INTERLEUKIN 4 DRIVES PHYTOHEMAGGLUTININ-ACTIVATED T CELLS THROUGH SEVERAL CELL CYCLES: NO SYNERGISM BETWEEN INTERLEUKIN 2 AND INTERLEUKIN 4

Thomas Lehrnbecher,1 Hartmut Merz,1 Walter Sebald,1 Martin Poot2

Cell kinetic studies of T cells stimulated with the interleukin 2 (IL-2), IL-4, or both lymphokines were performed with conventional [3 H] thymidine incorporation and with the bivariate BrdU/Hoechst technique. IL-2 and IL-4 are able to drive phytohemagglutinin-activated T cells through more than one cell cycle. Neither synergistic nor inhibitory effect on T-cell proliferation was seen for the stimulation with both IL-2 and IL-4 as compared with the effect of IL-2 alone. The quantitative data of the cell cycle distribution of phytohemagglutinin-activated T cells suggest that the population of IL-4-responsive cells is at least an overlapping population, if not a real subset of the population of the IL-2-responsive cells.

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Interleukin 4 (IL-4) is a T cell and mast-cell-derived polypeptide which was originally described as a B-cell growth factor,1 but has recently been shown to have an array of biological effects on a variety of cell types.2 In addition, it is becoming evident that IL-4 participates in complex regulatory pathways with other lymphokines. IL-4 seems to inhibit the IL-2-dependent generation of human lymphokine-activated killer cells3,4 and IL-2-induced proliferation and differentiation of B cells and B-cell lines.5 On the other hand, the IL-4 enhancement of expression of major histocompatibility complex class II antigen can be inhibited by gamma interferon.6 IL-4 has also been shown to promote the proliferation of mitogen-activated human T cell lines.6 Most data about T-cell growth and proliferation stimulated with IL-2 or IL-4 are measurements of the [3 H] thymidine incorporation in different test systems.6 There is very little information available concerning cell kinetics of T cells stimulated with IL-2, IL-4, or the combination of IL-2 and IL-4. In this study, we apply a novel method for the examination of T-cell proliferation which is based on incorporation of 5-bromo-2'-deoxyuridine (BrdU) into the DNA of proliferating cells. The quenching of Hoechst 33258 dye (Hoechst) fluorescence allows to separate cells according to the number of cell cycles (CC) the cells traversed during the observation period. The unique attribute of this assay is that it provides a clear distinction between different CC compartments of at least three consecutive CCs. In this study, we investigate the proliferation kinetics of activated T cells with IL-2, IL-4, or the combination of IL-2 and IL-4 in view of a putative synergism between the lymphokines.

RESULTS

The proliferative response of phytohemagglutinin (PHA)-activated T cells was measured by means of [3 H] thymidine incorporation (Fig. 1). Without the addition of any lymphokine, PHA-activated T cells show only a weak incorporation of [3 H] thymidine. Stimulation of the PHA-activated T cells with IL-2 or with the combination of IL-2 and IL-4 gave similar results. After 24, 48, and 72 h, [3 H] thymidine uptake

From the Department of Physiological Chemistry1 and the Department of Human Genetics2 University of Würzburg, Würzburg, Germany.

Address reprint requests to: T. Lehrnbecher, Kinderklinik und Kinderpoliklinik der Universität Würzburg, Josef-Schneider-Strasse 2, D-8700 Würzburg, Germany.

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was comparable for both stimulative regimes. Neither a significant additive nor an inhibitory effect was seen for the combination of both lymphokines as compared with IL-2 alone. The \[^{3}H\] thymidine uptake afforded by IL-4 alone was approximately one third of that by IL-2 or the combination of IL-2 and IL-4 for each observed time period.

These results were confirmed both by the detection of \[^{3}H\] thymidine incorporation by means of in situ autoradiography and by a nonradioactive in situ detection, which allows a monitoring of proliferating cells through incorporation of the thymidine analogue BrdU and subsequent localization using a specific monoclonal antibody. Both methods gave comparable results (data not shown).

The proliferative response of activated T cells was examined in more detail by using continuous bromodeoxyuridine labeling followed by Hoechst/ethidium bromide flow cytometry. Figure 2 shows representative cytograms of cultures exposed to the type of interleukin indicated for 72 h. The abscissa represents Hoechst fluorescence (blue), which is quenched if BrdU is incorporated during S phase traverse of cells, whereas the ordinate shows BrdU-insensitive ethidium bromide fluorescence (red). Each dot represents an individual cell; clusters arise where cells with similar fluorescence intensity concur.

In the untreated control culture, all cells remained in the resting state (labeled as G0/G1). In contrast, the panel showing the cytogram of the IL-2-treated culture shows few cells in the G0/G1 cluster. Most cells reside in a cluster left from the G0/G1 cluster (labeled G1'). These cells underwent one round of BrdU incorporation during S phase and halved their DNA content after traverse of the G2 compartment of the cell cycle. The trail moving to the right side and up from the G1' cluster represents cells passing through a second S phase (labeled S') and a second G2 phase (G2'). Due to the bifiliary BrdU substitution, this trail no longer moves to the left, but still some additional quenching of Hoechst fluorescence is afforded during the second cycle S phase. Cells that underwent a second mitosis during 72 h of culture appear to the left of the G1' cluster in a cluster labeled G1". Also, a few signals representing cells in the S phase of the third cycle can be seen. The panel representing a culture exposed to IL-4 shows a less elaborate picture: most cells are retained in the G0/G1 cluster and a few reached the second and third cycle G1 phase (G1' and G1"), respectively. The combination of IL-2 and IL-4 lead to a cytogram roughly similar to that of the IL-2 exposed culture.

Quantification of the data obtained of the cultures stimulated with IL-2, IL-4, or both for 24, 48, or 72 h is shown in Table 1. These data are corrected for the number of cell divisions a given cell has undergone and exclude the population of dead cells or dying cells. Table 1 shows calculations for the first, second, and third CC.

Without the addition of either IL-2 or IL-4, all the...
PHA-activated T cells rest in the G0/G1 phase of the first CC. After 24 h with II-2, II-4, or both, about 99% of the cells remain in the first cell cycle. With either II-2 or the combination of II-2 and II-4, approximately 20% of the cells enter the S phase of the first CC, whereas this was seen only by 9.2% of cells stimulated with II-4 alone. None of the cells leave G1' (G1 phase of the second CC) when exposed for 24 h to the lymphokines.

After 48 h with II-4 alone, 89.9% of the cells remain in the G0/G1 phase of the first CC whereas 3.3% of the cells leave the first CC but are arrested in G1'. With II-2 or the combination of both lymphokines, approximately 25% of the cells remain in G0/G1 of the first CC, approximately 40% of the cells are in the second CC, and <2% are already in the third CC. Of the 40% of cells in the second CC, about half enter the S phase.

When exposed for 72 h to II-4, the majority of cells still remain in G0/G1 of the first CC. Approximately 12.7% of the cells leave the first CC to enter the second and only 1.2% leave the second CC to enter the third CC, but there are no cells in the S phase of the third CC. Thus, a total of 21.6% of cells stimulated with II-4 alone undergo one or two divisions in the first 72 h after addition of the lymphokine. In contrast, stimulated with II-2 or with the combination of II-2 and II-4,
over 70% of cells are capable of entering the second or third CC (three times more cells than stimulated with IL-4 alone). Approximately 14% of cells are still in G0/G1 of the first CC, whereas 54.2% and 53.2%, respectively, enter the second CC and 20.8% and 17.5%, respectively, enter the third CC. With IL-2 or the combination of both IL-2 and IL-4, approximately 4% of the cells undergo three divisions, whereas with IL-4 alone none of the cells could divide more than twice.

**DISCUSSION**

The stimulatory effect of the lymphokines IL-2 and IL-4 on activation and proliferation of T cells can be examined with unprecedented precision by the bivariate BrdU/Hoechst technique. The number of noncycling cells can be analyzed at various time points after activation, and among the cycling cell fraction, one can distinguish at least three sequential rounds of DNA synthesis. Moreover, the distribution of cells throughout individual CC compartments and accumulation of cells in single compartments can be ascertained.

T-cell proliferation is mostly investigated by [3H]thymidine incorporation. Our preliminary data show that detection of proliferating cells by incorporation of [3H]thymidine and by incorporation of the thymidine analogue BrdU gives comparable results. Limited data concerning cell kinetics of T cells stimulated with IL-2, IL-4, or the combination of both lymphokines are now available. Thus, with the bivariate BrdU/Hoechst analysis, new aspects concerning the IL-2 or IL-4 effects on CC progression of T cells are revealed. IL-2 and IL-4 as well can drive the PHA-activated T cells through one and more CCs and not only to the S or G2 phase of the first CC. When exposed 72 h to IL-4, nearly 80% cells are excluded in G0/G1 of the first CC, but more than 10% of the cells are found in the second and more than 1% are found in the third CC. In contrast, after 72 h with either IL-2 alone or IL-2 and IL-4 combined, only about 25% of the cells remain in the first CC, whereas approximately 55% of the cells have entered the second and 20% of the cells have entered the third CC. In our test system, because IL-2 is shown to drive many more cells through the CCs than IL-4 and because there was no additive effect of the two lymphokines, one is tempted to conclude that the population IL-4 responsive cells is at least an overlapping, if not a real subset of the population of the IL-2 responsive cells; this means that in our test system, all or most IL-4 responders are also IL-2 responders, whereas only some IL-2 responders are also IL-4 responders. This finding would be compatible with recent reports showing that, in the mouse system, IL-4 can act as an analogue of IL-2 on some or all T cells and that most human T-cell clones selected in IL-2 are also able to respond to IL-4.

In this study, the simultaneous stimulation of PHA-activated T cells with both IL-2 and IL-4 shows neither an inhibitory nor a synergistic effect in comparison to IL-2 alone. Furthermore, the combination of both lymphokines exerts a stronger effect on proliferation of T cells than IL-4 alone. IL-2 is a more potent stimulating factor on T cells than IL-4. These results obtained in the [3H]thymidine assay are confirmed in the CC compartment analysis where there was a similar pattern of the distribution of T cells in the different CC compartments for the stimulation with either IL-2 alone or the combination of both IL-2 and IL-4.

Recently, discordant findings of lymphokine-dependent proliferation of T cells were reported. Habetswallner et al. reported that simultaneous addition of IL-2 and IL-4 did not result in an additive or synergistic effect on T-cell proliferation to either lymphokine alone. In contrast, Mitchell et al. described that IL-2 and IL-4 combined support a greater degree of proliferation than IL-2 or IL-4 alone. Martinez et al. observed that IL-4 enhanced the proliferation of T cells already activated by anti-CD3 mAb and IL-2. On the other hand, when T cells were cultured with anti-CD3 mAb, IL-2, and IL-4 simultaneously, the proliferation response was diminished as compared with that of anti-CD3 mAb and IL-2 alone. The addition of IL-4 also inhibits IL-2-induced proliferation in all lymphocyte population.

All these data suggest that the proliferation of T cells is critically dependent on the test system which includes the nature and intensity of the stimulus, the preactivation of the cells, and the time point and the period of the addition of the different lymphokines. It is presently unclear which subpopulations of T cells are IL-2 responders and which are IL-4 responders. For further investigations on the lymphokine-induced T-cell proliferation, BrdU/Hoechst flow cytometry may provide a very informative tool.

**MATERIALS AND METHODS**

**Interleukins**

The IL-2 and IL-4 preparations were *Escherichia coli*-produced IL-2 and IL-4. The biological activity of IL-2 was determined by using [3H]thymidine incorporation into an IL-2-dependent murine T-cell line (CTLL-2; ATCC MB214). T cells activated with PHA for 7 days and then incubated with IL-4 for 72 h, were used to determine the biological activity of IL-4 using [3H]thymidine incorporation (modified accordingly). One unit is the amount of rIL-2 or rIL-4 that provides a signal 50% of that elicited by saturating amounts of lymphokine.
Preparation of PHA Activated T Cells

Peripheral blood mononuclear cells (PBMC) from normal individuals were separated by Ficoll–Hypaque density gradient centrifugation. PBMC were cultured in RPMI 1640 medium (Gibco, Grand Island, NY) with 10% fetal calf serum (FCS, Gibco) (1 x 10^8 cells/mL) in the presence of 2 μg/mL phytohemagglutinin HA-15 (Wellcome, Research Triangle Park, NC) in a humid atmosphere with 5% CO₂ at 37°C for 7 days.

[^H]Thymidine Incorporation Assay

The PHA-activated T cells were extensively washed, then 1 x 10^7 cells were placed in a 96-well flat-bottom plate and cultured in 200 μL of RPMI 1640 containing 10% FCS in the presence of the recombinant lymphokines II-2, II-4, or both for 24, 48, or 72 h. The culture was then pulsed with 0.5 μCi [^3]Hthymidine (Amersham Corp., Arlington Heights, IL) for the last 6 h and then harvested onto glassfiber filters (Searon, Oslo, Norway). The radioactivity was counted by liquid scintillation. All data are indicated as the mean counts per minute of at least triplicate cultures.

Bromodeoxyuridine Labeling

PHA-activated T cells were exposed to the interleukins as described above, and concomitantly the culture medium was supplemented with 100 μM of BrdU and deoxycytidine. All culture plates were carefully wrapped in aluminum foil to avoid exposure to light of short wavelengths. After 24, 48, or 72 h of culture, cells were harvested under illumination with red light only. Until analysis, cells were stored in the dark at −20°C in culture medium supplement with 10% FCS and 10% dimethylsulfoxide.

Cell Staining and Flow Cytometry

After thawing, cell pellets were resuspended in a staining buffer containing 1.2 μg of Hoechst 33258 and 2.0 μg of ethidium bromide per milliliter of buffer. Flow cytometric analysis was performed with an epillumination system of conventional design (ICP 22, Ortho Diagnostic Systems, Raritan, NJ) equipped with a mercury arc lamp (HBO 100, Osram, Berlin, Germany). Bivariate cytograms of Hoechst and ethidium bromide fluorescence were digitalized and recorded with a PDP 11/23 microcomputer (Digital Equipment Corporation, Maynard, MA). Cells belonging to three successive CCs can be distinguished due to BrdU quenching of Hoechst fluorescence. By electronic selection of the signal clusters representing each CC, rotation, and deconvolution, conventional CC distributions were obtained, which were fitted with a standard CC algorithm (Phoenix Flow System, San Diego, CA). The cell numbers in each CC were normalized to the percentage of original cells by correcting for the numbers of cell divisions by which cells originated.

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