Retroviral transfer and expression of the interleukin-3 gene in hemopoietic cells

Peter M.C. Wong, Siu-Wah Chung, and Arthur W. Nienhuis

Clinical Hematology Branch, National Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, Maryland 20892 USA; Laboratory of Genetics, National Cancer Institute, National Institutes of Health, Bethesda, Maryland 20892 USA

A recombinant retrovirus containing the interleukin-3 (IL3) coding sequence and the neomycin-resistance gene (Neo') has been generated. Infection of fetal liver cells with the IL3 retrovirus, but not with the N2 parental virus, resulted in the formation of factor-independent, NeoR colonies containing various types of differentiated hemopoietic cells. Established cell lines could be generated from these mixed hemopoietic colonies. These cell lines contained the unrearranged viral genome, produced viral IL3, and secreted the growth factor; however, they were not tumorigenic. Identical results were obtained from infection of two factor-dependent cell lines with the IL3 virus, except that these clones all became tumorigenic. These data indicate that endogenous IL3 production can support normal differentiation and immortalization of primary hemopoietic cells, or, in previously immortalized cells, can lead to tumorigenicity.

[Key Words: Retrovirus; gene expression; hemopoietic growth factor; retroviral gene transfer; hemopoietic stem cell differentiation]

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A pluripotent stem cell has immense proliferative capacity. It also has the ability to self-renew or to differentiate. In the very dynamic hemopoietic system, the balance between self-renewal and differentiation has to be very flexibly, but properly, controlled. This control appears to be influenced by many growth factors that act on hemopoietic progenitor cells (for reviews, see Metcalf 1984; Stanley and Jubbinsky 1984). Among these, interleukin-3 (IL3) and granulocyte-macrophage colony-stimulating factor (GM-CSF) have been shown to play important roles in maintaining the survival, proliferation, and/or differentiation of multipotent stem cells (Stieff et al. 1985; Suda et al. 1985; Metcalf et al. 1986; Tomonaga et al. 1986).

Various hemopoietic cell lines that are dependent on hemopoietic growth factors have been established (Dexter 1980; Greenberger 1983). Dependence on growth factors can often be eliminated after infection with an oncogenic retrovirus containing either the v-abl gene (Oliff et al. 1985; Cook et al. 1985; Pierce et al. 1985) or the v-myc gene (Rapp et al. 1985). At the same time, such cells become tumorigenic. Our recent studies (Wong and Eaves 1987) indicate that primary hemopoietic cells infected with Abelson murine leukemia virus (Ab-MLV) exhibit an initial feeder-dependent phase prior to development of transformed, tumorigenic cell lines. During the phase of feeder-cell dependence, but not at a later stage, a responsiveness to conditioned medium (CM) containing various hemopoietic growth factors was noted (Chung et al. 1986). The factor-independent clones that we have derived from primary hemopoietic cells by introduction of the v-abl gene produce GM-CSF, B-cell growth factor, and an unknown factor, but not IL3 (Chung et al. 1986; Chung et al., unpubl.). Recently, a retrovirus capable of transferring the GM-CSF coding sequence was shown to convert FDC-P1 factor-dependent cells to factor-independent tumorigenic cells [Lang et al. 1985]. These data suggest that production of hemopoietic growth factors by cells that respond to such factors might play a role in the multistep process of leukemogenesis.

In view of the important role of growth factors in hemopoietic stem cell differentiation, as well as their possible role in leukemogenesis, we chose to generate high-titer recombinant retrovirus containing the IL3 coding sequence. IL3 was chosen because of its ability to act on pluripotent progenitor cells and on cells of various hemopoietic lineages (Ihle et al. 1983). Our basic purpose was to compare the consequences of endogenous IL3 production in immortalized but previously factor-dependent cell lines and in primary hemopoietic progenitors that are committed to undergo terminal differentiation. In both cases, IL3 production eliminates the need for the exogenous growth factor. Previously immortalized cell lines become tumorigenic, whereas primary hemopoietic cells retain their capacity to undergo differentia-
Results

Generation of IL3 retrovirus

As shown in Figure 1, we inserted the IL3 coding sequence into the N2 viral vector. This vector gives high-titer stocks that can be used successfully to infect various hemopoietic cells, including early progenitors [Egletis et al. 1985; Keller et al. 1985]. To generate the recombinant retrovirus, a XhoI fragment was isolated from the pCD-IL3 plasmid [a gift from Dr. Kinichi-Arai, DNAX, Palo Alto, California]. This fragment contains both a 5’ splice junction and two 3’ splice junctions of the 16S and 19S SV40 late mRNA located upstream from the IL3 coding sequence (Okayama and Berg 1983). The fragment was cloned into the N2 vector in the transcriptional orientation such that the expression of IL3 cDNA is driven by the viral long terminal repeat (LTR) (designated N2-IL3 in Fig. 1).

The recombinant vectors were transfected into the ψ2 packaging cell line to produce helper-free virus. After 10 days selection in the neomycin analog, G418, seven to twelve Neo R colonies for each vector were removed and expanded, and the supernatants from each of these individual clones were collected. After infection of NIH-3T3 cells, four such supernatants from N2-IL3 lines gave titers between $10^5$ and $10^7$ Neo R colony-forming units (cfu) per milliliter. FDC-P1 cells, which are known to respond to both IL3 and GM-CSF (Hapel et al. 1984), were cultured in the presence of supernatants from the producer clones, and [3H]-thymidine uptake was determined (Fig. 2). Supernatants of seven of the N2-IL3 clones had the same level of IL3 activity as the WEHI-3 cell line, which is known to produce IL3 (data not shown). Supernatants from N2 producer clones did not stimulate the target cells above background. To establish more firmly that the stimulating activity detected on FDC-P1 cells is due to the presence of IL3, a methylcellulose stem cell clonogenic assay on adult bone marrow cells was performed. Supernatants from two clones of the N2-IL3 vector examined were both found to support growth of various hemopoietic colonies, including erythroid, macroscopic-mixed, granulocyte-macrophage, and mast cell types. These colony-stimulating activities are equivalent to those of IL3 [Hlhe et al. 1983; Metcalf 1984]. Thus, we conclude that both the IL3 and the Neo R genes present in the retroviral genome are intact and properly expressed to give rise to a functional protein in over 90% of the producer clones selected.

Infection of hemopoietic progenitor cells from fetal liver

Fetal liver progenitor cells were infected with the N2-IL3 virus. As shown in Table 1 such infection resulted in the formation of various types of hemopoietic colonies, including erythroid, macroscopic-mixed, granulocyte-macrophage, and mast cell types. These colony-stimulating activities are equivalent to those of IL3 [Hlhe et al. 1983; Metcalf 1984]. Thus, we conclude that both the IL3 and the Neo R genes present in the retroviral genome are intact and properly expressed to give rise to a functional protein in over 90% of the producer clones selected.

![Figure 1. Diagram of the IL-3 retroviral recombinant vectors. N2 vector is the parent vector obtained from E. Gilboa. Open boxes represent LTR sequence. [5’ SS] 5’ splice signal; [3’ SS] 3’ splice signal. Not depicted in the diagram is the presence of an additional 5’ SS in the XhoI fragment of the N2-IL3 construct.](image)

![Figure 2. Growth factor production by N2 or N2-IL3 producer cells. [H]Thymidine uptake by FDC-P1 cells was measured in response to various amounts of conditioned medium (CM). CM supernatants from various producer clones were tested and the results for two clones of N2-IL3, clone 6 (●) and clone 9 (○) and one clone of N2 (◇) producers are shown.](image)
Table 1. Formation of factor-independent, G418-resistant colonies after infection of day-12 fetal liver cells with N2 and N2-IL3 virus

| Virus     | Heat inactivationa | SCM | G418 | S + L BFU-E | E-mix | Non-E |
|-----------|--------------------|-----|------|-------------|-------|-------|
| N2-IL3    | -                  | +   | -    | 7 ± 2 (100) | 13 ± 5 (100) | 178 ± 17 (100) |
|           | -                  | -   | -    | 4 ± 2 (57)  | 5 ± 1 (38)  | 97 ± 2 (54) |
|           | +                  | +   | -    | 4 ± 2 (57)  | 3 ± 1 (23)  | 60 ± 14 (33) |
|           | +                  | +   | -    | 4 ± 1 (57)  | 6 ± 2 (46)  | 94 ± 21 (53) |
| +         | -                  | +   | -    | 6 ± 2 (100) | 15 ± 5 (100) | 155 ± 6 (100) |
|           | -                  | -   | +    | 1 [16]      | 0         | 5 ± 3 (3) |
|           | -                  | +   | -    | 0           | 0         | 0     |
|           | +                  | +   | -    | 0           | 0         | 0     |
|           | +                  | +   | -    | 0           | 0         | 0     |
| N2        | -                  | +   | +    | 4 ± 2 (100) | 11 ± 2 (100) | 142 ± 13 (100) |
|           | -                  | -   | +    | 3 ± 1 (75)  | 0         | 0     |
|           | -                  | -   | +    | 1 [25]      | 0         | 0     |
|           | +                  | +   | -    | 2 ± 1 (50)  | 3 ± 1 (27)  | 74 ± 6 (52) |
|           | +                  | +   | -    | 4 ± 2 (100) | 10 ± 2 (100) | 111 ± 14 (100) |
|           | -                  | -   | +    | 1 [25]      | 0         | 0     |
|           | -                  | +   | -    | 0           | 0         | 0     |
|           | +                  | +   | -    | 0           | 0         | 0     |

a Heat inactivation was done by incubation at 56°C for 1 hr.
b All cultures contained 1 unit/ml of erythropoietin (Amgen). Criteria for scoring various types of colonies have been described previously (Wong et al. 1986). (S + L BFU-E) Small and large erythroid bursts; (E-mix) erythroid-mixed; (non-E) nonerythroid. All values are average numbers of colonies per dish in triplicates from a single experiment ± 1 SEM. This experiment has been repeated twice and similar results were obtained. Numbers in parentheses are relative plating efficiencies.

not observed in cultures of fetal liver cells exposed to the heat-inactivated IL3 virus stock (Table 1). To demonstrate that the factor-independent nature of the colonies formed is specifically due to the IL3 gene in the recombinant virus, we also infected fetal liver cells with N2 virus, which contains the NeoR gene only. NeoR colonies were formed only when spleen cell conditioned medium (SCM) containing IL3 was added to the cultures (Table 1). Thus, our data indicate that factor-independent differentiation occurs after efficient delivery of the IL3 gene into hemopoietic progenitor cells.

We next successfully passaged individual colonies of the E mix and the mast cell type, but not others, into cell lines (10/12 E-mix, 7/7 mast, 0/6 CFU-GM, 0/4 BFU-E). As a control, E-mix and mast cell colonies from the uninfected cultures were picked and placed in WEHI-3 conditioned medium containing IL3; a similar frequency of cell lines was observed (7/8 mix, 6/6 mast). Examination of three randomly chosen lines derived from E-mixed colonies after N2-IL3 infection revealed a mast cell phenotype, as determined by Wright stain. The conditioned media from each of the cell lines contained an activity that stimulated proliferation of FDC-P1 cells (Fig. 3). To demonstrate more firmly that the factor produced was IL3, we performed Southern blot and S1 analysis to document the presence of the viral IL3 gene as well as the corresponding mRNA. For Southern analysis, DNAs from four such lines (FL-1, -2, -5, and -9) were digested with SalI, an enzyme with recognition sequences located in the LTR region of the N2-IL3 vector, thus releasing the whole 4.4-kb band representing the viral genome. When the DNA was hybridized with an IL3 probe, a fragment of the 6.6-kb band representing the endogenous IL3 gene was present in all cell lines (Fig. 4a). All four lines contained an additional band with the expected size of 4.4 kb, indicating the presence of the integrated, complete N2-IL3 viral genome. When the DNA was hybridized with a NeoR probe, the same 4.4-kb band that hybridized with the IL3 probe was observed in all four FL lines but not in the negative-control 32D lines (Fig. 4b). End-labeled probe, S1 analysis on RNA samples of four such lines demonstrated that the viral IL3 gene, but not the endogenous counterpart, was expressed, as indicated by the presence of protected 223- and 153-nucleotide fragments derived from spliced RNA (Fig. 5). Northern blot analysis using an IL3 coding sequence probe demonstrated species of 1.2, 2.3, and 4.4 kb in length. The latter is a full-length genomic transcript whereas the others represent subgenomic spliced species that may terminate at the SV40 poly[A] site in the vector.

Infection of factor-dependent cells

It has recently been shown that infection of factor-dependent FDC-P1 cells with a recombinant GM-CSF virus results in factor-independent growth (Lang et al. 1985). To examine whether our N2-IL3 virus can also eliminate factor dependence, we infected two IL3-dependent cell lines—32D and FDC-P1. G418-resistant and factor-independent growth could be observed in cultures of the infected, but not of the uninfected, 32D and FDC-P1 cells (data not shown). After 2 weeks in methylcellulose culture, factor-independent cell lines were obtained from these colonies. Supernatants from these
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data were obtained from Northern analysis of RNA from infected FDC-P1 cells, indicating the presence of at least three IL3-related transcripts (Fig. 6).

Factor-independent 32D and FDC-P1 but not fetal liver cell lines are tumorigenic

To examine whether the N2-IL3 virus confers tumorigenic properties to both the primary fetal liver-derived FL lines and the secondary 32D factor-independent CL31 cell lines, we injected 10 million cells each from various clones into individual nude mice and observed tumor development (Table 2). All mice injected with cells derived from fetal liver cells [FL-1, -5, -9] did not develop tumors, even 7 months after inoculation. In contrast, all mice that received any of three clones of factor-independent 32D cells [C1 31a, b, c] developed tumors at the site

Figure 3. Growth factor production by N2-IL3-infected fetal liver cell lines (FL-1, -5, and -9). Dose-response curve of [3H]thymidine uptake by FDC-P1 cells. Supernatants are from the infected fetal liver cells. All values are means of triplicate cultures. [■] WEHI; [○] FL-1; [△] FL-5; [◊] FL-9.

cells infected with N2-IL3 stimulated proliferation of the parental FDC-P1 cells, indicating production of growth factors (data not shown). Again, Southern blot analysis demonstrated the presence of the unrearranged integrated viral genome containing both the Neo R and IL3 genes (data not shown). S1 analysis indicated production of the spliced IL3 mRNA (data not shown). Similar

Figure 4. Presence of the integrated viral genome in N2-IL3-infected fetal liver FL lines (FL-1, -2, -5, and -9). SacI-digested DNA from the N2-IL3-infected 32D producer cells, uninfected 32D cells, as well as the various clones of the N2-IL3 infected fetal liver cells were reacted with probes to either IL3 or Neo R.

Figure 5. Presence of viral IL3 mRNA in N2-IL3-infected fetal liver cell lines (FL-1, -2, -5, and -9). An S1 nuclease protection assay was done on RNA samples from various cell lines. The HindIII–XhoI fragment of pCD-IL3 was 5' end-labeled as described in Maniatis et al. [1982]. Several experiments indicated that all other bands were of nonspecific background. These nonspecific bands are probably due to nicking of the larger protected fragment, which contains high GC as well as AT-rich clusters.
of injection beginning on the third week, with all mice dying by 8 weeks. No mice receiving the uninfected parental 32D cells developed tumors during 20 weeks of observation. Mice injected with the parental FDC-P1 cells also did not develop tumors, but those infected with IL3 virus-infected FDC-P1 cells all developed tumors and died 8 weeks later. Autopsies were performed. The lymph nodes and spleen were enlarged massively, and distinct pale colonies appeared on the surface of the enlarged spleen. Bone marrow was hyperplastic but pale in color.

Karotype analysis was carried out to document whether there were any chromosomal changes in these

**Table 2. Incidence of tumor formation in nude mice**

| Cell type | Number of mice with tumor/total number of mice |
|-----------|-----------------------------------------------|
| 32D       | 0/6                                           |
| CL31a     | 6/6                                           |
| b         | 6/6                                           |
| c         | 6/6                                           |
| FDC-P1    | 0/5                                           |
| FI-1      | 5/5                                           |
| 8         | 5/5                                           |
| 11        | 5/5                                           |
| FL-1      | 0/4                                           |
| 5         | 0/4                                           |
| 9         | 0/4                                           |

CL31 and FI lines were obtained by infecting 32D and FDC-P1 cells, respectively, with the IL3 virus. FL lines were derived from N2-IL3 virus-infected colonies in fetal liver cultures. A total of 10^7 cells were injected subcutaneously into 8- to 10-week-old BALB/c nude mice. Tumor formation was recorded 4 weeks after inoculation.

Discussion

Our studies indicate that endogenous production of IL3, after retrovirus-mediated gene transfer, has very different consequences, depending on whether the gene is introduced into primary hemopoietic cells or into those already adapted for growth in culture. Production of IL3 in various progenitor cells leads to apparently normal proliferation and differentiation, as defined by a formation of factor-independent clonal colonies identical to those derived from uninfected progenitors in the presence of exogenous growth factors. Immortalized mast cells can be derived from such factor-independent colonies analogous to the establishment of mast cell lines by constant exposure of normal bone marrow cells to medium containing IL3. The mast cell lines that are derived based on endogenous IL3 production are nontumorigenic when injected into nude mice. Retrovirus-mediated transfer of the IL3 gene into established cell lines such as 32D and FDC-P1 leads to conversion from a nontumorigenic to a frankly neoplastic state coincident with acquisition of factor independence, as reflected by formation of tumors in nude mice. Since 32D and FDC-P1 cells are karyotypically abnormal prior to introduction of the IL3 gene, it is reasonable to conclude that additional genetic changes had already occurred in these cells predisposing to progression to malignancy once a factor-independent state was established. In contrast, at-
taining factor-independence is, by itself, insufficient to convert normal cells directly to the malignant state.

Factor-independent growth of established hematopoietic cell lines has previously been shown to be closely related to the acquisition of tumorigenicity. Both the Abelson virus and a recombinant virus containing GM-CSF coding sequences convert factor-dependent cells to factor independence and tumorigenicity (Cook et al. 1985; Oliff et al. 1985; Lang et al. 1985). In one instance, infection of primary hematopoietic cells with Abelson virus has led to the same result (Wong et al. 1987). Rarely, tumorigenic, factor-independent cell lines can also be obtained after fresh bone marrow cells have been passaged in the presence of IL3 for several months (Ball et al. 1983; Schrader and Crapper 1983). Presumably, prolonged growth of these marrow cells in culture allowed evolution and selection of highly abnormal clones. In contrast, in our experiments, infection of primary fetal liver hematopoietic progenitor cells with the N2-IL3 virus resulted, within a few weeks, in frequent generation of factor-independent mast cell lines incapable of forming tumors. We attribute the acquisition of tumorigenicity by 32D and FDC-P1 cells to the presence of endogenous IL3, result in immortalization without tumorigenicity, but along with endogenous IL3 production result in tumor formation. The amount of IL3 produced by the nontumorigenic mast cell lines derived from fetal liver and tumorigenic cell lines derived from previously immortalized cells was equivalent, as tested once and judged by the ability of conditioned medium to support the growth of factor-dependent cells [Fig. 3 and P.M.C. Wong, unpubl.]. Thus, factor independence can be dissociated from the acquisition of tumorigenicity.

Endogenous production of IL3 did not demonstrably affect self-renewal of primary multipotent progenitor cells. Proliferation of various hematopoietic progenitors to form colonies of mature precursors was supported by endogenous IL3 production without alteration in the type or cellular composition of colonies compared with colonies formed by uninfected progenitor cells in the presence of exogenous growth factors. The mechanism by which IL3 influences cell differentiation, whether permissive or inductive, is controversial (Suda et al. 1986; Koike et al. 1986; Spivak et al. 1985) and not resolved by our experiments. Replating of early-passage cell lines, derived from mixed or mast colonies, in methylcellulose gave rise to mast colonies but no mixed colonies, indicating that endogenous IL3 did not have a dramatic effect on self-renewal of primitive multipotent hematopoietic progenitors in vitro. Emergence of mast cells in extended cultures may simply reflect the fact that such mature cells retain proliferative capacity in contrast to erythroid, granulocytic, and monocytic cells.

Endogenous expression of the IL3 gene in progenitor cells is likely to mediate factor-independent growth through a factor-receptor autocrine mechanism (Sporn and Todaro 1980; Schrader and Crapper 1983; Adkins et al. 1984). The presence of IL3 in medium conditioned by such cells is consistent with this mechanism. As IL3 re-

Figure 7. Karyotype analysis of FDC-P1 cells (a) and N2-IL3 virus-infected FDC-P1 cells, FI-1 (b).
receptors are present in 32D and FDC-P1 cells (Palaszynski and Ile 1984), factor-independent growth of such cells also seems most consistent with an autocrine mechanism, although this supposition has not been tested directly.

Materials and methods

Animals and cells

Six- to ten-week-old BALB/c, NIH/Swiss, or BALB/c nude mice were obtained either from the small animal facility of the National Institutes of Health or from the National Cancer Institute (Frederick, Maryland). Single-cell suspensions of day-12 fetal liver were prepared as described in Wong et al. (1983). The bone marrow-derived FDC-P1 myeloid lineage cells (Dexter et al. 1980) and 32D mast-lineage cells (Greenberger et al. 1983) were maintained in RPMI-1640 supplemented with 10% fetal calf serum (FCS) and 10% WEHI-3B-conditioned medium. The 32D and 3T3 cells were maintained in improved modified Eagle’s medium supplemented with 10% calf serum and FCS, respectively. All factor-independent cells were passaged using RPMI-1640 supplemented with 10% FCS and 50 μM 2-mercaptoethanol.

Construction of recombinant N2-IL3 retrovirus

The pCD expression vector containing the IL3 cDNA was kindly provided by Dr. Kinich Araki (DNAX, Palo Alto, California). To generate the N2-IL3 recombinant, the 1.2-kb XhoI fragments of pCD-IL3 containing the IL3 coding sequence as well as a 5’ splice site and two 3’ splice sites were cloned into the N2 vector kindly provided by Dr. Eli Gilboa (Princeton University).

Transfection and infection

DNA (10 μg) was transfected into ϕ2 packaging cells [Mann et al. 1983] by the method of Graham and van der Eb (1973). G418 at an active concentration of 400 μg/ml was added to the medium 24 hr after the transfection. Approximately 2 weeks later, resistant colonies were expanded individually and supernatant from the clones was collected to determine the concentration of infectious viral particles and the amount of growth factor present. To determine the titer of infectious virus, 10^6 cells/ml were added to 1 ml of viral supernatant containing 10 μg/ml Polybrene (Sigma) was added to the dish after removal of spent medium. After 2 hr incubation with occasional shaking, 4 ml of fresh medium was added directly. The next day, the medium was replaced with another 4 ml of medium containing 1 mg/ml G418 (active concentration). G418-resistant colonies were scored 12 days later. Supernatant infection of fetal liver cells was done by incubating 1–2 × 10^6 cells with 1 ml of viral supernatant containing 5 μg/ml Polybrene. After an overnight incubation, cells were washed three times and plated in methylcellulose culture. For infecting 32D cells, coculturing was done, 3 × 10^6 virus producer cells were plated in 60-mm dishes and 24 hr later, 10^6 32D cells were seeded with 5 ml of fresh medium containing 5 μg/ml Polybrene. The following day, nonadherent cells were harvested, washed three times, and plated in methylcellulose culture.

Southern blot and S1 nuclease analysis

High-molecular-weight DNA from various clones was prepared (Maniatis et al. 1982) and 10–15 μg of DNA was digested with SacI, electrophoresed, transferred onto GeneScreen Plus, and hybridized to either an M13 probe or a nick-translated IL3 or Neo probe. Hybridization was carried out at 65°C overnight. Filters were washed 1 hr at 65°C in 2 × SSC/1% SDS followed by 1 hr in 0.1 × SSC at room temperature. The IL3 probe was either an EcoRI–HindIII fragment from pTZ-IL3 or a XhoI–HindIII fragment from pCD-IL3, both of which contain the 5’ end of IL3 cDNA. The Neo probe was a nick-translated 600-bp fragment from the pSV2-neo plasmid.

For S1 analysis, total cellular RNA was obtained by lysing approximately 2 × 10^6 cells in 4 ml guanidine thiocyanate solution. This was then overlaid on top of a cushion of 2.5 ml of cesium chloride (2 g/ml) in each of six SW-41 tubes. They then spun at 32k for 16 hr. The RNA pellet at the bottom was dissolved in water. After two phenol/chloroform extractions, the RNA was precipitated and stored with ethanol at −70°C until used. S1 analysis was carried out according to Ley et al. (1982). Northern analysis was done according to Maniatis et al. (1982).

Methylcellulose culture and proliferation assay

Day-12 fetal liver cells were prepared and cultured in methylcellulose medium at 37°C for 6–8 days as described previously [Wong et al. 1983, 1982]. Adult bone marrow cells as well as 32D cells were cultured similarly, except the duration of incubation was 10–14 days. In culturing 32D cells, WEHI-3B-conditioned medium (at an optimal final concentration of 10%) was used in place of SCM (a final optimal concentration of 1% was used) and erythropoietin (Epo) at a final concentration of 1 unit/ml was not added. The preparation of SCM was done according to the following. Single-cell suspension in alpha medium (Gibco) was prepared from pooled spleens of 8- to 12-week-old BALB/c mice. Cells were adjusted and added to the culture mixture in a final concentration of 4 × 10^6/ml. The culture mixture was in alpha medium, which contained 1:300 dilution of the pokeweed mitogen stock (Gibco), 1% bovine serum albumin (Gibco), and 10^-4 M of 2-mercaptoethanol. After 4 days of incubation at 37°C in a humid incubator with 5% CO2, the supernatant was harvested after centrifugation [1000g]. Aliquots were made and stored at −20°C until used. The optimal activity was determined by both the cell proliferation assay and the clonogenic assay. The source of Epo (1 unit/ml culture) used was TcEpo from Amgen (California). Scoring criteria for various colonies were described previously [Wong et al. 1986]. Proliferation assays based on [3H]thymidine incorporation was done according to Chung et al. (1986). Conditioned media from various cell lines was collected by harvesting media exposed to cells after 2–3 days of incubation at an initial concentration of 1–5 × 10^6 cells/ml.

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P M Wong, S W Chung and A W Nienhuis

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