Characterization of Human Thymic Lymphocytes Forming Rosettes with Stromal Cells

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NONOYAMA, S., NAKAYAMA, M., ABE, J., KOHSAKA, T., KOBAYASHI, N. and YATA, J. Characterization of Human Thymic Lymphocytes Forming Rosettes with Stromal Cells. Tohoku J. Exp. Med., 1992, 168 (3), 467-474 — The interaction of thymic lymphocytes and stromal cells is believed to be important for T cell development in thymus. In this study, thymic rosettes (TR), which are cell-cell complexes of thymic lymphocytes and stromal cells, were isolated from human thymic tissue, and were characterized. Treating human thymus with collagenase in mild condition, human TR were successfully isolated. Subsequently, TR were purified by the 1G sedimentation method. Human TR consisted of a stromal cell in center surrounded by lymphocytes. The stromal cells were positive for CD14, CD11b, and HLA-DR but negative for thymic epithelial cell specific mAb, UH-1, suggesting that they are macrophage/dendritic cells. The lymphocytes which formed TR (TRL) were mainly double positive (CD4+CD8+) and CD1+ cells, and few of them expressed bright CD3, indicating that TRL are in the intermediate maturation stage. TRL expressed activation markers (Tal and HLA-DR) in a significantly higher percentage of cells than did unselected thymocytes. Blocking test revealed that CD11a and CD2 are involved in the binding of TRL and the stromal cells as adhesion molecules. —— adhesion molecule; T cell differentiation; human thymic rosettes

Stromal cells in the thymus are known to play important roles in T cell differentiation. They produce several growth factors, which are essential for immature T cells to proliferate and differentiate into mature T cells (Le et al. 1987; Murray et al. 1989). In addition, immature T cells are known to bind to stromal cells in vitro, and thymic T cells are known to be activated through the binding process (Denning et al. 1987). It has been also reported that the self antigens are presented by stromal cells (Lorenz and Allen 1989), suggesting that recognition of self antigen by T cells is educated by stromal cells. Thus, binding to stromal cells is considered to be an important process for immature T cells to

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differentiate into mature T cells.

The actual cell-cell binding in vivo has been demonstrated by isolating cell
complex between stromal cells and thymic lymphocytes (thymic rosette (TR))
(Kyewski et al. 1982) from murine thymi. These complexes are considered to
represent a in vivo thymic microenvironment, in which immature T cells commu-
nicate with stromal cells.

However, the similar cell-cell complexes have not been demonstrated in
human thymus. This is probably because a method to isolate them has not been
established yet. The method used in murine system fails to isolate human TR.
In this study, we modified the method and successfully isolated human TR. We
report the characterization of human thymic rosettes.

MATERIALS AND METHODS

Isolation of human TR

Human thymic tissues removed from children during cardiovascular surgery for congeni-
tal heart disease were used. The thymus tissue was gently teased, and digested by 100 U/
ml of collagenase (Boeringer-Mannheim, FRG) at 4°C for 30 min. Thus obtained cells
contained TR. TR were purified by the 1G sedimentation method described elsewhere
(Wekerle et al. 1980). Repeating the 1G sedimentation method five times, TR were
obtained in high purity.

Surface marker analysis

Surface marker expression of TRL was analyzed by flowcytometer. TR were separated
into free lymphoid cells (thymic rosette lymphocytes, TRL) and stromal cells by gently
pipetting them in phosphate buffered saline (PBS) at 37°C. The contamination of
lymphocytes other than TRL was less than 1% at this period. Cells were stained by the
direct immunofluorescence method, using the following mAb: anti-T3-FITC (CD3), anti-
T4-PE (CD4), anti-T8-FITC (CD8), anti-I2-FITC (anti-HLA-DR), and anti-Ta1-PE
(CD26). All mAbs were purchased from Coulter (Hialeah, FL, USA). The stained cells
were analyzed with an EPICS 753 flowcytometer (Coulter). The cells to be analyzed were
gated only for the lymphoid cells.

To analyze surface marker expression of stromal cells, immunomagnetic beads were
used. Human TR and monocytes were cultured with human IgG at 4° for 15 min to block
Fc receptor. Monocytes were obtained from human PBMC by adhesion to plastic wall.
Cells were then cultured with various kind of mAb at 4° for 15 min. After washing with
PBS containing 1% FCS, cells were incubated with immunomagnetic beads coated with
sheep anti mouse IgG (Dynal, Great Neck, NY, USA) at 4° for 15 min. Numbers of the
beads attached to the cells were counted under a microscope. If more than three beads were
attached, the cell was considered to be positive for the cultured mAb. Twenty human TR
were counted for each mAb. B1 (CD20), My4 (CD14), and Mo1 (CD11b) monoclonal
antibodies were purchased from Coulter. The anti-UH-1 IgG mAb, specific for human
thymic epithelial cells, was kindly provided by Dr. K. Hirokawa (Tokyo Metropolitan
Institute of Gerontology).

Blocking test

TR were separated into free lymphoid cells (thymic rosette lymphocytes, TRL) and
stromal cells by gently pipetting them in phosphate buffered saline (PBS) at 37°C. The
blocking test was then performed as described previously (Nonoyama et al. 1989). Briefly,
the separated cells were cultured in hanging drop in 20 μl of RPMI 1640 medium sup-
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implemented with 10% FCS for 10 min at 37°C. Binding between TRL and stromal cells were reconstituted during this culture. The blocking test was performed by adding various kind of mAbs into the hanging drop culture. MAbs used are as follows: anti-CD11a (Sanbio, Uden, The Netherlands), Anti-CD2 (Coulter, Hialeah, FL, USA), and anti-TCR α/β (Becton Dickinson, Mountain View, CA, USA). The concentration of mAbs were all 1 μg/ml. Rosettes which contain more than 3 lymphocytes were counted as positive. The percentage of conjugate forming cells was calculated as:

\[
\frac{\text{The number of reconstituted TR}}{\text{The number of original TR}} \times 100
\]

RESULTS

Isolation of human thymic rosettes

Human TR could be isolated by collagenase treatment in a mild culture condition of 4°C 30 min. The condition used for isolation of murine TR, that is treating thymic tissue with collagenase and dispase at 37°C for 30 min, was found to destroy human TR. TR were further purified by the 1G gradient method. After repeating 1G gradient method, contaminating lymphocytes which did not form rosettes decreased to be less than 1% of the entire rosette-forming population. Fig. 1 shows the typical shape of human TR, which consist of a stromal cell in center surrounded by several small lymphocytes.

Surface marker analysis of TRL and stromal cells

The purified TR were then separated into free cells by pipetting them in PBS at 37°C. They were then collected and analyzed for surface phenotype by flowcytometry (Table 1). The stromal cells were gated out on the flowcytometry

Fig. 1. Human thymic rosettes isolated by collagenase treatment and purified by the 1G sedimentation method.
As shown in Table 1, the majority of TR forming lymphocytes (TRL) were CD4+ CD8+ (DP). The percentage of this subset (90.4% ± 3.8%, mean ± S.D.) in TRL is significantly higher than that (80.8% ± 6.2%) in unselected thymocytes (p < 0.05, student t-test). Most of human TRL also expressed CD1, which recognize cortical thymocytes. The percentage of CD1 positive cells in TRL was significantly higher than that observed in unselected thymocytes (p < 0.05). In contrast, CD3 bright+ subset in TRL was only 4.0%, which was significantly less than that in unselected thymocytes. These results indicate that TRL are mainly in an intermediate maturational stage.

HLA-DR is an activation marker for T cells (Yu et al. 1980). The percentage of HLA-DR+ cells in TRL (58.5% ± 10.6%) is significantly higher than that in unselected thymocytes (p < 0.05, student t-test). Another activation marker, Tal (CD26) (Fox et al. 1984) was also highly expressed in TRL. The percentage of Tal+ cells in TRL (87.0% ± 3.6%) is again significantly higher than that in unselected thymocytes (74.0% ± 5.5%) (p < 0.05). The typical results of HLA-DR and Tal expression are shown in Fig. 2.

Surface marker expression on stromal cells in TR was examined using mAb and immunomagnetic beads. Stromal cells were positive for CD14, CD11b but negative for CD20 and UH-1, specific for human thymic epithelial cells. This result indicates that stromal cells in TR were macrophage/dendritic cell origin.

### Table 1. Cell surface phenotypes of thymic rosette forming lymphocytes (TRL) and unselected thymocytes (UST)

<table>
<thead>
<tr>
<th></th>
<th>CD4−CD8−</th>
<th>CD4+CD8+</th>
<th>CD4+CD8−</th>
<th>CD4−CD8+</th>
</tr>
</thead>
<tbody>
<tr>
<td>TRL</td>
<td>0.3±0.2*</td>
<td>90.4±3.8</td>
<td>5.6±2.7</td>
<td>2.9±1.4</td>
</tr>
<tr>
<td>UST</td>
<td>1.4±0.5</td>
<td>80.8±6.2</td>
<td>11.9±3.0</td>
<td>5.5±2.5</td>
</tr>
</tbody>
</table>

*The data are shown as a mean(%)±s.d. of seven independent experiments.

*p < 0.05.

### Table 2. Surface phenotypes of stromal cells in TR

<table>
<thead>
<tr>
<th>Antibody</th>
<th>My4</th>
<th>Mo1</th>
<th>B1</th>
<th>UH-1</th>
<th>Mouse IgG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stromal cells in TR</td>
<td>+</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Monocytes*</td>
<td>+</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

*Monocytes were purified from human PBMC by adhesion to plastic wall.
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Blocking test

The results of the blocking test are shown in Fig. 3. TR were first separated into free thymocytes and stromal cells by pipetting them in PBS. The contamination of lymphocytes which did not form TR was less than 1%. After the short-time culture in hanging drop system (upside down), TRL bound to stromal cells and formed rosettes again. This reconstitution of TR was partially inhibited by either anti-CD2 mAb or anti-CD11a mAb ($p < 0.05$, $t$-test). Almost complete inhibition was achieved by the combination of anti-CD2-mAb and

Fig. 2. The expression of HLA-DR and Tal on thymic rosette forming lymphocytes (TRL) and unselected thymocytes (UST). X-axis, fluorescence intensity; y-axis, cell numbers.

Fig. 3. Results of the blocking test. "% conjugate forming cells" indicates the percentage of conjugate forming cells after reconstitution of thymic rosettes in vitro.

Blocking test

The results of the blocking test are shown in Fig. 3. TR were first separated into free thymocytes and stromal cells by pipetting them in PBS. The contamination of lymphocytes which did not form TR was less than 1%. After the short-time culture in hanging drop system (upside down), TRL bound to stromal cells and formed rosettes again. This reconstitution of TR was partially inhibited by either anti-CD2 mAb or anti-CD11a mAb ($p < 0.05$, $t$-test). Almost complete inhibition was achieved by the combination of anti-CD2-mAb and
anti-CD11a mAb. Anti-TCR $\alpha/\beta$ mAb did not affect the binding in this system.

**Discussion**

This report demonstrates that the TR, a cell-cell complex of thymocytes and stromal cell, is present in human thymus as in murine thymus. Modifying the method used for the isolation of murine TR, human TR was successfully isolated. The morphology of human TR is similar to that of murine TR (Kyewski et al. 1982), which consists of a stromal cell surrounded by several lymphocytes. A possibility that human TR is formed during the preparation process remains. However, the observation by Kyewski and co-workers that thymocytes of each TR were either Thy1.1 or Thy1.2 even if Thy1.1 thymus and Thy1.2 thymus were co-digested to isolate TR (Kyewski et al. 1982) suggests that TR preexist in vivo and were not formed during isolation.

Cell surface phenotyping indicated that 90% of TRL were CD4+CD8+ (double positive) cells, while 80% of unselected thymocytes were CD4+CD8+. Significantly lower percentage of TRL were double negative (immature) cells or single positive (mature) cells ($p < 0.05$). The results using anti-CD3 and anti-CD1 mAb were consistent with this observation. Significantly higher percentage of TRL expressed CD1+, which reacts with cortical thymocytes, than unselected thymocytes. Mature CD3 bright+ cells were rarely found in TRL. This result shows the similarity of human TRL to murine TRL, which co-express CD4 and CD8, and have low density T cell receptor. It is suggested that TRL are in intermediate stage between intrathymic precursor cells and mature medullary-type thymocytes.

We previously reported that binding to thymic epithelial cells is limited to the thymocytes in an intermediate stage (Nonoyama et al. 1989). Since the stromal cells in TR are macrophage/dendritic cells (Kyewski et al. 1987), the present study demonstrates that the binding of thymocytes to macrophage/dendritic cells also occurs in an intermediate maturation stage. These results indicate that thymocytes in intermediate stage have high affinity to two types of thymic stromal cells, which are thymic epithelial cells and macrophage/dendritic cells.

TRL expressed activation antigens, HLA-DR and Ta1, in a significantly higher percentage than unseparated thymocytes. This higher HLA-DR expression is not due to the difference of maturation stage between TRL and UST, since the expression of HLA-DR on DP cells is less than that on the other stage of thymocytes (Nonoyama 1989). This result is consistent with the observation that murine TRL are activated and express activation markers (Kyewski et al. 1984). Since macrophage/dendritic cells activate peripheral T cells after making cell complexes in vitro (Langhoff and Steinman 1989), it is likely that macrophage/dendritic cells in TR cause activation of the TRL.

CD11a ($\alpha$ chain of LFA-1) and CD2 molecules were determined to be the
adhesion molecules used for the formation of TR. These two molecules are also
known to be the adhesion molecules for the binding of thymocytes and thymic
epithelial cells (Denning et al. 1987; Nonoyama et al. 1989). The data presented
here further confirm the importance of CD11a and CD2 for the interaction of
thymocytes and stromal cells. In addition, since these adhesion molecules are
also known to conduct an activation signal into cells (Denning et al. 1987; Noesel
et al. 1988), it is possible that these molecules play some roles in the activation of
TRL.

These findings demonstrate the interaction of T cells and stromal cells in
thymus. Distinct phenotype of TRL suggests that interaction between thymic T
cells and stromal cells occurs in a particular stage of T cells differentiation.

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