Interleukin-2 Receptor (Tac Antigen) Expressed on Adult T Cell Leukemia Cells

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Abstract

We studied the expression of the interleukin-2 (IL-2) receptor and the proliferative response to exogenous IL-2 of peripheral blood leukemic cells from patients with adult T cell leukemia (ATL) in order to see whether IL-2 receptor expressed on ATL cells is different from normal IL-2 receptor and whether it plays a role in the neoplastic growth in ATL.

Peripheral blood leukemic cells from 42 patients with ATL examined expressed IL-2 receptors that were detected by anti-Tac monoclonal antibody when examined immediately after the separation of cells or after the culture for 24 or 48 h. The number of anti-Tac binding sites ranged from 3,100 to 11,400 in fresh cells and from 3,600 to 96,000/cell in short-term cultured leukemic cells, whereas phytohemagglutinin-P (PHA-P)–stimulated normal T cells exhibited 6,900–35,000 anti-Tac binding sites per cell. ATL-derived and human T cell leukemia/lymphoma virus, type I (HTLV-I)–infected cell lines such as MT-1 and Hut102 expressed a much higher number of anti-Tac binding sites.

Leukemic cells from 15 patients with ATL examined showed no or very poor proliferative response to various concentrations of immunopurified IL-2, although they expressed Tac antigen (Ag). Radiolabeled IL-2 binding experiments demonstrated that ATL leukemic cells could bind IL-2, and they expressed both high and low affinity IL-2 receptors, although the number of high affinity IL-2 receptor was much less than that of low affinity IL-2 receptor and that of anti-Tac binding sites.

In contrast, leukemic T cells from a patient with T cell chronic lymphocytic leukemia (CLL), in whom HTLV-I infection was not demonstrated, responded as well as PHA-P–stimulated normal T cells, and their IL-2 receptors, unlike ATL cells, were modulated (down regulated) by anti-Tac antibody.

No differences were noted between ATL cells and normal activated T cells in one-dimensional sodium dodecyl sulfate polyacrylamide gel electrophoresis analysis of the IL-2 receptor. Thus, leukemic cells in ATL spontaneously and continuously express IL-2 receptor, which appears to be abnormally regulated and unresponsive to IL-2. These results, taken together with those on normal IL-2 receptors on HTLV-I–negative T-CLL cells, suggest that abnormal expression of the IL-2 receptor in ATL is closely associated with HTLV-I infection and may play a role in the neoplastic growth of ATL cells.

Introduction

Adult T cell leukemia (ATL), which has characteristic clinical and hematologic features and is endemic southwest of Japan (1–3), is a leukemia of peripheral, mature, and OKT4(+) T cells (4). Studies subsequent to the discovery and isolation of a novel human retrovirus, human T cell leukemia/lymphoma virus, type I (HTLV-I) (5, 6), have demonstrated a close association between HTLV-I and the leukemogenesis of ATL (7–10), although the mechanism of the cell transformation that may be induced by HTLV-I infection remains unclear.

We previously reported that leukemic cells in ATL are derived from peripheral mature T cells with a helper/inducer subset phenotype and express Tac antigen (Ag) (interleukin-2 [IL-2] receptor) (4) that is recognized by anti-Tac monoclonal antibody (11–13). Popovic et al. (14) also noted the strong expression of Tac Ag on cord blood cells that were transformed by co-culturing with HTLV-producing cell lines. A recent report by Waldmann et al. also showed that HTLV-positive ATL cells, but not HTLV-negative Sézary cells, express IL-2 receptors (15). In addition, we found that Tac Ag on ATL cells, unlike those on normal activated T cells, is not modulated by anti-Tac antibody (16), which suggests the abnormal regulation of Tac Ag expression in ATL cells. The proliferation of normal peripheral T cells initiated by antigen or lectin is mediated by IL-2, which is produced by activated T cells and binds to inducible IL-2 receptors. Further characterization of the IL-2 receptor on ATL cells is needed to understand whether it is merely a marker of ATL cells or if it is involved in their neoplastic growth.

In the present study, we examined the molecular size and IL-2 binding activity of IL-2 receptors expressed on peripheral blood leukemic cells from patients with ATL, and the proliferative response of ATL cells to exogenous IL-2, comparing those of normal activated T cells and HTLV-I negative leukemic T cells.

We will show that peripheral blood leukemic T cells in ATL respond very poorly to exogenous IL-2, although they express Tac antigen of similar molecular weight to that of normal activated T cells and can bind IL-2. In contrast, leukemic T cells from a patient with chronic lymphocytic leukemia.

1. Abbreviations used in this paper: ADF, ATL-derived factor; Ag, antigen, ATL, adult T cell leukemia; FCS, fetal calf serum; HTLV-I, human T cell leukemia/lymphoma virus (type I); IL-2, interleukin-2; PHA-P, phytohemagglutinin-P; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; T-CLL, T cell chronic lymphocytic leukemia.
leukemia which were negative for HTLV-I provirus integration, expressed Tac antigen and showed a good proliferative response to IL-2.

Methods

Patient population. 42 patients with ATL and one patient with chronic lymphocytic leukemia of T cell origin were studied. The age of ATL patients ranged from 24 to 76, and 25 of the 42 patients were male. The diagnosis of ATL was made on the basis of clinical features (2), hematologic characteristics (2), serum antibodies to ATL-associated antigens (6) in all cases, and the HTLV provirus integration in DNA of leukemic cells in seven cases. Peripheral white blood cell count ranged from 15,900 to 249,000/mm³.

The patient with T cell chronic lymphocytic leukemia (T-CLL) was a 54-yr-old male with generalized lymph node enlargement and a high white blood cell count (range, 29,200-100,000 cells/mm³). Peripheral blood smears showed lymphoid cells with nonconvoluted nuclei.

Cell separation and culture. Peripheral blood mononuclear cells were separated from heparinized blood by Ficoll-Hypaque density gradient centrifugation. Both fresh leukemic cells and frozen cells, which had been stored in liquid nitrogen, were used. The recovered cells with good viability (>85%) were used. More than 80% of the cells in the cell suspension examined were leukemic cells as determined by the morphological characteristics of cells on the May-Giemsa-stained slides.

Leukemic cells cultured from 1 to 7 d in RPMI 1640 medium containing 10% fetal calf serum (FCS) (M.A. Bioproducts, Walkersville, MD) or autologous serum or plasma were also examined for the expression and IL-2 binding of IL-2 receptor. Normal peripheral blood mononuclear cells cultured with 0.1% phosphomannaglutinin-P (PHA-P) (Difco Laboratories, Detroit, MI) for 2–7 d were used for the quantitation of Tac antigen and for IL-2 binding assay. ATL-derived and HTLV-I-infected cell lines, MT-1 (17) and Hut102 (18), which were maintained with RPMI 1640 medium (Nissui Pharmaceutical Co., Ltd., Tokyo, Japan) containing 10% FCS, were also used.

Detection of IL-2 receptor by immunofluorescence. Anti-Tac monoclonal antibody that recognizes human IL-2 receptor (13) was used to detect IL-2 receptor. Its production and characterization were described elsewhere (11, 13). One-million cells were incubated with a saturating amount of anti-Tac antibody at 4°C for 30 min, washed twice with Hanks' balanced salt solution containing 0.1% bovine serum albumin and 0.1% sodium azide, and then incubated with fluorescein isothiocyanate-conjugated F(ab)₂ anti-mouse IgG (Cappel Laboratories, Cochranville, PA) at 4°C for 30 min. Control staining was performed with normal BALB/c mouse serum instead of anti-Tac antibody. Washed and resuspended cells were observed under a fluorescence microscope, or subjected to flow cytometric analysis using a FACS analyzer (Becton-Dickinson Co., FACS Systems, Sunnyvale, CA) or a Spectrum III (Ortho Diagnostic Systems, Westwood, MA).

Radiolabeled anti-Tac binding assay. Anti-Tac antibody purified by gel and DEAE cellulose chromatography from hybridoma ascites was radiolabeled by the chloramine T method (specific activity 5,000–16,000 cpm-ng). Bindable fraction of radiolabeled anti-Tac was ~90% of all activity when determined by the binding of radiolabeled anti-Tac to an excess number of cells. 50 μl of serial dilutions of [125I]-anti-Tac, 50 μl of medium, and 400 μl of cell suspension (1.0–4.0 × 10⁶/ml) in RPMI 1640 medium containing 2% FCS were mixed and incubated on ice for 30 min. After washing and resuspending cells in 500 μl of medium, 200 μl of the cell suspension in duplicate was centrifuged through a 150-μl layer of a mixture of 20% olive oil (Nakarai Chemicals, Kyoto, Japan) and 80% Di-n-butyl phthalate (Nakarai Chemicals). The tips of the tubes containing the cell pellet were cut off and the radioactivity was counted in a gamma counter. Nonspecific binding was estimated by incubating cells with both radiolabeled anti-Tac and >1,000-fold excess amount of unlabeled anti-Tac. Nonspecific binding was <3.7% of total activities bound to cells in these experiments. Specific binding was obtained by subtracting nonspecific binding. To obtain accurate values of bound/free antibody, the values were corrected for bindable proportion of total radioactivity. Scatchard analysis of binding assay was performed by plotting values of bound/free vs. bound anti-Tac antibody.

Radiolabeled IL-2 binding assay. Escherichia coli-derived recombinant IL-2 was kindly provided by Takeda Chemical Industries Inc. (Osaka, Japan). The purity of recombinant IL-2 was >99.8%, and the biological activity which was comparable to Jurkat cell line-derived immunoaffinity-purified IL-2 was determined by the proliferation assay using IL-2-dependent cell lines and PHA-P-stimulated normal peripheral blood lymphocytes (data not shown). IL-2 was radiolabeled using 125I-Bolton and Hunter's reagents (B&H reagent, 2,000 Ci/mmol, moniodinated, NEX-120, New England Nuclear, Boston, MA). Specific radioactivity of labeled IL-2 was 2,000–5,000 cpm/ng. Effective radioactivity bindable to cells was ~90% of the total activity.

Radiolabeled IL-2 binding assay was performed according to the method described by Robb et al. (19, 20) with a slight modification. Cells were incubated at 37°C in IL-2-free fresh RPMI 1640 medium twice for 60 min to promote dissociation and/or degradation of endogenously bound IL-2. Serial dilutions of radiolabeled IL-2 and 2–5 × 10⁶ cells in a total volume of 0.5 ml of RPMI 1640 medium containing 25 mM Hepes, pH 7.2, and 10 mg/ml of bovine serum albumin (Armour Pharmaceutical Co., Kankakee, IL) were incubated at 37°C for 60 min. After incubating, washing, and resuspending cells, 200 μl of cell suspension in duplicate was centrifuged through a 150-μl layer of a mixture of 20% olive oil and 80% Di-n-butyl phthalate. The tips of the tubes containing the cell pellet were cut off and the radioactivity was counted in a gamma counter. Nonsaturable binding was determined by incubating cells with 250-fold excess amount of unlabeled IL-2. Specific binding was obtained by subtracting nonsaturable binding. The number of IL-2 binding sites and the affinity was estimated by Scatchard analysis.

Proliferative response to IL-2. Normal peripheral blood mononuclear cells cultured with 0.1% PHA-P for 7 d and peripheral leukemic lymphocytes were studied for the proliferative response to IL-2. One-tenth of a million cells were placed in a 96-well microtiter plate (Falcon Labware, Oxnard, CA) and cultured in 200 μl of RPMI 1640 medium containing 10% FCS and 30 μg/ml of gentamicin in the presence of various concentrations of immunooaffinity-purified IL-2 (reference 19, generously provided by Dr. K. Smith; 1 U biological activity = 8.2 ng protein) at 37°C in a humid atmosphere with 5% CO₂ for 72 h. The cultured cells were pulsed with [³H]thymidine (2 Ci/ml, Amersham Corp., Arlington Heights, IL) for the last 6 h, followed by precipitation onto glass fiber filters, and the radioactivity was counted by liquid scintillation. MT-1 cells were cultured both at 5 × 10⁶/ml and 2.5 × 10⁷/ml in a 96-well plate in the presence of IL-2.

Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). Tac Ag immunoprecipitated by anti-Tac antibody from labeled cell lysates was analyzed by one-dimensional SDS-PAGE according to the method described in detail previously (21). In brief, leukemic cells from ATL patients (viability >90%) were surface labeled with [125I]Na (New England Nuclear) by the lactoperoxidase method, washed and lysed in a cell extraction buffer (0.5% Nonidet P-40, 10 mM Tris-HCl, 0.15 M NaCl, 1 mM phenylmethylsulfonyl fluoride, pH 7.2). Tac antigen was immunoprecipitated from labeled cell lysates by incubation with anti-Tac antibody and staphylococcal protein A-coupled Sepharose 4B (Pharmacia Fine Chemicals, Piscataway, NJ). The immunoprecipitate was electrophoresed on 7.5% polyacrylamide gel and visualized by autoradiography.

Results

Expression of IL-2 receptor (Tac Ag) on ATL leukemic cells. Peripheral blood leukemic cells from 32 of the 42 patients with ATL examined expressed IL-2 receptors (Tac Ag) on
their cell surface when stained with anti-Tac antibody and examined immediately after separation of cells. 9–60% of the fresh cells examined were reactive with anti-Tac antibody in these 32 patients. Less than 5% of the fresh cells from the remaining 10 patients were reactive with anti-Tac antibody. Leukemic cells from these 10 patients, however, expressed the receptor when examined after 24 or 48 h culture in medium containing 10% FCS or autologous serum or plasma. 20–60% of short-term cultured leukemic cells were reactive with anti-Tac antibody. Leukemic cells cultured with medium generally expressed more IL-2 receptors on their cell surface than did fresh cells (Fig. 1). In case M.S., the number of anti-Tac binding sites was 5,500/cell on fresh leukemic cells and 96,000/cell on leukemic cells cultured for 5 d. In other words, 17-fold increase in the number of Tac Ag sites was detected after 5 d of culture. Cytofluorometric studies performed at the same time also revealed a marked increase of fluorescence intensity (Fig. 1).

In contrast, fresh peripheral blood lymphocytes from 20 normal individuals did not express IL-2 receptor (<2% of the cells were reactive with anti-Tac). Furthermore, only a small proportion (2.9–5.1%) of normal lymphocytes cultured with medium containing 10% FCS for 48 h expressed Tac antigen as detected by immunofluorescence method. Radiolabeled anti-Tac binding assay of short-term (48 h) cultured normal peripheral blood lymphocytes demonstrated 500–890 (mean 730, n = 4) anti-Tac binding sites per cell, which were much less than those of short-term cultured leukemic cells from ATL patients (Table I).

**Radiolabeled anti-Tac antibody binding.** The specificity of radiolabeled anti-Tac antibody binding was studied by the binding inhibition in the presence of an excess amount of unlabeled anti-Tac, IL-2, OKT3 antibody, and OKIa1 antibody. More than 96% of radiolabeled anti-Tac binding to PHA-P-stimulated normal lymphocytes or Hut102 cells was inhibited by the presence of >1,000-fold excess amount of unlabeled anti-Tac. In addition, ~98% of radiolabeled anti-Tac binding was inhibited by excess IL-2. OKT3 and OKIa1 antibodies, however, did not inhibit (<5%) radiolabeled anti-Tac binding. The reversibility of radiolabeled anti-Tac binding was tested by adding an excess amount of unlabeled anti-Tac to Hut102 cells that had been incubated by radiolabeled anti-Tac. 58% of the bound anti-Tac was released after 120 min incubation at 37°C.

Fig. 2 shows a typical binding curve and a Scatchard plot of radiolabeled anti-Tac binding study of 5 d cultured ATL leukemic cells. The number of anti-Tac binding sites per cell on leukemic cells from ATL patients, HTLV-I–infected cell lines, PHA-P–stimulated T cells, and leukemic cells from a patient with T-CLL is shown in Table I. In patient M.S., leukemic cells were examined for anti-Tac binding on several occasions in his clinical course. HTLV-I–infected cell lines, MT-1 and Hut102, expressed a high number of Tac Ag sites (560,000 and 410,000 sites/cell, respectively). Fresh peripheral blood leukemic cells from ATL patients expressed 3,100–11,400/cell anti-Tac binding sites. More Tac Ag molecules were detected when leukemic cells were examined after culture for 5 or 6 d, although the increase varied in each case. Peripheral blood T cells usually expressed the maximal number of anti-Tac binding sites after 2 to 3 d of activation with PHA-P, which declined thereafter. In seven normal subjects, 6,900–35,000 Tac Ag sites per cell were detected on T cells cultured in the presence of PHA-P for 2–7 d. The affinity of anti-Tac binding in ATL cells and HTLV-I–infected cell lines (K_D = 4–7 x 10^7 M^-1) was not different from that of PHA-P–stimulated normal lymphocytes.

Peripheral blood leukemic T cells from a patient with T-CLL expressed 2,000 anti-Tac binding sites per cell. HTLV-I provirus integration in DNA of leukemic cells was not demonstrated by the Southern blot hybridization method (Fukui, K., and T. Honjo, unpublished data) and no serum antibody to ATL-associated antigens was detected in this patient (data not shown).

**Proliferative response of leukemic cells from ATL patients to IL-2.** Peripheral blood T cells from 10 normal individuals who had been cultured for 7 d in the presence of 0.1% PHA-P responded well to added immunofluorinity-purified IL-2 (Fig. 3). On the contrary, fresh leukemic cells from 11 ATL patients and cryopreserved leukemic cells from 4 ATL patients which expressed Tac Ag did not respond well to various concentrations of IL-2. The stimulation index ([^{3}H]thymidine uptake cultured with IL-2/[^{3}H]thymidine uptake cultured with medium alone) at an IL-2 concentration of 0.5 U/ml ranged from 0.2 to 6.5 in ATL leukemic cells and from 75 to 806 in PHA-P–stimulated normal T cells. MT-1 cells established from an ATL patient (17) did not respond to IL-2 in spite of the markedly enhanced expression of Tac Ag (Fig. 4, Table I). Leukemic cells from a patient with T-CLL, on the other hand, showed a good proliferative response to IL-2, which was comparable with that of PHA-P–stimulated normal T cells (Fig. 4).

We also studied ATL cells cultured for 3–7 d, which expressed more IL-2 receptors than fresh leukemic cells in order to assess whether their poor proliferative response of ATL leukemic cells were due to an inadequate number of IL-2 receptors on fresh leukemic cells. Cultured leukemic cells,
**Table I. Number of Anti-Tac and IL-2 Binding Sites**

<table>
<thead>
<tr>
<th></th>
<th>Anti-Tac (sites per cell)</th>
<th>IL-2 (sites per cell)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh</td>
<td>Cultured</td>
</tr>
<tr>
<td>ATL*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Case M.S.</td>
<td>5,500</td>
<td>96,000</td>
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<td></td>
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<td></td>
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<tr>
<td></td>
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</tr>
<tr>
<td>S.M.</td>
<td>4,400</td>
<td>34,000</td>
</tr>
<tr>
<td>F.K.</td>
<td>5,100</td>
<td>6,100</td>
</tr>
<tr>
<td>Y.O.</td>
<td>11,400</td>
<td>15,000</td>
</tr>
<tr>
<td>K.U.</td>
<td>3,100</td>
<td>3,600</td>
</tr>
<tr>
<td>HTLV-I-infected</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MT-1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T-CLL (case O.O.)</td>
<td>2,000</td>
<td>410,000</td>
</tr>
</tbody>
</table>

Normal lymphocytes

- Cultured with medium for:
  - 2 d (mean, n = 4)
  - Stimulated by PHA-P for:
    - 2 d (mean, n = 2)
    - 3 d (mean, n = 3)
    - 7 d (mean, n = 2)

The number of anti-Tac and IL-2 binding sites were determined by Scatchard analysis of radiolabeled anti-Tac and radiolabeled IL-2 binding experiments. *Peripheral blood leukemic cells from ATL patients were examined both after separated from blood or recovered from fresh-frozen state and after cultured for 3–6 d.

however, still showed a poor response to exogenous IL-2 even after the number of cell surface anti-Tac binding sites increased (data not shown). In addition, peripheral blood leukemic cells from three patients with ATL who were examined did not respond to IL-2 even at a concentration of 32 U/ml (two patients) or 130 U/ml (one patient).

**Radiolabeled IL-2 binding.** Unstimulated normal peripheral blood lymphocytes and Tac-negative T or B cell lines did not bind significant amount of radiolabeled IL-2 (data not shown). As radiolabeled anti-Tac binding, the reversibility of radiola-

![Figure 2. Radiolabeled anti-Tac binding to ATL leukemic cells cultured for 5 d. The data shown were obtained by subtracting nonspecific binding, which was estimated by incubating cells with an excess amount of unlabeled anti-Tac. The result was transformed into a Scatchard plot which is shown in an inset. The calculated number of binding sites was 34,000/cell and \( K_d = 6.6 \times 10^8 \text{ M}^{-1} \) in this experiment.](image)

![Figure 3. Proliferative response to IL-2 of peripheral blood leukemic cells from 15 patients with ATL. Leukemic cells were cultured with various concentrations of immunoaffinity-purified IL-2 for 72 h and [\(^3\)H]thymidine uptake was determined. Each point represents mean of triplicate culture. Proliferative response to IL-2 of peripheral blood lymphocytes from 10 normal individuals cultured with 0.1% PHA-P for 7 d is shown as the shaded area. The leukemic cells from 15 ATL patients examined showed no or very poor proliferative response to IL-2 as compared with PHA-P-stimulated normal lymphocytes. TdR, thymidine.](image)
beled IL-2 binding was studied. The reversible proportion of the binding was \( \approx 90\%\) of the total activity bound to the cells.

In order to determine whether no or very poor proliferative response of ATL leukemic cells to exogenous IL-2 was due to poor binding of IL-2 to the IL-2 receptor, we tested the binding of radiolabeled IL-2 to leukemic cells. As shown in Figs. 5 and 6, radiolabeled IL-2 binding assay and its Scatchard analysis demonstrated IL-2 receptors with two different affinities in PHA-P–stimulated normal peripheral lymphocytes, HTLV-I–infected cell lines, and leukemic cells from ATL patients. Peripheral blood lymphocytes stimulated by PHA-P for 2–7 d exhibited 1,300–3,500 IL-2 receptors/cell with high affinity \( K_d = 2.6–6.9 \times 10^8 \text{M}^{-1}\) and 8,000–25,500 receptors/cell with low affinity \( K_d = 3.7–8.8 \times 10^8 \text{M}^{-1}\). In HuT102 cells, the majority of IL-2 receptors were low affinity receptors. In ATL case M.S., fresh leukemic cells expressed 250/cell high and 2,250/cell low affinity IL-2 receptors. The leukemic cells cultured with medium for 5 d expressed 2,100/cell high and 37,500/cell low affinity IL-2 receptors, the total of which was roughly equal to the number of anti-Tac binding sites (34,000/cell).

Radiolabeled IL-2 binding assay of both fresh and short-term cultured leukemic cells from another ATL patient (case S.Y.) also showed both low and high affinity IL-2 receptors (data not shown), as in case M.S. No major differences were noted in the affinity of two classes of IL-2 receptors between ATL leukemic cells \( K_d = 1.1 \times 10^8 \text{M}^{-1}\), 9.5 \( \times 10^8 \text{M}^{-1}\), HuT102 cells, and PHA-P–stimulated normal lymphocytes.

These radiolabeled IL-2 binding experiments showed that both ATL leukemic cells and HTLV-1–infected cell lines expressed comparable number of high affinity IL-2 receptors to that of PHA-P–stimulated normal lymphocytes, which could bind IL-2 at the concentrations used in the proliferative response to IL-2, and also expressed a large number of low affinity IL-2 receptors.

**Figure 4.** Proliferative response to IL-2 of HTLV-I–uninfected leukemic cells from a patient with T-CLL and ATL-derived, HTLV-I–infected MT-1 cell line. Peripheral blood leukemic cells from the T-CLL patient and MT-1 cells were cultured with various concentrations of IL-2 for 72 h and \(^{3}H\)thymidine uptake was determined. Each point represents mean of triplicate culture. The result shown is a representative of three different experiments. Leukemic cells from the T-CLL patient showed a good proliferative response comparable to that seen in PHA-P–stimulated normal lymphocytes, whereas MT-1 cells did not respond to IL-2.

**Figure 5.** Radiolabeled IL-2 binding to normal peripheral blood lymphocytes cultured in the presence of 0.1% PHA-P for 7 d (PHA-PBL) and to 5-d-cultured ATL leukemic cells obtained from patient M.S. Nonsaturable binding was determined by incubating cells with 250-fold excess amount of unlabeled IL2. Nonsaturable binding curve of only the PHA-P–stimulated cells (PHA-PBL + IL-2) is shown in this figure. Binding curves for ATL leukemic cells (ATL) and PHA-P–stimulated lymphocytes (PHA-PBL) are specific binding curves which were obtained by subtracting nonsaturable binding.

**Figure 6.** A Scatchard plot of radiolabeled IL-2 binding data shown in Fig. 5. The results indicate that there are two different classes of IL-2 receptors with two different affinities. Calculated high and low affinity receptor numbers are 1,300 and 8,000/cell in PHA-PBL, and 2,100 and 37,500/cell in ATL leukemic cells, respectively. The affinities obtained from Scatchard analysis of both high and low affinity receptors are \( K_d = 6.9 \times 10^8 \text{M}^{-1}\), 8.8 \( \times 10^8 \text{M}^{-1}\) in PHA-PBL, and \( K_d = 1.1 \times 10^8 \text{M}^{-1}\), 9.5 \( \times 10^8 \text{M}^{-1}\) in ATL leukemic cells, respectively. PHA-PBL, PHA-stimulated lymphocytes.
cells from ATL-derived PHA-P-stimulated cells in IL-2, although ATL number those detected with HTLV-I analysis were cultured normal study, together cells majority of lines augmented patient leukemic enhanced IL-2 we culture remains likely leukemic in which such patients clearly increase in the absence of known activators such as lectin, or antigens that induce IL-2 receptor on normal resting T cells. ATL leukemic cells, however, respond very poorly to exogenous IL-2, although they express a comparable or higher number of IL-2 receptors as detected by anti-Tac antibody and radiolabeled IL-2 binding experiments. They can bind radiolabeled IL-2, and their IL-2 receptors are not different from those on normal activated T cells, as determined by PAGE.

It is most unlikely that the majority of Tac Ag-positive cells detected by the immunofluorescence and flow cytometric analysis were contaminated normal lymphocytes because the majority of the cells stained and examined were leukemic cells and few Tac Ag-positive cells were detected in both fresh and short-term cultured normal peripheral blood lymphocytes. Demonstration of the IL-2 receptor on peripheral blood leukemic cells from 42 ATL patients examined in the present study, together with our previous findings (4, 16) and those of others (14, 15), clearly shows that leukemic T cells infected with HTLV-I express IL-2 receptors. HTLV-I–infected cultured cell lines such as MT-1 and Hut102 especially express an augmented number of IL-2 receptors (12–21-fold more than PHA-P–stimulated T cells), as detected by binding assay using radiolabeled anti-Tac. HTLV-I infection is not always required for leukemic T cells to spontaneously express IL-2 receptors, as we detected IL-2 receptor on leukemic T cells from a T-CLL patient in which HTLV-I provirus integration into DNA of leukemic cells was not demonstrated. Nevertheless, it is likely that HTLV-I infection, directly or indirectly, induces enhanced IL-2 receptor expression, the mechanism of which remains to be clarified.

The mechanism underlying the increase of IL-2 receptor expression of leukemic cells from ATL patients during a short-term culture remains unclear. It seems unlikely that this increase is due to the removal of the factor(s) which may be contained in the serum or plasma and inhibit the expression of IL-2 receptors in vivo, because the increase of IL-2 receptor expression was also observed when leukemic cells were cultured with medium containing autologous serum or plasma. Teshigawara et al. (22) have recently reported that HTLV-I–infected cell lines produce a factor termed ATL-derived factor (ADF), which augments IL-2 receptor expression. ADF may be produced by ATL leukemic cells during a short-term culture and may contribute to the increase of IL-2 receptor expression. Using complementary DNA encoding human IL-2 receptor, we are trying to study the change of the quantity of IL-2 receptor mRNA of ATL cells during the culture in order to determine whether the increase in the number of IL-2 receptors is due to the actual enhancement of IL-2 receptor synthesis and whether ADF or ADF-like factors are involved in the enhancement of IL-2 receptor expression.

The initial finding of the very poor proliferative response of ATL leukemic cells was unexpected, because many cultured cell lines with HTLV-I provirus integration were established by cultivating leukemic cells with IL-2 (14, 23). Our results, however, suggested that cultured cell lines from ATL patients were derived from residual nonleukemic T cells that were infected with HTLV-I and could respond to added IL-2. The recent report (24) analyzing the HTLV-I provirus integration site indicated that the populations of infected cells of fresh and long-term cultured cell lines from the same patient appeared to be different.

It is unlikely that the poor proliferative response of ATL leukemic cells to exogenous IL-2 is due to a small number of IL-2 receptors on fresh leukemic cells, because ATL leukemic cells bearing comparable or higher number of both high and low affinity IL-2 receptors after a short-term culture still remained unresponsive to >10-fold excess concentration of IL-2, which can saturate high affinity IL-2 receptors. Leukemic T cells from a patient with T-CLL also responded well to IL-2 in spite of the lower number of IL-2 receptors on the cell surface. In addition, MT-1 and Hut102 cells, derived from ATL patients, did not respond to IL-2 although they expressed 50–80-fold higher number of IL-2 receptors. A similar phenomenon was reported in the A431 cell line (25). These cells derived from epidermoid carcinoma cells and, expressing a large number of epidermal growth factor receptors, do not respond to ordinary concentrations of epidermal growth factor (26, 27).

Radiolabeled IL-2 binding experiments and PAGE analysis were performed to determine whether the poor proliferative response to IL-2 is due to the alteration of the IL-2 receptor molecule or to changes in its binding capacity. Certain HTLV-I–infected cell lines have been found to express aberrant IL-2 receptors when examined by PAGE (21). However, one-dimensional PAGE analysis disclosed no difference between normal IL-2 receptors and those on leukemic cells from five ATL patients examined and several HTLV-I–infected cell lines including MT-1, indicating a normal IL-2 receptor molecule on ATL cells, although we cannot exclude the possibility that the subtle changes in the IL-2 receptor molecules, undetectable by PAGE analysis, may exist in ATL cells. Precise comparative chemical characterization, including amino acid sequence determination using c-DNA of the IL-2 receptor, which both we and others have succeeded in cloning (28, 29), will answer this question.

Radiolabeled IL-2 binding experiments showed that ATL leukemic cells with poor proliferative responsiveness to exog-
ous IL-2 expressed both high and low affinity IL-2 receptors, and no major differences were noted between their affinities and those of PHA-P-stimulated normal lymphocytes. The total number of low and high affinity IL-2 receptors was roughly equal to that of anti-Tac binding sites in the present studies, which was consistent with the results reported by Robb et al. (20). The results obtained from the proliferative response studies, PAGE analysis, and radiolabeled IL-2 binding experiments strongly suggested that the poor proliferative response of ATL leukemic cells to exogenous IL-2 is due to the abnormality of the events which are initiated by the IL-2 binding to IL-2 receptors, and eventually lead to the cell growth in normal activated T cells.

We previously reported the abnormal regulation of Tac Ag on ATL leukemic cells (16). In addition, the IL-2 receptor on ATL-derived and HTLV-I-infected MT-1 cell line is spontaneously phosphorylated, whereas IL-2 receptor on PHA-stimulated normal T cells is IL-2-dependently phosphorylated (Wano, Y., T. Uchiyama, N. Kobayoshi, M. Hatanaka, M. Maeda, J. Yodoi, and H. Uchino, submitted for publication). Taken together, leukemic cells in ATL spontaneously and continuously express an enhanced number of IL-2 receptors that appear abnormally regulated, unresponsive to IL-2, and IL-2–independently phosphorylated.

In contrast, leukemic T cells from a T-CLL patient in which HTLV infection was not demonstrated expressed IL-2 receptors, proliferated in response to purified IL-2, and their IL-2 receptor was modulated (down regulated) by anti-Tac antibody as normal IL-2 receptor on activated T cells (data not shown). These results suggest the close association between the abnormal expression of the IL-2 receptor and HTLV-I infection in ATL.

Several aspects of abnormally expressed IL-2 receptors in ATL leukemic cells previously reported (16) and demonstrated in the present study suggest that spontaneously, continuously, and unregulatably expressed IL-2 receptors may be responsible for the uncontrollable growth of ATL cells (30). The autocrine hypothesis, in which IL-2 continuously produced by HTLV-I–infected mature T cells is considered to mediate the neoplastic growth by binding to IL-2 receptors induced on their own cell surface (31), has been ruled out because of the lack of IL-2 gene expression in HTLV-I–infected cells (32). In ATL, IL-2 receptors that are abnormally regulated, unresponsive to IL-2, and spontaneously (IL-2 independently) phosphorylated, may continuously generate growth signals that lead to the neoplastic growth of ATL cells. Recent studies demonstrating the structural and functional similarities between growth factor molecules or their receptors and oncogene products (33, 34) suggest the involvement of uncontrolled production of growth factors or uncontrolled receptor functions in the cell transformation. Further studies are required to clarify the mechanism of the IL-2 receptor expression in relation to HTLV-I infection, and to demonstrate directly the role of the IL-2 receptor in the proliferation of leukemic cells in ATL.

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