Interepithelial lymphocytes of the rat submandibular gland

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Abstract: The distribution of interepithelial cells in the rat submandibular gland was investigated histologically and ultrastructurally. All the interepithelial cells were small lymphocytes with a smooth or shallow-indented nuclear outline, and were located below the level of the epithelial cell nuclei of acini and ducts. No other cells such as Langerhans cells or cerebriform cells, such as are commonly found in stratified squamous epithelia, were detected. A mean of 6.7 interepithelial lymphocytes per 1,000 salivary gland epithelial cells was found: 7.1 in acini, 4.6 in intercalated ducts, 6.7 in granular and striated ducts, and 8.3 in excretory ducts.

Introduction

Epithelia of external and internal surfaces of the mammalian body commonly include cells at various levels that differ from the majority of epithelial cells in possessing a clear halo around the nucleus. Such 'clear cells' are designated interepithelial or intraepithelial cells, and are variously classified as melanocytes, Langerhans cells, Merkel cells, lymphocytes, or others. Of these, an important function as a first immunological barrier has been ascribed to interepithelial lymphocytes which are distributed in both single-layered epithelia and stratified squamous epithelia, and some regions of the epithelia could probably have a function in the maturation and instruction of the lymphocytes equivalent to the function of the bursa Fabricii in birds.

The distribution and frequency of interepithelial lymphocytes occurring in normal condition have been studied in the intestinal tracts9, epidermis4,8, and oral mucosa4,7. However, there are no reports of such cells in the salivary gland epithelia. The purpose of the present paper is to report the distribution and frequency of interepithelial lymphocytes in the submandibular gland of the rat.

Materials and Methods

10 male Wistar rats weighing 250 to 300 g were used. The animals were maintained under conventional condition with exogenous antigen exposure.

Both right and left submandibular glands were fixed in 10 per cent neutral-buffered formalin immediately after sacrifice by ether anesthesia, embedded in paraffin, and 4 µ thin sections were stained with hematoxylin and eosin, periodic acid-Shiff and a silver impregnation method (Watanabe's method). In addition activity of N-AS-D-chloroacetate esterase was examined.

Specimens for ultrastructural investigation were fixed immediately in 4.0 per cent glutaraldehyde by immersion and post-fixed in 2.0 per cent osmium tetroxide, dehydrated, and embedded in resin. Semithin sections were stained with toluidine blue. Then the ultrathin sections were cut, stained with uranyl acetate and lead citrate, and examined under an Akashi LEM-2000 ultramicroscope.

Under the ultramicroscope, all recognizable interepithelial mononuclear cells were counted in relation to the number of salivary gland epithelial cells, and numbers of the interepithelial mononuclear cells per 1,000 salivary gland epithelial cells were computed.
in acini, intercalated, granular and striated, and excretory ducts.

**Results**

Light microscopy:
The common light microscopic findings of interepithelial mononuclear cells stained with hematoxylin and eosin were chromatin-dense and small round nuclei surrounded by clear halo, and indistinct cytoplasm (Figs. 1 and 2). The most of these interepithelial mononuclear cells were located below the level of the epithelial cell nuclei, and they occurred singly in both acini and ducts. Although a few lymphocytes and plasma cells were scattered in interacinal and periductal fibrous connective tissue stroma in part, they were not noted in the fibrous connective tissue stroma adjacent to the epithelial regions with interepithelial mononuclear cells (Figs. 1 and 2). No cells with activity of N-AS-D-chloroacetate esterase were found in either epithelia and fibrous connective tissue stroma.

Ultrastructural finding:
Ultrastructurally, 142 interepithelial mononuclear cells in the rat submandibular salivary gland epithelia were documented. All of the interepithelial mononuclear cells in the rat submandibular salivary gland epithelia were small lymphocytes, and no other cells such as Langerhans cells and Merkel cells distributed normally in the stratified squamous epithelia were found. These interepithelial lymphocytes were round to oval in shape with smooth cell-outlines, 4.0 to 6.0 μm in apico-basal diameter, and located in basal regions of the acini and ducts (Figs. 3 and 4). The cytoplasm of lymphocytes was sparse, containing free ribosomes, a few mitochondria and endoplasmic reticulum. The nuclei of lymphocytes were spherical in shape and contained abundant heterochromation, but nucleoli were inconspicuous. The margin of these nuclei were smooth (Fig. 3) or with shallow-indentations (Fig. 4). However, lymphocytes with deeply indented nuclei called 'cerebriform cells' were not found.

Quantitative evaluation:
The result of the count of interepithelial lymphocytes under the ultramicroscope in the

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Figs. 1 and 2. Light photomicrographs of interepithelial mononuclear cells (arrows) in acinus (1) and excretory duct (2) of rat submandibular salivary gland. No mononuclear cells or other inflammatory cells in the fibrous connective tissue stroma. Hematoxylin and eosin, Fig. 1 ×400 and Fig. 2 ×350.
Figs. 3 and 4. Photographs showing the ultrastructure of interepithelial lymphocytes in the acinus (3) and striated duct (4) of rat submandibular salivary gland. Fig. 3 shows small lymphocyte with a smooth nuclear outline, and Fig. 4 shows small lymphocytes with slight nuclear indentations. Arrows indicate basal lamina. Fig. 3 ×13,200 and Fig. 4 ×8,720.
quantitative distribution of interepithelial lymphocytes (IEL) in the rat submandibular salivary gland epithelia.

![Fig. 5. Quantitative distribution of interepithelial lymphocytes (IEL) in the rat submandibular salivary gland epithelia.](image)

A mean of 6.7 interepithelial lymphocytes per 1,000 salivary gland epithelial cells is found in rat submandibular gland: 7.1 in acini, 4.6 in intercalated duct, 6.7 in granular and striated ducts, and 8.3 in excretory duct.

### Discussion

A considerable amount of morphological research has been reported on the intercellular mononuclear cells of the epithelia, and most of such investigations so far have concentrated on the intestinal mucosa and epidermis in normal and pathological conditions. Recently, interepithelial mononuclear cells of the oral mucosa have been studied in detail.

The interepithelial mononuclear cell population of single-layered epithelia is almost completely composed of lymphocytes, and in stratified squamous epithelia other interepithelial mononuclear cells such as Langerhans cells and Merkel cells are distributed in addition. Furthermore, interepithelial lymphocytes have been found in normal visceral organs, i.e., in kidney reported by Bohle et al. and in thyroid gland reported by Toujas and Guelfi. Although their origin, significance, function and fate are still known only in fragments, it is thought that one of the important roles of interepithelial mononuclear cells, especially lymphocytes and Langerhans cells, is to form a first immunological barrier. In the oral mucosa, Burkhardt et al. and Bos and Burkhardt have reported an interesting quantitative and qualitative examination of the interepithelial mononuclear cell population in mice under different antigen exposure, with special attention to the lymphocytes and Langerhans cells. They say that interepithelial lymphocytes must be considered an integral component of the oral epithelium, since interepithelial lymphocytes are noted in germfree animals, and since interepithelial lymphocytes do not show a significant increase in specific pathogen-free or conventionally maintained animals compared to germfree ones. On the other hand, the number of interepithelial lymphocytes in the intestinal mucosa shows a significant decrease in germfree animals compared to conventionalized ones, but they are not absent in germfree animals.

Then there seem to be a fundamental difference in function between intestinal and oral interepithelial lymphocytes.

In the present study of rat submandibular gland, all of the interepithelial mononuclear cells were small lymphocytes, and no other cells distributed in stratified squamous epithelia were found. The nuclei of these interepithelial lymphocytes were round in shape with or without shallow indentation, but cells with deeply indented nuclei called ‘cerebriform cells’ were not found. Although animals used in the present examination were maintained conventionally, the appearance of interepithelial lymphocytes might not be the expression of a subclinical inflammation, since neither mono- nor poly-nuclear cells infiltrated in the interacinar and periductal fibrous connective tissue stroma adjacent to the epithelial regions with interepithelial lymphocytes.

The pathological and biological significance of the present report of interepithelial lymphocytes in normal salivary gland epithelia is of interest. It is well known that during embryonic growth salivary glands are closely associated with the lymphoid tissue. Such a relation between salivary gland epithelia and lymphoid tissues may give rise to some salivary gland lesions such as benign lymphoepithelial lesions, Warthin’s tumor, lymphoepithelial cyst, etc.

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抄録：唾液腺における上皮内リンバ球の分布を詳細に検索した報告は未だない。そこで筆者はラット額下腺を用いて超微構造的にこの点について検討を試みた。

検索には体重250〜300gの雄Wistarラットを用い、額下腺を摘出後直ちに固定し、通法の如く超薄切片を作製した。超微構造的に上皮内リンバ球の形態を観察するとともに、唾液腺上皮腺房部、介在部、顆粒管および線条部の主導管の4部位に分けて各部位における上皮細胞1,000個に対する上皮内リンバ球の数を算出した。

今回の検索で超微構造的に142個の上皮内リンバ球が観察された。これら上皮内リンバ球のすべては小リンバ球であり、球状の核を有し、胞体には遊離リボソームの他にはごくわずかなミトコンドリアと粗面小胞体をみるようになかった。また、核に深い切れ込みを有するいわゆるcerebriform型とよばれるリンバ球はみられなかった。なお、唾液腺上皮に特殊な胞体内顆粒や細胞間結合装置を有する上皮内細胞は認められなかった。

上皮内リンバ球の唾液腺上皮各部位における分布は上皮細胞1,000個に対して腺房部では7.1個、介在部では4.6個、顆粒管と線条部では6.7個、主導管では8.3個であった。

References


