Development of Autoimmune Exocrinopathy Resembling Sjögren’s Syndrome in Estrogen-Deficient Mice of Healthy Background

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Although a number of autoimmune diseases are known to develop in postmenopausal women, the mechanisms by which estrogen deficiency influences autoimmune lesions remain unclear. We speculate that antiestrogenic actions might be a potent factor in the formation of pathogenic autoantigens. Previously, we have identified 120-kd α-fodrin as an important autoantigen in Sjögren’s syndrome (SS). When healthy C57BL/6 (B6) mice were treated with an ovariectomy (Ovx), we found a significant increase in TUNEL*-apoptotic epithelial cells in the salivary gland cells associated with α-fodrin cleavage during 2 and 3 weeks after Ovx. By contrast, no apoptotic cells were found in estrogen receptor-α knockout mice. In in vitro studies using primary cultured mouse salivary gland cells and human salivary gland cells, we found a cleavage product of 120-kd α-fodrin in cells that had undergone tamoxifen (Tam)-induced apoptosis through caspase activation, especially caspase-1. Adoptive transfer of α-fodrin-reactive T cells into Ovx-B6 and -SCID mice resulted in the development of autoimmune exocrinopathy quite similar to SS. These results suggest that estrogen deficiency exerts a crucial influence on autoantigen cleavage, and may cause, in part, autoimmune exocrinopathy in postmenopausal women. (Am J Pathol 2003, 163:1481–1490)

Loss of ovarian function following menopause results in functional failures of the immune system, bone metabolism, and endocrine system. Estrogenic action has been suggested to be responsible for the strong female preponderance of many autoimmune diseases, including systemic lupus erythematosus (SLE), scleroderma, rheumatoid arthritis (RA), and Sjögren’s syndrome (SS).1–4 Sex hormones influence both humoral and cell-mediated immune responses in a number of experimental models.5–9 Previous reports indicate that the increase in autoantibody production as a result of estrogen deficiency is mediated by cytokines such as interleukin-6 (IL-6), interferon-γ (IFN-γ), and tumor necrosis factor-α (TNF-α), and that estrogen plays an important role in the regulation of B-lymphocyte development in mouse bone marrow.10–12

Recently, we have demonstrated that the dysfunction of regulatory T cells as a result of estrogen deficiency may play a crucial role on acceleration of organ-specific autoimmune lesions, and that estrogenic action influences target epithelial cells through Fas-mediated apoptosis in a murine model for SS.13 Although autoimmune diseases are triggered by various environmental factors, such as hormonal changes, microbial infections, stress, and aging,14,15 much less is known about the role of estrogen deficiency on the formation of autoantigen. We hypothesize that estrogen deficiency may influence the formation of pathogenic autoantigen in target organs through a T-cell-independent pathway.

Previously, we have identified 120-kd α-fodrin as an important autoantigen in both NFS/sld murine SS model and in patients,16 but the mechanisms of α-fodrin cleavage in the salivary gland cells remain unclear. Our recent study has been strongly suggestive of essential roles of caspase cascade for α-fodrin cleavage leading to tissue destruction in primary SS.17 α-fodrin is a ubiquitous, heterodimeric calmodulin-binding protein that is cleaved by calcium-activated protease (calpain) in apoptotic cells and caspase through Fas-mediated apoptosis in Jurkat cells.18–20 The fodrin α-subunit of various cells has also been shown to be cleaved in association with apoptosis.21–23 Several reports have demonstrated that estrogen may play an inhibitory role on apoptosis in endothelial cells, breast cancer cells, cardiac myocytes, prostate cells, and neuronal cells.24–27 Moreover, it has been noted that some enzymatic activities are elevated in postmenopausal women compared with normal healthy women.28,29

The aim of this study was to analyze the effect of estrogen deficiency on the formation of pathogenic autoantigen. Moreover, caspase activity in mouse salivary

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gland cells stimulated by an antiestrogenic action has been analyzed, indicating that estrogen deficiencies may play a pivotal role in autoantigen cleavage initially triggered in the salivary and lacrimal gland.

Materials and Methods

Mice and Treatments

Female C57BL/6 (B6) mice (H-2b) were purchased from Japan SLC (Shizuoka, Japan), and maintained in a specific pathogen-free (SPF) mouse colony and given food and water *ad libitum*. Estrogen receptor-α knockout mice (ERαKO) on a B6 strain background were purchased from Taconic (Germantown, NY). Female SCID C.B-17-scid/scid mice (H-2d), purchased from Japan SLC (Shizuoka), were used to confirm cell transfer experiment. Normal female B6 (H-2b), and BALB/c mice (H-2d), purchased from Japan SLC (Shizuoka), were used to obtain antigen-stimulated T cells. Mice were ovarioctomized (Ovx) at 4 weeks of age and compared with sham-operated (Sham, in both strain) mice. At 1 to 6 weeks after Ovx, all organs were removed from the mice and analyzed.

Histological Analysis

All organs were removed from mice, fixed with 4.0% phosphate-buffered formaldehyde (pH 7.2), and prepared for histological examination. The sections were stained with hematoxylin and eosin. The disease incidence was determined using the histological score of inflammatory lesions by White and Casarret, estimated by three independent, well-trained pathologist in a blinded manner.

In Situ End-Labeling of Fragmented DNA (TUNEL)

Apoptotic cells were detected in sections using the *in situ* TUNEL Kit (Wako Pure Chemical Industries, Ltd., Osaka, Japan). Briefly, sections were incubated with proteinase K (20 μg/ml) for 10 minutes, and then presoaked in TdT buffer (0.5 μmol/L cacodylate, 1 μmol/L CoCl₂, 0.5 μmol/L dithiothreitol, 0.05% bovine serum albumin, 0.15 mol/L NaCl) for 10 minutes. Sections were incubated for 2 hours at 37°C in 25 ml of TdT solution, containing 1X terminal transferase buffer, 0.5 nmol of biotin-dUTP, and 10 U of TdT (Wako Pure Chemical). After the TdT reaction, sections were soaked in TdT blocking buffer (300 nmol/L NaCl, 30 nmol/L tri-sodium citrate-2-hydrate), incubated with HRP-conjugated streptavidin for 30 minutes at room temperature, and developed for 10 minutes in phosphate-buffered citrate (pH 5.8) containing 0.6 mg/ml DAB. Nuclei were counterstained with hematoxylin.

Western Blot Analysis

Western blot analysis with anti-human α-fodrin (Affiniti, Mamhead, UK) was performed. Briefly, the cells were incubated in 20 mmol/L Tris-HCl (pH 8.0), 20 mmol/L NaCl, 0.5% Triton X-100, 5 mmol/L ethylenediaminetetraacetate (EDTA), and 3 mmol/L MgCl₂ lysis buffer. After centrifugation for 20 minutes at 12,000 × g at 4°C, supernatant was extracted and used for sample. Ten micrometers of each sample per well was used for sodium dodecyl sulfate-polyacrylamide gel electrophoresis. Protein binding was visualized with enhanced chemiluminescence Western blotting reagent (Amersham Biosciences, Arlington Heights, IL). Control for protein loading was provided with anti-human α-tubulin, or GAPDH monoclonal antibody (Sigma Chemical Co., St. Louis, MO). To detect caspase-1 in cultured human cells and mouse tissues, Western blot analysis was performed by the indicated methods using anti-human caspase-1 (Sigma Chemical Co.), and anti-human active form caspase-1 (p20 subunit; Upstate Biotech, Charlottesville, VA) polyclonal antibody. An anti-human caspase-1 polyclonal antibody is known to cross-react with mouse lysate.

Primary Culture of Mouse Salivary Gland (MSG) Cells

Mouse salivary gland (MSG) epithelial cells were prepared as previously described. Briefly, mouse salivary glands were minced into 1-mm² pieces, washed with Hank’s balanced salt solution (HBSS) without Ca²⁺ and Mg²⁺, and placed in a 60-mm dish containing HBSS with 0.76 μg/ml EDTA, 4.9 μg/ml L-ascorbic acid, and 4.9 μg/ml reduced glutathione. Fragments were washed with Dulbecco’s modified Eagle’s medium/soybean trypsin inhibitor, and placed in a mixture of collagenase (750 U/ml of type I) and hyaluronidase (500 U/ml of type IV) dissolved in DMEM/F12 containing 10% fetal bovine serum (FBS). The digest suspension was passed through a 100-μm nylon mesh filter. Adherent cells after culture in DMEM containing 10% FBS for 24 hours at 37°C were isolated as salivary gland epithelial cells. Apoptotic cells were detected by flow cytometry with an EPICS flow cytometer (Beckman Coulter, Miami, FL) using the Annexin V-FITC Apoptosis Detection kit (Genzyme, Cambridge, MA).

Cell Transfection

We used polymerase chain reaction (PCR) techniques to generate derivatives of a human caspase-1 promoter-
luciferase construct. The following forward and reverse oligomers were used as primers to create a promoter of the human caspase-1 gene: 5’TACGCGAGTTGAGAACTCTTACCTG3’, 5’GATCTAGAGGCTTTTCTCCTCCCT’. The PCR-amplified promoter fragments, including Pst-1/Xba-1 site, were cloned into the multiple cloning site of the pGL3-basic vector (Promega, Charbonnieres, France), upstream of the luciferase gene. The caspase-1 promoter-luciferase gene were transfected into HSG cells using LipofectAMIN (Promega). The vector pGL3-basic (lacking a promoter) and the vector (Promega) pGL3-control served as negative and positive controls, respectively. Briefly, the transfection medium, containing 10 μg of plasmid DNA and 60 μL of Lipofectin reagent in 2 ml of serum-free DMEM was incubated for 20 minutes at room temperature and then diluted with serum-free DMEM to a final volume of 5 ml and added to HSG cells, plated the day before. The transfection process occurred at 37°C for 5 hours, then 5 ml of DMEM containing 20% fetal calf serum (FCS) was added to the cells.

**Luciferase Assay**

The transfected cells were incubated for 24 hours and stimulated for the last 2 hours with Tam (1 × 10^−7M). Pretreatment with 17β-estradiol (1 × 10^−10M) was performed during last 12 hours. After rinsing with phosphate-buffered saline, cells were lysed with reporter lysis buffer (Promega), and cell extracts were used for luciferase assay with the Promega kit in a luminometer (Promega). To control transfection efficiency, pSVβ-galactosidase plasmid (Promega) was cotransfected with the luciferase reporter constructs in a 1:4 ratio. The results showed that the difference in the relative efficiency of transfection between constructs was negligible.

**Caspase Activities**

Caspase activities in Tam-induced apoptosis in HSG cell extracts and mouse various tissues were assayed using Caspase-Family Colorimetric Substrate Set (BioVision Inc., Palo Alto, CA). Briefly, tissue or cell lysates were incubated with pNA-conjugated substrates (200 mmol/L, caspase-1, -2, -3, -5, -6, -8, and -9 substrate: Ac-YVAD-pNA, Ac-IETD-pNA, and Ac-LEHD-pNA) at 405 nm in a microtiter plate reader using a 96-well plate. Absorbance of each sample was read to 6 weeks after the cell transfer. To confirm cell transfer experiments, Ovx- and Sham-SCID mice (n = 7 and n = 5) were transferred with α-fodrin-reactive T cells obtained from BALB/c mice in the same manner.

**Adoptive Transfer**

To obtain α-fodrin-reactive T cells, B6 mice were injected subcutaneously with 20 μg recombinant α-fodrin protein (JS-1) and Freund’s complete adjuvant (ICN Biochemicals) at 4 weeks of age, and i.p. injections of 20 μg JS-1 and Freund’s incomplete adjuvant (ICN Biochemicals) were performed at 6 weeks of age. OVA (10 μg/head)-reactive T cells were obtained as the same manner for control experiments. After 2 weeks later (8 weeks of age), mice were sacrificed and the splenic T cells were obtained as donor cells. As recipients, female B6 mice were ovariectomized (Ovx, n = 7) or sham-operated (Sham, n = 5) at 4 weeks of age. After 2 weeks (6 weeks of age), Ovx- and Sham-mice were transferred i.p. with 5 × 10^6 α-fodrin-reactive T cells. The transfected mice were analyzed at 4 and 8 weeks after the cell transfer. To confirm cell transfer experiments, Ovx- and Sham-SCID mice obtained as donor cells. As recipients, female B6 mice were ovariectomized (Ovx, n = 7) or sham-operated (Sham, n = 5) at 4 weeks of age. After 2 weeks (6 weeks of age), Ovx- and Sham-mice were transferred i.p. with 5 × 10^6 α-fodrin-reactive T cells. The transfected mice were analyzed at 4 and 8 weeks after the cell transfer. To confirm cell transfer experiments, Ovx- and Sham-SCID mice (n = 7 and n = 5) were transferred with α-fodrin-reactive T cells obtained from BALB/c mice in the same manner.

**Results**

**Apoptosis Induced in Salivary Gland Cells by Ovariectomy (Ovx)**

To examine the in vivo effects of estrogen deficiency in normal B6 and ERαKO mice, Ovx was performed at the age of 4 weeks. A radioimmunoassay confirmed that β-estradiol was not detectable in the sera of Ovx-mice (Ovx-B6 and Ovx-ERαKO, not detected; Sham-B6, 27.4 ± 2.8 pg/ml; Sham-ERαKO, 26.8 ± 3.6 pg/ml). At 1 to 6 weeks after Ovx, an in situ apoptosis detection assay was performed using all organs. Although we found a significant increase in TUNEL+ apoptotic cells in the salivary gland sections of B6 mice, but not ERαKO mice, at 2 or 3 weeks after Ovx (Figure 1), no significant apoptosis was observed in any other organs of Ovx and Sham mice. To define TUNEL labeling due to the effect of Ovx in vivo, we examined Annexin V-flow cytometric analysis using primary cultured MSG cells, indicating that a small proportion of apoptotic cells (12.5%) in Ovx mice was found, but not in Sham mice (3.1%) (Figure 1A). No significant difference in number of TUNEL+ apoptotic cells of Ovx and Sham mice was observed at 0, 1, 4, 5, and 6 weeks after Ovx (Figure 1B). Thus, apoptotic changes in the salivary glands of normal mice were observed transiently at 2 or 3 weeks after Ovx, supposing...
Figure 1. A: Detection of TUNEL*-apoptotic cells in the salivary gland sections from Ovx- and Sham-B6 mice, but not from ERαKO mice, at 3 weeks after Ovx (arrows). Annexin V-flow cytometric analysis using MSG cells demonstrated 12.5% positive cells detected in Ovx-B6 mice, but 3.1% positive in Sham-B6 mice. Data are representative in triplicate. B: A significant increase of apoptotic epithelial cells was observed in the salivary gland tissues from Ovx-B6 mice, not from ERαKO mice, restricted at 2 and 3 weeks after Ovx. The percentage of epithelial cells staining positively with TUNEL was enumerated using a 10×20-grid net micrometer disk covering an objective of area 0.16 mm². Data were analyzed in 10 fields per section and expressed as mean percent ± SD in five mice examined per group. (*P < 0.05, Student’s t-test.)
that antiestrogenic action to the epithelial tissues seems to be transient in vivo.

Effect of Estrogen Deficiency on α-Fodrin Proteolysis

We next investigated whether estrogen deficiency is involved in the formation of pathogenic autoantigens in salivary glands. To analyze α-fodrin proteolysis in the salivary glands from Ovx-B6 mice, Western blot analysis was performed using tissue samples. Although an intense band expressing 120-kd α-fodrin was found in the salivary gland samples from Ovx-B6 mice, no cleavage products of α-fodrin were observed in other organs from Ovx mice or any of the samples from Sham mice, while expression of α-tubulin as a control was present (Figure 2). Thus, estrogen deficiency induces in vivo proteolysis of α-fodrin in association with apoptosis in the normal salivary gland cells.

Tamoxifen (Tam)-Induced Apoptosis and α-Fodrin Cleavage in Salivary Gland Cells

It has been reported that the antiestrogen tamoxifen (Tam) induces cell death in the human breast cancer cell line MCF-7.32 To examine whether Tam induces apoptosis in the mouse and human salivary gland (MSG and HSG) cells, the cells were treated with 1 × 10^-10 to 1 × 10^-6 (M) Tam for 48 hours. We found a time- and concentration-dependent increase in number of apoptotic MSG and HSG cells until 48 hours (data not shown). In contrast to the colon cancer cells (HT-29 and Colo201) or Jurkat cells, apoptosis was induced in MSG, HSG, and MCF7 cells treated with Tam (Figure 3B). Of importance is that the 240-kd α-fodrin in Tam-induced apoptotic MSG and HSG cells was cleaved into 120-kd fragment in a time-dependent manner on Western blotting (Figure 3B). In Tam-induced apoptotic MCF7 cells, negligible levels of cleaved products of α-fodrin were found on Western blotting. We next examined whether estrogen could inhibit Tam-induced apoptosis of MSG and HSG cells. As shown in Figure 3C, Tam-induced apoptosis of MSG and HSG cells was significantly reduced by the pretreatment with estrogen. These data indicate that apoptosis of mouse and human salivary gland cells followed by α-fodrin cleavage into 120-kd fragment could be induced by an antiestrogenic action.

Participation of Caspases in Estrogen-Deficient Salivary Gland Cells

We next investigated whether apoptotic proteases are involved in the α-fodrin cleavage during Tam-induced apoptosis in MSG and HSG cells. A significant increase in caspase-1 activity was detected with relatively elevated caspase-3 and -8 activity on Tam-induced apopto-

Figure 2. Detection of α-fodrin cleavage in various tissues from Ovx- and Sham-B6 mice on Western blot analysis showing distinct band of 120 kd in the salivary gland tissue alone from Ovx-B6 mice. α-tubulin protein as internal control was present. Data are representative of five mice in each group.

Figure 3. A: Effect of Tam (1 × 10^-7 M) on apoptosis in various mouse and human cells (MSG, HSG, MCF-7, Jurkat, HT-29, and Colo201). Staurosporin (1 μmol/L) was used as common apoptotic reagent. B: α-fodrin cleavage into 120 kd in MSG and HSG cells induced by Tam. Western blot analysis was performed with mouse monoclonal Ab to α-fodrin. C: Tam-induced apoptosis of MSG and HSG cells was significantly reduced by the pretreatment with estrogen. Apoptotic cells were detected by flow cytometer using propidium iodide staining and Annexin V-FITC. The data are the mean ± SD from three individual experiments.
tic HSG cells (Figure 4A). In addition, Tam-induced apoptosis in HSG cells was inhibited considerably by the treatment with caspase inhibitors zVAD, DEVD, and WEHD (Figure 4B). We next examined the level of the caspase activities (caspase-1, -2, -3, -5, -6, -8, and -9) in various organs from Ovx- and non-Ovx-B6 mice. Figure 4C shows that a significantly elevated caspase-1 activity in the salivary gland tissues is observed in Ovx-B6 mice with slight elevation of caspase-3, and caspase-8 activity. No significant differences in caspase activities were observed in other organs.

Figure 4. A: A significant increase in caspase-1 activity with relatively elevated caspase-3, and caspase-8 activity on Tam-induced apoptosis in HSG cells. B: Tam-induced apoptosis in HSG cells was inhibited considerably by the treatment with caspase inhibitors zVAD, DEVD, and WEHD. C: A significantly elevated caspase-1 activity in the salivary gland tissues is observed in Ovx-B6 mice with slight elevation of caspase-3, and caspase-8 activity. No significant differences in caspase activities were observed in other organs.
with slight elevation of caspase-3 and 8 activity. No differences in caspase activities were observed in other organs. These results suggest that estrogen deficiency stimulates caspase activity, especially caspase-1, in the salivary gland tissues in vivo. We then examined whether Tam could influence the promoter activity of caspase-1 in HSG cells. After transfection with the plasmid containing the caspase-1 promoter-ligated upstream of the luciferase gene, HSG cells were stimulated with Tam, and then luciferase assay was performed. Figure 5A shows an increased promoter activity of caspase-1 after stimulation with Tam in HSG cells, but not in MCF-7, Jurkat, HT-29, and colo201 cells. The increase in caspase-1 promoter activity observed in HSG cells was significantly reduced by the addition of estrogen. We observed a time-dependent expression of caspase-1 (p20) activities until 48 hours on Western blot analysis (Figure 5B). Figure 5C shows that a distinct expression of caspase-1 (p20) in the salivary gland tissues is observed in Ovx-B6 mice. These results suggest that the salivary gland cell apoptosis could be induced by caspase activation, especially caspase-1, in estrogen deficient state.

Adoptive Transfer of α-Fodrin-Reactive T Cells into Ovx-B6 and -SCID Mice

We examined the adoptive transfer experiments using α-fodrin-reactive T cells into Ovx-B6 mice as shown in protocol (Figure 6A). Before the cell transfer, we confirmed the proliferative T-cell response against JS-1, not against lysozyme, albumin, and ovalbumin, of JS-1-immunized mice (Figure 6B). Consequently, inflammatory lesions developed exclusively in the salivary and lacrimal gland at 4 and 8 weeks after the transfer with $5 \times 10^6$ α-fodrin-reactive T cells, not with $5 \times 10^6$ OVA-reactive T cells, while no inflammatory lesions in any other organs were detectable (Table 1 and Figure 6C). When $5 \times 10^6$ α-fodrin-reactive CD4$^+$ or CD8$^+$ fractionated T cells were transferred into Ovx-B6 and Ovx-SCID mice, no inflammatory lesions were observed in any organs (data not shown). Proliferative T-cell response against recombinant α-fodrin (JS-1) was clearly observed in spleen cells from transferred Ovx-B6 mice (Figure 6D). In Ovx-SCID mice, inflammatory lesions were also induced in the salivary and lacrimal gland at 8 weeks after the transfer with $5 \times 10^6$ α-fodrin-reactive T cells, while no inflammatory lesions in any other organs were detectable (Table 1). Significant proliferative T-cell response against recombinant α-fodrin (JS-1) was observed in spleen cells from transferred Ovx-SCID mice (Figure 6E).

Discussion

The mechanisms responsible for the development of autoimmune diseases during the postmenopausal stage are still unclear. Previous reports concerning gender differences in autoimmunity have suggested that estrogen influences the cytokine production of effector cells and autoantibody production. The distinct immune environments in males and females underlie many of the gender-related differences in autoimmunity. These environments are established by the cytokines that are released by immune cells, particularly T helper (Th) lymphocytes. Sex hormones, pituitary hormones including prolactin, and growth hormones, as well as liver-derived insulin-like growth factor-1 affect autoimmune diseases by modulating cytokine productions. Women have higher levels of these hormones than men. Estrogen withdrawal after menopause leads to an increase in the production of cytokines, such as granulocyte-macrophage colony-stimulating factor, IL-1, IL-6, and TNF-$\alpha$. Although many studies have described the effects of estrogen on cytokine productions in effector cells, much less is known about the effect of estrogen deficiency in...
A

Donor
Female B6 or
BALB/c mice

4w 6w 8w

JS-1/CFA (s.c.) JS-1/IFA (i.p.)

T cell transfer

Recipient
Ovx-B6 or SCID

4w 6w 10w 14w

Analysis

B

C57BL/6

Salivary

Lacrical

Ovx alone

Ovx+Transfer

C

D

E

B6

SCID

[Graphs and images showing data comparison and experimental results]
target organs of postmenopausal women. Thus, it is required to determine how estrogen deficiency influences the expression of autoantigens in target cells before the infiltration of lymphocytes into the salivary glands.

In this study, we have demonstrated a significant apoptosis associated with α-fodrin cleavage in the salivary gland cells of estrogen deficient healthy B6 mice. Moreover, inflammatory lesions developed exclusively in the salivary gland cells of estrogen deficient healthy B6 mice. It is assumed that estrogen acts as a negative regulator of caspase-1 activity in the salivary gland cells.

When MSG and HSG cells were induced to undergo apoptosis using Tam, the 240-kd α-fodrin was cleaved into a single detectable fragment of 120 kDa. Among the substrates cleaved during apoptosis are nuclear autoantigens such as PARP, U1-70-kd, the nuclear lamin, and DNA-dependent kinase. In organ-specific autoimmune diseases, no evidence in vivo cleavage of self-proteins during apoptosis has been demonstrated. Our data suggest that antiestrogenic actions have a potent effect on the proteolysis of α-fodrin autoantigen in the salivary gland through up-regulation of caspase-1 activity. These results strongly suggest that α-fodrin fragments induced by Ovx may play an important role in the development of autoimmune lesions as a pathogenic autoantigen. Molecular mechanisms responsible for tissue-specific apoptosis induced by estrogen deficiency are further investigated.

In conclusion, we have demonstrated that antiestrogenic actions, including estrogen deficiency or Tam stimulation, may have a crucial influence on apoptosis and

Table 1. Frequency of Inflammatory Lesions in the Salivary and Lacrimal Glands in Transferred C57BL/6 and SCID Mice Treated with Ovx

<table>
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<th>Treatment</th>
<th>No. of mice</th>
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<th>Parotid</th>
<th>Lacrimal</th>
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<td>5/7</td>
<td>7/7</td>
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<tr>
<td>Sham+transfer</td>
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<tr>
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<tr>
<td>8 weeks after transfer (C57BL/6)</td>
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<td>5</td>
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*Histological evaluation for frequency of inflammatory lesions in the salivary and lacrimal glands was done according to the method proposed by White and Casarett.30
†Female C57BL/6 (B6) and SCID mice were ovariectomized (Ovx) at 4 weeks of age.
‡Ovx-B6 and SCID mice were transferred intraperitoneally with $5 \times 10^6$ α-fodrin-reactive T cells at 6 weeks of age.
§Ovx-B6 mice were transferred with $5 \times 10^6$ OVA-reactive T cells. α-fodrin-reactive T cells were obtained from B6 or BALB/c mice as described in detail in the text.
α-fodrin proteolysis through an increased caspase activity in the salivary gland cells, suggesting a novel mechanism for the development of organ-specific autoimmunity in postmenopausal women.

References