Cytokine and adhesion molecule expression in the minor salivary glands of patients with Sjögren’s syndrome and chronic sialoadenitis

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Abstract

Objective—To investigate the role of cytokines and cell adhesion molecules in the pathogenesis of Sjögren’s syndrome (SS).

Methods—Using an indirect immunoperoxidase technique we assessed the expression of the cytokines interleukin-1α (IL-1α), interleukin-1β (IL-1β), interleukin-8 (IL-8), transforming growth factor β (TGFβ) and granulocyte macrophage colony stimulating factor (GM-CSF), of the adhesion molecules intercellular adhesion molecule-1 (ICAM-1), lymphocyte function associated antigen-1 (LFA-1), the activated molecular form of LFA-1 (NK1-L16), CD2, and LFA-3, and of a panel of cellular markers in the minor salivary glands.

Results—In SS and chronic sialoadenitis (CS), the ductal epithelial cells and acini expressed all the cytokines examined. The percentage of glandular mononuclear cells which stained positive for cytokines did not differ significantly between SS and CS. NKI-L16 was detected on 33.6 (SD 10.1) % and 15.3 (4.3) % of LFA-1 cells in SS and CS, respectively (p < 0.002).

Conclusion—SS and CS did not differ in the pattern of cytokines examined. The characteristic cell clustering seen in the salivary glands in SS may be caused by the upregulation of NKI-L16.


Sjögren’s syndrome (SS) is an autoimmune disease characterised by a periductal mononuclear cell (MNC) infiltrate resulting in distinct focal cellular aggregates of the salivary and lachrymal glands, leading to a chronic inflammatory process with secondary symptoms and signs of dry mouth and dry eyes and by the production of autoantibodies.1–3 The infiltrating cells are predominantly CD4 T cells, but plasma cells and macrophages are also present.4–5 Despite extensive histological, immunopathological and virological studies, the triggering factors responsible for the autoimmune lesion in the glandular tissue of SS remain obscure.

Cytokines have been implicated in the pathogenesis of many inflammatory lesions. Previous studies, applying immunohistochemical techniques to the salivary glands of patients with SS, demonstrated that interferon gamma (IFNγ) was extensively expressed in the acini, ducts, and MNCs, but only 2.4% of infiltrating cells expressed interleukin-2 (IL-2).6,7 This was further confirmed by in situ hybridisation and the reverse transcriptase polymerase chain reaction (RT-PCR),8,10 in which mRNA for IFNγ and IL-2 was detected in the salivary glands of all patients studied, while interleukin-4 (IL-4) was absent.8 These observations suggest continuing T cell activation of the Th1 cell type in the SS lesions, as reflected by the local production of IL-2 and IFNγ. Furthermore, biopsies of salivary glands from SS patients with extensive lymphoid infiltrates contained mRNA for IL-2 and IFNγ, while biopsies from few lymphoid infiltrates had large amounts of transforming growth factor β (TGFβ).10 It was suggested, therefore, that the greater concentrations of TGFβ in minimal lesions may play a role in downregulating the inflammatory process. Support for this hypothesis comes from a mouse model in which disruption of the TGFβ gene by homologous recombination in murine embryonic stem cells makes the animals unable to mount an efficient anti-inflammatory response. The salivary glands of these mice were characterised by a periductal lymphoplasmacytic infiltrate not unlike that associated with SS.11

Chronic sialoadenitis (CS) is a non-specific inflammatory process of the salivary glands characterised by a sparse diffuse MNC infiltrate in the gland leading to glandular atrophy with secondary signs and symptoms of dry mouth, but not the presence of autoantibodies. Thus this disease is the ideal control for immunopathological studies in SS. We decided, therefore, in the first instance to investigate the expression of cytokines such as TGFβ, IL-1α, IL-1β, IL-8 and granulocyte macrophage colony stimulating factor (GM-CSF) in SS and CS patients to see if these two conditions could be distinguished. We also examined salivary glands from five healthy normal control subjects. In order to clarify the typically different (focal versus diffuse) distribution and organisation of the MNC infiltrate in SS and CS, we then studied the expression of specific adhesion molecules (lymphocyte function associated antigen-1 (LFA-1)/intercellular adhesion molecule-1 (ICAM-1) and CD2/LFA-3) involved in homo- and heterotypic cluster formation. Finally, we examined the expression of an activation dependent epitope of the LFA-1
molecule recognised by the monoclonal antibody (MAb) NKI-L16.12

Patients and methods

Patients

We studied six patients with definite SS (four primary and two secondary with rheumatoid arthritis; mean age 64-6 (range 40-79 years; all female) who fulfilled the American College of Rheumatologists criteria and European criteria for SS.2,3 This group was characterised by symptoms and signs of ocular and oral involvement, and the presence of antinuclear antibodies, and SS-A and SS-B antibodies. Four of five patients tested were rheumatoid factor positive and all demonstrated a typical pericentral infiltrate on histological analysis of the minor salivary glands. We also studied six patients with CS (mean age 58-3 range 57-76 years; five females and one male) with sicca syndrome and the histopathological features of CS. This group was characterised by oral involvement (xerostomia), no signs or symptoms of lachrymal involvement, absence of autoantibodies (rheumatoid factor, SS-A and SS-B) and the presence of a lymphocytic infiltrate in the salivary glands. Five biopsies from healthy volunteers were also studied.

Tissue Preparation and Staining

Biopsies containing at least two foci per 4 mm² were obtained from minor salivary glands from the midline of the lower lip and embedded in optimal temperature cutting compound (OCT, Miles, CA) and snap frozen in isopentane and liquid nitrogen. Samples were stored at -70°C until sectioned for immunohistological staining. Sections 5 μm thick were cut with a cryostat (Leitz) at -22°C. Sequential sections were mounted on poly-l-lysine coated slides and dried overnight at room temperature. Sections were fixed in acetone for 10 minutes, wrapped in tin foil, and stored at -70°C until further use.

Cytokines and cellular antigens were determined by an indirect immunoperoxidase technique. The MAbs used to determine cellular content were OKT3, OKT4, OKT8, (anti-CD3, anti-CD4, anti-CD8, respectively; ATCC, USA), B-Ly1, and EBMT1 (anti-CD20, a B cell marker and anti-CD68, a macrophage marker, respectively; Dako laboratories, Copenhagen, Denmark). Anti-factor VIII related antigen (Dako) was used to stain endothelial cells. The MAbs used to determine cytokine expression were anti-IL-1α (gift of Dr R Thorpe, NIBSC, Potters Bar), anti-IL-1β (gift of Dr D Boraschi, Italy), anti-GM-CSF (gift of Dr K Ruedi, Sandoz, Basel, Switzerland), anti-IL-8 (gift of Dr M Ceska, Vienna) and anti-TGFβ (gift of Dr Feldman, Genentech, USA). The MAbs used to detect the adhesion molecules were anti-LFA-1 (TSI/22 hybridoma clone, gift of Dr T Springer, USA), NKI-L16 (gift of Prof Figdor, Netherland), anti-ICAM-1 (gift of Dr D Haskard, London, UK), anti-CD2 and anti-LFA3 (Becton Dickinson, CA, USA).

Briefly, the salivary gland tissue sections were incubated for 10 minutes with a 1:20 dilution of normal rabbit serum in a humidified chamber at room temperature. The sections were then incubated with the primary MAbs, at appropriate dilutions in phosphate buffer saline (PBS) for two hours. The excess MAb was removed by washing with PBS. The secondary antibody, horseradish peroxidase conjugated rabbit anti-mouse (Dako laboratories, Copenhagen, Denmark) diluted in PBS at 1:100, was added and sections were incubated for 30 minutes. After rinsing with PBS the sections were developed in a solution of diaminobenzidine tetrahydrochloride 0.7 mg/ml, and counterstained with haematoxylin. Finally, sections were dehydrated by transferring through alcohol and CNP30, and mounted in DPX.

Negative control staining was performed on all salivary gland specimens studied, following an identical procedure except that incubation with the primary MAb was omitted; no staining was noted in these sections. Tonsil tissue was used as a positive control. Other controls included MOPC21—a non-immune mouse IgG1 myeloma protein MOPC21, with an isotype similar to the cytokine MAbs used, which was used at equivalent protein concentration.

All comparisons between specimens were made with data derived from the same staining run.

Confirmation of Cytokine Immunoreactivity

The specificity of the cytokine MAbs has been extensively studied13 and demonstrated by the providers of these reagents (IL-1α,14 IL-1β,15 GM-CSF,16 IL-817 and TGFβ18). The specificity of the binding of these antibodies to salivary gland tissue was confirmed by: staining with cytokine MAb (staining was concentration dependent); using an irrelevant antibody, MOPC21, at equivalent protein concentration; comparing the pattern and degree of staining of different MAbs for the same cytokine product on salivary gland and tonsil which yielded the same pattern of staining; and verifying increased cytokine production by phytohaemagglutinin stimulated versus resting peripheral blood MNCs.

Microscopic Evaluation

All microscopic evaluations were performed in a coded fashion by one observer (AC) who was blinded to the name of the patients and diagnosis. To rule out intra-observer variation, the samples were randomised, recoded, and read on a second occasion. (No significant differences were demonstrated between the two readings.) Samples were also assessed by another investigator (GY) to estimate inter-observer variation. (No significant differences were found between the two investigators.)

Number of Infiltrating Cells

For each patient, the total number of infiltrating cells was estimated by counting all
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...the cells throughout the minor gland biopsied and expressing the results per mm².

**CELL SUBSETS**
The number of cells staining positively for a particular cell marker was counted in a total of 500 cells, and the results expressed as a percentage.

**CYTOKINE STAINING**
Cytokine expression was assessed in three areas of the salivary glands: infiltrating cells, ducts, and acini. In the infiltrate, cytokine staining was assessed as a percentage of positively staining cells in a total of 500 cells, while in the acini and ducts the intensity of staining was estimated by devising a scale of 0–8, where 0 = no staining and 8 = maximal staining.

**VASCULAR PROLIFERATION**
Vascular proliferation was estimated by counting the number of blood vessels staining with factor VIII and expressing the results per mm².

![Figure 1](http://ard.bmj.com/)

**Figure 1** Phenotypic analysis of the mononuclear cellular infiltrate in Sjögren syndrome (■) and chronic sialoadenitis (□).

![Figure 2](http://ard.bmj.com/)

**Figure 2** Interleukin-8 staining in Sjögren's syndrome (A), chronic sialoadenitis (B), and normal controls (C). Transforming growth factor beta staining in Sjögren's syndrome (D), chronic sialoadenitis (E), and normal controls. A, B, D, E: 10 mm represents 34.5 μm; C, F: 10 mm represents 19.5 μm.
Adhesion molecule expression was assessed in the three areas of all salivary gland tissue sections and their distribution analysed. For the expression of NKI-L16, the number of NKI-L16 positive cells in a total number of 500 LFA-1 positive cells was counted and the results expressed as a percentage.

RESULTS

Histological features and cellular composition

Minor salivary gland biopsies from patients with SS demonstrated typical focal lymphocytic clusters of more than 50 cells per focus. In contrast, in CS the infiltrating cells were randomly scattered throughout the gland. No infiltrating cells were observed in normal controls. Significantly greater numbers of infiltrating cells per mm² of tissue were seen in SS (1717 ± 1 (1083-3) cells/mm²) compared with CS (537 ± 0 (584-2) cells/mm²) (p < 0.031). Phenotypic analysis of the infiltrating cells in the two pathological groups revealed no statistically significant differences (fig 1). It is worth noting however, that the CD4:CD8 ratio was greater in SS (2.8) compared with CS (1.8), reflecting the greater number of CD4 cells seen in SS (336 (16.4) cells/mm²) compared with CS (21.2 (13.5) cells/mm²). The number of CD8 cells was similar in the two groups (14.6 (10.3) and 16.6 (12.1) cells/mm², respectively). Greater numbers of CD20 cells were counted in SS (20.0 (9.5) cells/mm²) than in CS (13.3 (8.9) cells/mm²). There was a tendency to greater numbers of CD68 cells in CS (16.2 (10.9) cells/mm²) compared with SS (8.0 (5.4) cells/mm²). The number of blood vessels per mm² of tissue in SS (134.1 (21.8) vessels/mm²) was not significantly different from that in CS (129.5 (10.4) vessels/mm²), but in both diseases more blood vessels were present compared with normal subjects (100.5 (10.8) vessels/mm²) (p = 0.015 and p = 0.004 versus SS and CS, respectively).

Cytokine expression

Mononuclear cellular infiltrate. IL-1α, IL-1β, TGFβ and GM-CSF were expressed in the MNC infiltrate of both SS and CS (fig 2). No statistically significant differences were noted in the percentages of positively staining cells in the two patient groups (fig 3). IL-8 expression was minimal (<2%) in the MNC infiltrate of all minor salivary glands examined.

Ducts. All the cytokines studied were strongly expressed in the epithelial cells of the ducts in SS and CS, but at a much lower level in normal controls. As can be seen in figure 4, the intensity of cytokine staining was significantly greater in the disease groups compared with control subjects, except for GM-CSF. No statistically significant differences were noted in the intensity of staining between SS and CS. There was no correlation between the intensity of ductal cytokine staining and the number of infiltrating cells surrounding the same duct (data not shown).

Acini. All cytokines studied were expressed in the acini of SS, CS, and normal controls. The intensity of cytokine staining in the acini...
LFA-3 was expressed by acini, ducts, blood vessels, and MNC in both SS and CS. ICAM-1 was equally detected on blood vessels and MNC of both diseases. Likewise, LFA-1 and CD2, found on MNC, were equally expressed in SS and CS. However, there was a striking difference in expression of NKI-L16, which was found on 33.6 (10.1)% and 15.3 (4.3)% of LFA-1 positive cells in SS and CS, respectively (p < 0.002) (fig 6).

Discussion

In this study, analysis of cytokine expression by the infiltrating MNC revealed a similar percentage of IL-1α, IL-1β, IL-8, GM-CSF and TGFβ positively staining cells in Sjögren’s syndrome and chronic sialoadenitis. In both disease groups, the ducts and acini showed more intense staining of all cytokines examined compared with normal control subjects. No significant difference was noted between SS and CS. Acinar staining was less intense than ductal staining, supporting the previous suggestion of a minor role for the acinar epithelial cells compared with ductal epithelial cells in the salivary gland inflammatory process as shown by the expression of class II molecules in the epithelial cells. The HLA-DR upregulation may be caused by IFNγ produced locally at the site of inflammation, as it is supported by the reported increase in IFNγ staining in SS compared with controls. The aberrant expression of class II molecules could be related to the potential antigen presenting function of epithelial cells, as described in the epithelial cells from gut and respiratory mucosae, epidermis, and renal proximal tubules.

The strong expression of TGFβ which we observed in the epithelial, acinar, and infiltrating cells clearly demonstrates that in SS
there is no obvious TGFβ deficiency, but the activation state of TGFβ needs to be determined before any firm conclusions can be made. We did not demonstrate a correlation between the intensity of cytokine ductal staining and the number of surrounding MNCs. Our findings do not support other workers’ observations of high expression of TGFβ in salivary glands of patients with SS, with a minimal infiltrate compared with those with an intense infiltrate. It is possible that an unknown cytokine or other inflammatory mediator not examined in this work may be responsible for the periductal clustering of the infiltrating cells so typical in SS. Furthermore, it is necessary to study agonists and antagonists simultaneously, as the balance between these two may determine the nature of the disease.

Although there were greater numbers of CD4 and CD20 cells in SS compared with CS, the difference did not reach statistical significance. However, the number of MNCs throughout the SS salivary glands was three times greater. The organisation of MNC into foci in SS and sparsely diffuse infiltrates in CS was a striking difference between the two groups. In chronically inflamed non-lymphoid tissues, infiltrating lymphocytes form aggregates which are functionally and structurally similar to follicles found in lymph nodes or tonsil. The presence of focal lymphoid aggregates in the rheumatoid synovium is associated with greater production of cytokine by the synovial membrane. In this study, we confirmed the presence of similar large lymphocyte aggregates in the salivary glands in SS, but were not able to show greater cytokine staining in the aggregates seen in SS.

The histological differences noted between SS and CS cannot be attributable to a greater degree of vascularisation in SS, as the numbers of blood vessels in SS and CS were similar. However, the differences may lie in the expression of adhesion molecules: we demonstrated a striking increase in the percentage of NKI-L16/LFA-1 cells in SS compared with CS. Cell adhesion by LFA-1 is established by the binding of LFA-1 to its ligands (ICAM-1 and ICAM-2) on the opposing cell, resulting in homotypic and heterotypic aggregation. In this study, salivary gland expression of NKI-L16, the activated LFA-1 epitope, correlated well with the capacity of cells to aggregate and to form the characteristic foci seen in the salivary glands of patients with SS. A preponderance of memory cells with CD45RO marker has been shown in SS salivary glands; these cells have been reported to adhere better to endothelial cells and to form large homotypic cell clusters compared with CD45RO negative cells. It has been suggested that the LFA-1/ICAM-1 and vascular leukocyte antigen-4 (VLA-4)/VCAM-1 pathways might be the main routes for T cells to migrate to sites of salivary gland inflammation in SS. This suggestion is supported by the extensive coexpression of LFA-1 or VLA-4 and CD45RO by CD4 T cell infiltrates adjacent to ICAM-1 or VCAM-1 postcapillary venules. These findings may point to a cytokine mediated upregulation of ICAM-1 and VCAM-1, facilitating the recruitment of LFA-1/NKI-L16 and VLA-4 cells and resulting in the infiltration of the salivary glands in SS.

In conclusion, this study shows that SS is characterised by periductal clustering of MNC, while CS is characterised by a sparse infiltrate. No significant differences in the state of infiltrating MNC and cytokine expression in the salivary glands of the two patient groups, but there was upregulation of NKI-L16 in the MNC of SS, suggesting that the typical MNC aggregation seen in SS may be mediated by activation of LFA-1.

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