diameter ≤1 cm) and obvious ray resistance in their edge detected
by X-ray. Lesion tissues were pathologically examined to contain numerous collagen fibrils,
which were cysts or tissues covered with fully or partly non-keratinized stratified squamous
epithelium.

Methods

Collection and treatment of tissue samples

The periapical granuloma and radicular cyst samples were obtained from lesion
tissues during apicetomy. Control group samples were obtained by using a sterile blade and
scraping the parodontium of healthy premolar teeth extracted because of orthodontic purposes.
The periodontal tissues were fixed in 10% formalin for over 48 h, dehydrated, embedded,
sectioned into 5-μm continuous slices, and then stained with hematoxylin and eosin (HE) for
histopathology analysis by microscopy.

Detection of CD334-SCF in FBs, CD31-SCF in ECs, and CD14-SCF in MPs by DIF

The first antibodies used in this study with a dilution rate of 1:100 were as follows:
anti-SCF (Santa Cruz Biotechnology, Santa Cruz, CA, USA), CD14 antibody (Santa Cruz
Biotechnology), CD31 antibody (Santa Cruz Biotechnology), and CD334 antibody (Santa
Cruz Biotechnology). The secondary antibodies used in this study with a dilution rate of 1:200
were as follows: goat-anti-mouse IgG (H+L) Alex Flour 555® (Cell Signaling Technology,
Danvers, MA, USA) and goat-anti-rabbit IgG (H+L) Alex Flour 488® (Cell Signaling
Technology). Tissue slices were dewaxed, repaired with saline sodium citrate buffer,
blocked with goat serum, incubated using the first antibody, and washed with phosphate-
buffered saline thrice for 5 min. The slices were then incubated with the secondary antibody
in the dark, stained with 4’,6-diamidino-2-phenylindole, and immediately observed under
fluorescence microscope. CD334+, CD31+, and CD14+ showed green light signals, while
SCF+ showed a red light signal, located in the cell membrane and/or cytoplasm. The light
was orange when red and green light signals merged in the same field. Slices were blindly
observed under a fluorescence microscope by two pathologists according to the counting
method reported by Batista et al. (2005), gaining double-positive cells per square (cells/
mm²). The average value recorded by this method corresponded to the average density
of CD334+SCF double-positive FBs, CD14+SCF double-positive MPs, and CD31+SCF
double-positive ECs in slices.

Statistical analysis

Data are reported as means ± SD and were analyzed by SPSS 13.0 software. The
Kruskal-Wallis test was used to compare the average densities of CD334+SCF double-positive
FBs, CD14+SCF double-positive MPs, and CD31+SCF double-positive ECs. The Nemenyi
test was conducted to compare each cell type independently among the patient and control
groups. P < 0.05 was considered statistically significant.
RESULTS

Histological analysis

No obvious infiltration of inflammatory cells was observed in the control group (Figure 1A). However, there was excessive infiltration of inflammatory cells such as neutrophilic granulocytes, lymphocytes, and MPs, and capillary proliferation (Figure 1B) in the periapical granuloma group. In the radicular cyst group, we observed intercellular edema and excessive infiltration of inflammatory cells (mainly neutrophilic granulocytes) in the epithelial layer (Figure 1C) and collagen fibrils without infiltration of inflammatory cells in the fibrous tissue layer (Figure 1D).

Figure 1. Results from histological analysis. A. Healthy control group; B. periapical granulomas group; C. radicular cysts group; D. fibrous tissue layer.

DIF results

SCF were found to be expressed in FBs, ECs, and MPs in each group. A few CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs were observed to be scattered in the control group (Figure 2, Figure 3, and Figure 4). In the periapical granuloma group, CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, CD14-SCF double-positive MPs, and erythrocytes were dramatically increased (Figure 5, Figure 6, and Figure 7). Besides, in the radicular cyst group, CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, CD14-SCF double-positive MPs, and mature erythrocytes were increased (Figure 8, Figure 9, and Figure 10).
Figure 2. Double immunofluorescence results of fibroblasts in the healthy control group. A. CD334⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).

Figure 3. Double immunofluorescence results of endothelial cells in the healthy control group. A. CD31⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).
Figure 4. Double immunofluorescence results of macrophages in the healthy control group. 
A. CD14⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).

Figure 5. Double immunofluorescence results of fibroblasts in the periapical granuloma group. 
A. CD334⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).
Figure 6. Double immunofluorescence results of endothelial cells in the periapical granuloma group. A. CD31⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).

Figure 7. Double immunofluorescence results of macrophages in the periapical granuloma group. A. CD14⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).
Figure 8. Double immunofluorescence results of fibroblasts in the radicular cyst group. A. CD334⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).

Figure 9. Double immunofluorescence results of endothelial cells in the radicular cyst group. A. CD31⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).
Density of CD334+SCF double-positive FBs in each group

As shown in Figure 11, there were significant differences in the density of CD334-SCF double-positive FBs between the control group and periapical tissue groups (P < 0.01), while no differences were observed between the periapical granuloma and radicular cyst groups (P > 0.05).

![Figure 10](image)

**Figure 10.** Double immunofluorescence results of macrophages in the radicular cyst group. A. CD14⁺; B. SCF⁺; C. DAPI⁺; D. Merged (scale bars = 50 μm, 400X).

**Figure 11.** Density of CD334+SCF double-positive fibroblasts in each group. Mean ± SD. *P < 0.01 vs healthy control group; °P > 0.05 vs periapical granuloma group.

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Density of CD31+SCF double-positive ECs in each group

As shown in Figure 12, there were significant differences in the density of CD31-SCF double-positive ECs between the control group and periapical tissue groups (P < 0.01), while no differences were observed between the periapical granuloma and radicular cyst groups (P > 0.05).

![Figure 12](image)

**Figure 12.** Density of CD31+SCF double-positive endothelial cells in each group. Mean ± SD. *P < 0.01 vs healthy control group; †P > 0.05 vs periapical granuloma group.

Density of CD14+SCF double-positive MPs in each group

As shown in Figure 13, there were significant differences in the density of CD31-SCF double-positive ECs between the control group and periapical tissue groups (P < 0.01). The density was considerably higher in the periapical granuloma group than in the radicular cyst group (P < 0.01).

![Figure 13](image)

**Figure 13.** Density of CD14+SCF double-positive macrophages in each group. Mean ± SD. *P < 0.01 vs healthy control group; †P < 0.01 vs periapical granuloma group.
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Density of CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs in each group

The results are shown in Figure 14, and the differences have been described above.

**DISCUSSIONS**

Chronic periapical disease is characterized by the inflammation and destruction of periapical tissues. The bacteria and its virulence factor in an infectious root canal enter periapical tissues and stimulate the host inflammatory reaction and immune mechanism by apical foramina, produce various pro-inflammatory cytokines such as tumor necrosis factor α (TNF-α), interleukin-1β (IL-1β), and interleukin-6, and directly cause osteoclast formation and bone absorption (Jakovljevic et al., 2015).

In human chronic periapical disease tissues, excessive infiltration of lymphocytes (LYMs), plasmocytes, FBs, MPs, and MCs has been observed with EC swelling and capillary proliferation (Fan and Zhou, 2012). It has been proven that active MPs and LYMs are the cells responsible for periapical tissue inflammation and bone absorption at the early stage of disease (Takeichi et al., 2008). MPs are involved in phagocytosis in the inflammation process and can be promoted by microorganism lipopolysaccharides as well as chemical mediators to produce IL-1β and TNF-α, activate osteoclast, and cause periapical bone absorption and destruction (Nair, 2004). IL-1β has the ability to affect T cells and ECs in periapical tissues for an improved local immune reaction, while TNF-α can stimulate FB proliferation and promote local tissue fibrosis (Solanki et al., 2013).

A study reported that ECs play an important role in human chronic periapical disease, and various pro-inflammation cytokines could destroy the close association of ECs, improving endothelial permeability in the inflammatory reaction and immune process (Hama et al., 2008). ECs have also been found to have the ability to regulate inflammation in periapical tissues by controlling the movement of inflammatory cells and increasing blood vessel permeability (Takeichi et al., 2008).
In the inflammatory process, FBs could proliferate to form fibrotic tissues and encompass lesion areas, forming a type of fibrous coat to restrict the spread of inflammation into deep tissues (Fan and Zhou, 2012). LPS in infected pulp tissues of pulp canal could activate LYMs and FBs producing pro-inflammatory cytokines such as TNF and IL-1 (Chang et al., 2005). Some studies have suggested that FBs could be activated to secrete inflammatory cytokines, such as NLRP3 and IL-33, after stimulation of inflammation and participate in infiltration, cell transport, and tissues fibrosis in the tissue repair process (Liu and Liu, 2013; Sipert et al., 2014; Velickovic et al., 2016).

SCF, an important stem cell cytokine with multiple functions, has been extensively studied (Fang et al., 2012). Over-expression of SCF/c-Kit is associated with the inflammatory reaction, proliferation spread, and wound repair (Kalra et al., 2013). Expression of the SCF/c-Kit receptor and its ligand in epithelial tissues has been found to be associated with neoplastic transformation (Gagari et al., 2006). Therefore, SCF has a close relationship with allergic asthma, liver tissue fibrosis, small cell lung cancers, and so on (Choudhary et al., 2016). Another study reported that the SCF/c-Kit pathway could promote tissue angiogenesis and reconstruction (Kim et al., 2014). In the inflammatory process, FBs and ECs produce SCF, while mature MCs and eosinophil granulocytes express SCF by adhering to fibronectin or ECs and show degranulation (Kim et al., 2014).

Studies have shown that there is excessive infiltration of FBs, ECs, and MPs in periapical tissues of patients with chronic periapical disease. However, the mechanism of SCF-positive FBs, ECs, and MPs in the incidence, development, and progression of the disease has not been elucidated. In the current study, DIF results showed that the densities of CD334-SCF double-positive FBs, CD31-SCF double-positive ECs, and CD14-SCF double-positive MPs were all significantly higher in the chronic periapical groups than in the control group. No significant differences were observed in the densities of CD334-SCF double-positive FBs and CD31-SCF double-positive ECs in the periapical granuloma and radicular cyst groups. The density of CD14-SCF double-positive MPs was considerably higher in the periapical granuloma group than in the radicular cyst group. Our study indicated that controlling or regulating the functions of FBs, ECs, and MPs with SCF expression could be a new treatment target to control the development of human chronic periapical disease.

Conflicts of interest

The authors declare no conflict of interest.

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