Is There an Interspecific Diversity of the Thymic Microenvironment?

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Thymic epithelial cells (TEC) heterogeneity suggests the existence of functional subsets. Anti-cytokeratin (Anti-CK) monoclonal antibodies (MAb), markers of epithelial differentiation, have been used to detect TEC subsets in rodents and humans. These MAb revealed a different topography of CK-defined TEC subsets in mice and humans, leading us to carry out a comparative study of mammalian thymuses. Our study showed that the distribution pattern of cytokeratins in the thymic epithelium is complex and unique, with coexpression of CK typical of simple and stratified epithelia. Moreover, we demonstrated an interspecific diversity of CK expression within the thymic lobules. Interestingly, such diversity was not a general phenomenon for the expression of any thymic microenvironmental proteins, because the location of extracellular matrix components was essentially similar in the mammalian species studied.

KEYWORDS: Mammalian thymuses, thymic microenvironment heterogeneity, cytokeratins, thymic epithelium, extracellular matrix.

INTRODUCTION

Intrathymic lymphocyte differentiation events, including the selection of the T-cell repertoire, essentially occurs in the context of the so-called thymic microenvironment (Sprent et al., 1988; von Boehmer et al., 1989; Fowlkes and Pardoll, 1989; Blackman et al., 1990; Boyd and Hugo, 1991). As such microenvironment is rather complex—being composed mostly by the thymic epithelial-cell network together with other stromal cells and extracellular matrix—the precise role of each nonlymphoid component on the general process of thymocyte maturation remains to be determined. Thus, the characterization of these components would be helpful to further study the interactions between these thymic compartments, including using in vitro systems. Considering that thymic epithelial cells (TEC) pleiotropically influence steps of T-cell maturation (Kyewski, 1986; Schuurman, 1988; Haynes, 1990; van Ewijk, 1991), and that these cells are morphologically and antigenically heterogenous (Haynes, 1984; von Gaudecker, 1986; Kendall, 1988), one can imagine that TEC subsets might be involved in specific events of intrathymic T-cell differentiation. Yet, functionally defined TEC subsets were not isolated so far, despite the several TEC lines (Itoh, 1979; Nieburgs et al., 1985; Potworowski et al., 1986; Mizutani et al., 1987; Naquet et al., 1989) and the variety of available monoclonal antibodies (MAb) that are phenotypic markers of so-called TEC subsets (Haynes, 1984; van Vliet et al., 1984; de Maagd et al., 1985; Kaneshima et al., 1987; Takacs et al., 1987; Ison and Boyd, 1990). In this respect, a nomenclature for these anti-TEC MAb, subdividing them on the basis of their labeling patterns in 5 main groups called CTES—Clusters of Thymic Epithelial Staining—was proposed (Kampinga et al., 1989). Interestingly, these markers revealed a close antigenic similarity between the thymic epithelium and epidermis, with Hassall’s corpuscles corresponding to a final degree of TEC differentiation, as the stratum corneum keratinocytes (Haynes, 1984; Schmitt et al., 1987). In this context, anticytokeratin (anti-CK) MAb, epithelial differentiation markers, might be regarded as useful tools in the study of TEC subsets, as actu-
ally evidenced in some rodents (Nicolas et al., 1985, 1986; Colic et al., 1988a, 1988b; Savino and Dardenne, 1988a; Farr and Brady, 1989) and humans (Savino and Dardenne, 1988b). Nonetheless, the location of a given CK-defined TEC subset within the thymic lobules was different in mouse as compared to man (Savino and Dardenne, 1988a, 1988b), suggesting a species-specific diversity of CK expression in mammalian thymuses. These data prompted us to carry out a comparative study of CK-defined TEC subsets in mammals. This analysis actually showed an interspecies heterogeneity in the intrathymic location of CKs, including in early stages of thymus ontogeny (Meireles de Souza and Savino, 1993). Differently, such diversity was not observed regarding the expression of extracellular matrix proteins, herein applied as representatives of another thymic microenvironmental component. Additionally, the present work demonstrated that the pattern of CK expression by the thymic epithelium is complex and rather unique, with cytokeratins typically found in simple or stratified epithelia being coexpressed.

RESULTS

Detection of the Thymic Epithelial Cell network by Pan-Specific Anti-CK Antibodies

We analyzed the thymuses of mammals representatives of different phylogenetic branches, as seen in Fig. 1. In all species studied, both polyclonal and monoclonal antibodies pan-specific
for cytokeratins (HTK and KL4, respectively) stained the whole network of epithelial cells, distinguishing the cortical and medullary TEC, these latter being more voluminous. Moreover, subcapsulary-subseptal TEC were evidenced (data not shown).

**Topography of TEC Expressing Cytokeratins Typical of Simple Lining Epithelia**

In a first set of experiments, we studied the expression of CK8 and CK18 (strictly found in simple epithelia) and CK19, mainly detected in this type of epithelial tissues (Quinlan et al., 1985; Sun et al., 1985). In all cases, an interspecific diversity in CK distribution was seen. In primate thymuses, CK8 TEC were only observed in the medulla, a pattern exclusively found in this mammalian order. In contrast, the cortical region of rodent thymuses was consistently labeled by the anti-CK8 MAb, besides the medullary CK8 TEC in rat. Similarly, the rabbit thymic epithelium was also stained by the anti-CK8 MAb in both thymic regions. Differently, CK8 TEC were restricted to the cortico-medullary junction in the sheep thymus, and could not be detected in opossum thymus sections (Fig. 2).

Although CK8 and CK18 are a naturally occurring pair, their colocalization was solely integral in the primate and rodent orders. The major disparity of CK8 and CK18 immunodetection was found in the opossum thymus, because it was CK8-, being entirely labeled by the anti-CK18 MAb.

Regarding the intrathymic distribution of CK19, we also evidenced an interspecific diversity in all orders studied. Rabbit and opossum TEC networks were virtually CK19 negatives, whereas the thymic epithelium of sheep, hamster, and human were completely stained. In the other mammalian species tested, CK19 TEC were mostly medullary (Fig. 3).

![Figure 2](image-url)  
**FIGURE 2.** CK8/18-expression pattern in mammalian thymuses: (a) rat thymus section, showing the whole TEC network stained by the anti-CK8 MAb (×200); (b) CK8 cortical zone of rabbit thymus (×160); (c) hamster thymus section, showing CK18 subseptal and cortical TEC (×320); (d) CK18 cortical and medullary TEC in the opossum thymus (×250). C: cortex, M: medulla.
Localization of “Stratified” Cytokeratins in Mammalian Thymuses

We also evaluated the localization of cytokeratins typically found in rather complex stratified lining epithelia (Quinlan et al., 1985; Sun et al., 1985). One of the anti-CK MAb used for that, named KL1, recognizes the CK1/CK10 pair, the marker of terminal differentiation in the epidermal epithelium (Viac et al., 1983). Using this reagent, we detected the same distribution pattern of CK expression in both primate thymuses, with the whole thymic epithelium being labeled. Other mammals presented KL1 staining preferentially at the medullary region, with the number of KL1⁺ cells varying from species to species. These data are summarized in Fig. 4.

In addition to CK1/CK10, we studied the expression of CK13, typical of stratified epithelia of internal organs. In human, Rhesus monkey, and hamster thymuses, subcapsular-medullary TEC were consistently labeled by the anti-CK13 MAb. This pattern differed in mouse, rabbit and opossum species, in which a medullary restriction of CK13 localization was observed. Conversely, sheep thymic epithelium was totally stained by the anti-CK13 MAb, whereas rat thymic epithelial cells remained negative to this reagent (Fig. 5).

Immunoblot Detection of Thymic Cytokeratins

As the immunocytochemical reactivity patterns of anti-CK MAb presented an interspecific diversity in the intralobular localization of distinct cytokeratins, it might be possible that the same MAb recognized distinct CK in different species, though these proteins are evolutively conserved (Moll et al., 1982a; Fuchs and Marchuk, 1983;
Blumenberg, 1988). Thus, we used a one-dimensional immunoblot assay to approach the molecular specificity of some representative anti-CK MAb in thymic extracts from Rhesus monkey, hamster, rat, rabbit, and sheep. For that, we applied the monoclonal reagents KL1 and anti-CK18, respectively, examples of markers of stratified and simple lining epithelia.

The results obtained with KL1 clearly showed that, although this MAb presented different reactivity patterns in situ, it recognized the same CK pair of apparent molecular weight (MW) of 58 and 66 kDa in the various extracts (Fig. 6a). In parallel with these findings, the anti-CK18 MAb reacted with a 45-kDa CK protein band in monkey, rat, and sheep extracts, whereas their in situ staining pattern differed among these mammals (Fig. 6b).

FIGURE 4. KL1 reactivity pattern in mammalian thymuses: (a) Rhesus monkey TEC network totally stained by KL1 (× 160); (b) hamster thymus section, showing subseptal and medullary KL1+ TEC (× 160); (c) rabbit thymic medulla, with the majority of TEC being labeled by KL1 (× 250); KL1+ medullary TEC in thymuses from: (d) sheep (× 250), (e) rat (× 200), and (f) opossum (× 250). C: cortex, M: medulla; arrows indicate KL1+ subseptal TEC.
FIGURE 5. CK13-expression pattern in mammalian thymuses: (a) sheep thymus section, showing CK13' cortical and medullary TEC (x 160); (b) monkey thymus showing CK13 staining in the medullary region, including Hassall's corpuscles (large arrow) and subseptal TEC (short arrow) (x 250); (c) CK-13 labeling of hamster thymic medulla, including Hassall's corpuscles (arrows) (x 400); (d) opossum thymus section, showing CK13' medullary TEC (large arrow) and CK13' Hassall's corpuscles (short arrow) (x 250). C: cortex, M: medulla.

Distribution of Extracellular Matrix Components in Mammalian Thymuses

Considering the interspecific diversity of the thymic epithelium, as defined by the cytokeratin expression, we carried out a comparative analysis of thymic extracellular matrix proteins in the different mammalian species. We chose typical basement membrane proteins, namely, laminin, fibronectin, and type-IV collagen, as representatives of a distinct component of the thymic microenvironment, yet being largely produced by the thymic epithelium. Additionally, we had previously described their topographic distribution in human and mouse thymuses (Berrih et al., 1985; Lannes-Vieira et al., 1991).

In contrast to the CK interspecific diversity, the distribution of extracellular matrix proteins was strikingly similar in mammals, with labeling

FIGURE 6. Cytokeratins recognized by (A) KL1 and (B) anti-CK18 MAb in one-dimensional immunoblot assay of thymic extracts derived from (a) Rhesus monkey, (b) hamster, (c) rat, (d) rabbit, and (e) sheep. L: low-molecular-weight standard markers, H: high-molecular-weight standard markers.
being restricted to the typical basement membranes adjacent to capsule, septa, and blood vessels, together with a fine medullary network (Fig. 7). The rat thymus was the only one that
diverged, because it presented short and thin fibronectin, type-IV collagen, and laminin fibers both at the cortical and medullary regions.

DISCUSSION

The heterogeneity of the thymic microenvironment, and in particular of its epithelium component, is already largely accepted. Nonetheless, an interspecific diversity in the localization of molecules expressed by TEC has been much less investigated. In this regard, the present work brings consistent data revealing that the topography of TEC-expressing specific cytokeratins varies amongst mammalian species belonging to distinct phylogenetic orders (summarized in Fig. 8). This was initially suspected by our previous data on CK distribution in the mouse and human thymuses (Savino and Dardenne, 1988a, 1988b), and more recently by Colic et al. (1990) studying other mammalian species. Nevertheless, these findings were strictly based on immunohistochemistry, thus being unable to discard the possibility of epitope cross reactivity among distinct cytokeratins in different species. Actually, we showed herein that the same specific molecular weight cytokeratins could be evidenced with KL1 and anti-CK18 MAb in thymic extracts from several species, independently of the diversity in their immunohistochemically defined intrathymic localization. These data suggest that, for a variety of mammalian species, there is a topographical diversity of CK expression, or at least...
of CK epitopes. The approach of in situ hybridization using CK-specific mRNA or cDNA probes, or the use of polyclonal antibodies produced against synthetic CK peptides (Roop et al., 1984; Ruggiero et al., 1990) would be useful to clarify this question. Nonetheless, even if the integral CK molecules were not heterogeneously distributed among species, a CK epitope diversity was definitely detected. Interestingly, these epitopes could be functional, because modulations of CK staining patterns were observed in many physiological and pathological situations (Meireles de Souza and Savino, 1993).

The present work also demonstrates that the pattern of CK expression by the thymic epithelium is complex and rather unique, with cytokeratins typically found in simple or stratified epithelia being coexpressed. It is noteworthy that the colocalization of CK8, CK18, CK19, CK1/CK10 pair, and CK13 was exclusively found in this epithelium (compare to data presented in Quinlan et al., 1985). Whereas the CK1/CK10 pair is expressed by highly keratinized epithelia, as the epidermis, the CK13 is typical of many internal stratified epithelia, like the esophagus, trachea, and tongue, and in general are mutually exclusive. The complexity and uniqueness of the thymic epithelium CK expression were further emphasized by Heid et al. (1988) describing the presence of hair-type keratins in the thymus. Taken together, the previous findings suggest that the general classification of epithelial tissues based on their pattern of cytokeratin expression cannot be applied to the thymus. The interspecific diversity of CK expression, when analyzed in terms of clusters of thymic epithelial-cell staining, grouped the CK-defined TEC subsets in different CTES (Kampinga et al., 1989), depending on the mammalian species analyzed, as depicted in Table 1. Thus, CK8 TEC are classified as group I in rat, group II in mouse and hamster, and group XX in primates. Yet, anti-TEC MAb can be classified in different CTES groups according to the species (Kampinga et al., 1989). This point indicates that the applicability of CTES nomenclature is rather restrict.

In spite of interspecific diversity of cytokeratin localization within thymic lobules, important similarity was observed in species belonging to the same mammalian order in the phylogenetic tree. This concept can be inferred from Fig. 8. In particular, human and Rhesus monkey diverged only in the CK19 TEC distribution. Moreover, the anti-CK8, anti-CK18, and KL1 MAb reactivity patterns seemed to be exclusive features of this order.

We also observed a compartmentalization of the same CK in cortical or medullary regions in different mammals. Particularly, we verified that TEC recognized by KL1 or anti-CK13 MAb (and in many instances by the anti-CK19 MAb) were predominantly found in medulla. Actually, in the cases when these MAb labeled the whole (or most of) medullary region, subcapsullary TEC were also stained. This characteristic is shared with many other anti-TEC MAb (Haynes, 1984; de Maagd et al., 1985; Takacs et al., 1987; Kampinga and Aspinall, 1990; Izon and Boyd, 1990), suggesting profound similarities in subcapsullary-medullary TEC that might be related to a hypothetic common embryonic origin (Crouse et al., 1985; Lobach and Haynes, 1987). Nevertheless, studies on cytokeratin expression did not clarify the question of whether the thymic epithelium is derived from endoderm or endoectoderm (Le Douarin et al., 1984; von Gaudecker, 1986; Lampert and Ritter, 1988). In fact, the localization of the same CK in different thymic regions in diverse species may imply that anti-CK MAb are not appropriate embryonic markers.

In contrast to cytokeratin expression data, the study of the extracellular matrix molecule localization in mammalian thymuses revealed a conserved pattern, except for the rat thymus. This conservation suggests that these molecules may be involved in important events of thymic physiology, as recently proposed (Savino and Lannes-Vieira, 1991).

### Table 1

Classification of Mammalian Thymuses for Cytokeratin Expression Patterns Based on the "Clusters of Thymic Epithelial Staining" Nomenclature

<table>
<thead>
<tr>
<th>Anti-CK8</th>
<th>Anti-CK18</th>
<th>Anti-CK19</th>
<th>Anti-CK13</th>
<th>KL1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>XX</td>
<td>XX</td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td>Monkey</td>
<td>XX</td>
<td>XX</td>
<td>II.A</td>
<td>II</td>
</tr>
<tr>
<td>Hamster</td>
<td>III</td>
<td>III</td>
<td>V.C</td>
<td>XX</td>
</tr>
<tr>
<td>Mouse</td>
<td>III</td>
<td>III</td>
<td>V.E</td>
<td>V.A</td>
</tr>
<tr>
<td>Rat</td>
<td>I</td>
<td>I</td>
<td>V.E</td>
<td>V.C</td>
</tr>
<tr>
<td>Rabbit</td>
<td>III.A</td>
<td>XX</td>
<td></td>
<td>V.E</td>
</tr>
<tr>
<td>Sheep</td>
<td>XX</td>
<td>I</td>
<td>I</td>
<td>IV</td>
</tr>
<tr>
<td>Opossum</td>
<td>–</td>
<td>I</td>
<td>–</td>
<td>V.C</td>
</tr>
</tbody>
</table>

*XX: classification according to CTES nomenclature (Kampinga et al., 1989); < negative.
Lastly, if we consider these results together with the concept that thymocyte differentiation appears to occur in a similar way in different species (Fowlkes and Pardoll, 1989), and that functional TEC subsets might be related to diverse events of T-lymphocyte maturation, we are impelled to think that anti-CK and anti-TEC MAbs (yet presenting an interspecific diversity of staining) may not be functional markers of TEC subpopulations. These MAbs should be rather regarded as markers of microenvironmental variant series antigens in analogy to the recently proposed species-specific cell-surface antigens—the variant series—of developing thymocytes (Aspinall et al., 1991).

MATERIALS AND METHODS

Thymuses

Human thymus biopsies were derived from children aging from 6 months to 4 years that were submitted to surgeries due to cardiac malformations.

Besides human specimens, this study comprised material from different mammalian species, including animals from five phylogenetically different orders, as depicted in Fig. 1. Thymuses from Rhesus monkey (Macaca mulatta), hamster (Mesocricetus auratus), rat (Ratus norvegicus), rabbit (Oryctolagus cuniculus), sheep (Ovis aries), and opossum (Didelphis marsupialis) were provided by the animal facilities of the Oswaldo Cruz Foundation, and C57BL/6 mice (Mus musculus) were obtained from the animal house of the Department of Immunology, Sao Paulo University. All animals were young adults.

Antibodies

A panel of antibodies recognizing different cytokeratins was used. The polyclonal antibody HTK (Biosoft, Paris) and the monoclonal antibody KL4 (Immunotech, Marseille), pan-specific to cytokeratins, proved to label the whole thymic epithelium network (Savino and Dardenne, 1988a). The monoclonal reagent KL1, specific to CK1/CK10 pair (Viac et al., 1983), was purchased from Immunotech (Marseille), whereas other MAbs, respectively, monospecific to CK8, CK18 (both typical of simple epithelia), or CK19, were Amersham products (Buckinghamshire, UK). The MAb anti-CK13, marker of esophageal differentiation, was obtained from Progen Biotechnik (Heidelberg). General characteristics of these antibodies recognizing distinct thymic epithelial cells are summarized in Table 2.

Extracellular matrix components of the thymic microenvironment were analyzed with rabbit antisera specific to type-IV collagen, fibronectin, or laminin (Grimaud et al., 1980), purchased from Institute Pasteur (Lyon, France). The distri-

<table>
<thead>
<tr>
<th>Antibody (origin)</th>
<th>Immunogen</th>
<th>Specificity</th>
<th>Recognized TEC&lt;sup&gt;a&lt;/sup&gt; in mice&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>HTK (Biosoft)</td>
<td>Human epidermis</td>
<td>Various CK&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Whole TEC network</td>
</tr>
<tr>
<td>KL4 (Immunotech)</td>
<td>Human epidermis</td>
<td>Various CK</td>
<td>Whole TEC network</td>
</tr>
<tr>
<td>RPN1166 (Amersham)</td>
<td>PtK1 cells (rat canguru)</td>
<td>CK8</td>
<td>Cortical TEC</td>
</tr>
<tr>
<td>RPN1166 (Amersham)</td>
<td>PtK1 cells (rat canguru)</td>
<td>CK18</td>
<td>Cortical TEC</td>
</tr>
<tr>
<td>RPN1165 (Amersham)</td>
<td>SV40-transformed human keratinocyte</td>
<td>CK19</td>
<td>Few medullary TEC</td>
</tr>
<tr>
<td>K,13.1 (Progen)</td>
<td>CK13 isolated from human esophagus</td>
<td>CK13</td>
<td>Not previously tested</td>
</tr>
<tr>
<td>KL1 (Immunotech)</td>
<td>Human epidermis</td>
<td>CK1/CK10</td>
<td>Few medullary TEC</td>
</tr>
</tbody>
</table>

<sup>a</sup>TEC: thymic epithelial cells; CK=cytokeratin.
<sup>b</sup>References: Nicholas et al. (1985); Savino and Dardenne (1988a).
bution patterns of the molecules specifically detected by these antibodies in the human and mouse thymuses were previously reported (Berrih et al., 1985; Lannes-Vieira et al., 1991).

We further used biotinilated or fluoresceinated goat antisera as secondary antibodies. Biotinilated immunesera anti-mouse Ig (GAM/BIOT) and anti-rabbit Ig (GAR/BIOT) were obtained from Amersham, and fluoresceinisothiocyanate-coupled anti-mouse Ig serum (GAM/FITC) and anti-rabbit Ig goat serum bound to tetramethylrhodamineisothiocyanate (GAR/TRITC) were purchased from Biosys (Compiegne, France).

**Immunohistochemistry**

Acetone-fixed, 4-μm thick thymus frozen sections were subjected to indirect immunofluorescence or immunoperoxidase assays according to previous descriptions (Nicolas et al., 1985). Specimens were incubated for 1 hr with a certain primary antibody and washed in PBS. With regard to the immunofluorescence technique, primary antibodies were revealed with appropriate GAM/FITC or GAR/TRITC, incubated for 1 hr. After washing in PBS, slides were mounted in glycerol/PBS and examined through a Leitz Ortoplan fluorescence microscope (Wetzlar, Germany). The rabbit serum pan-specific to cytokeratins and the mouse MAb monospecific to certain cytokeratins were used in double labeling assays.

For streptavidin-biotin immunoperoxidase staining, specimens were subjected to biotinilated secondary antibodies (GAM/BIOT or GAR/BIOT) for 1 hr, washed in PBS, and then exposed to the streptavidin/peroxidase complex (Amersham, Buckinghamshire, UK) for 1 hr. After washing in PBS, peroxidase activity was revealed by a 10-min incubation with aminoethylcarbazole (Sigma Co., Saint Louis, MO) in the presence of H₂O₂. Slides were counterstained with Mayer’s hematoxylin and mounted in gelatin/glycerol/fenol.

**Immunoblot Analysis of Cytokeratins**

The molecular specificity of anti-CK MAb, namely, anti-CK18 and KL1, representatives of different patterns of TEC labeling, were tested on thymus extracts (including those from monkey, hamster, rat, rabbit and sheep) by one-dimen-

sional immunoblot analysis. Extracts enriched in cytokeratins were obtained by a modification of the technique previously described by Woodcock-Mitchell et al. (1982). Thymic fragments of each species were homogenized in glass potters with 25 mM Tris-HCl (pH 7.2) and 1 mM PMSF at 4°C. Supernatants containing mostly thymocytes were discarded and pellets were subjected to the lysis buffer containing 25 mM Tris-HCl (pH 7.2), 1 mM EDTA, 1 mM EGTA, 1 mM PMSF, and 1% Triton x 100, at 4°C. This suspension was centrifuged at 10,000 g for 40 min at 4°C, the supernatant discarded, and the pellet resuspended in 25 mM Tris-HCl (pH 7.2), 1 mM PMSF at 4°C, being then sonicated for 15 min (Branson 2200, Branson Ultrasonics C., Danbury, CN). After one further centrifugation at 10,000 g, the pellet was submitted to a 5-min incubation in Tris-HCl, 1% SDS, 5% 2-mercaptoethanol at 95°C. Extracts were aliquoted and stocked at −20°C.

Protein bands were resolved by SDS-PAGE in 10% acrylamide gels, as originally described by Laemli (1970). Proteins were then blotted onto nitrocellulose sheets (Schleicher and Schuell, Dassel, Germany) according to Towbin et al. (1979). Transfer was checked with Ponceau S Red, and nonspecific binding sites were saturated by 1-hr horizontal agitation with PBS containing 10% fetal calf serum (FCS) and 0.05% Tween 20, at room temperature. Bands were then incubated overnight within a certain anti-CK solution at 4°C. The anti-CK18 was diluted 1:10 in saturation solution and KL1 was diluted 1:200. After washing in PBS/FCS/Tween for 30 min, nitrocellulose sheets were incubated with GAM/BIOT (1:500) for 2 hr, washed, and finally incubated in Streptavidin-peroxidase complex (1:500) for 2 hr. After a further washing, enzyme activity was demonstrated with 3',3-diaminobenzidine (0.3 mg/ml Tris-HCl, pH 7.6) in the presence of 0.03% H₂O₂.

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