COLONY FORMATION BY NORMAL AND MALIGNANT HUMAN B-LYMPHOCYTES

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Summary.—A method is described that permits colony formation in culture by B lymphocytes from normal blood and from blood, marrow or lymph nodes of patients with myeloma or lymphoma. The method depends on: (1) exhaustively depleting cell suspensions of T lymphocytes, (2) a medium conditioned by T lymphocytes in the presence of phytohaemagglutinin (PHA-TCM), and (3) irradiated autologous or homologous T lymphocytes. Under these conditions the assay is linear. Cellular development of B lymphocytes can be followed; differentiation to plasma cells is seen in cultures of cells from normal individuals and myeloma patients, but not lymphoma patients. Malignant B lymphocytes in culture produced immunoglobulin of the class identified in the patient’s blood, or in freshly obtained cells. We conclude that the assay is suitable for studying the growth, differentiation and regulation of normal and malignant B lymphocytes in culture.

A number of methods have been described for obtaining colonies of normal or malignant B lymphocytes in culture. All of these have disadvantages; for example, the techniques of Fibach et al. (1976) for growing lymphoid colonies containing T cells and B cells from human peripheral blood, and that of Radnay et al. (1979) for growing normal and chronic lymphocytic leukaemia B cells require pre-incubation with pokeweed mitogen or phytohaemagglutinin (PHA) followed by plating in agar. Population changes during the preincubation phase made it difficult to obtain reliable quantitative data with those methods. Hamburger & Salmon (1977) have described a method for growing immunoglobulin-producing myeloma cells (malignant B cells). Their method requires a stimulator derived from cultures of adherent spleen cells from mice previously treated with mineral oil, and 18–21 days of incubation. Growth of myeloma cells was not always obtained with this technique, and the plating efficiency was routinely < 0.1%.

In this paper we describe a method that permits colony formation by either normal or malignant B lymphocytes by the same technique and without pre-incubation; the method also mirrors, at least to some extent, the known cellular interactions involved in the regulation of immune responses (Moretta et al., 1979). The method depends on the use, as stimulators, of medium conditioned by populations of semi-purified T cells and as well as irradiated T cells. It permits the development of colonies in 7 days with a plating efficiency that usually exceeds 1%.

MATERIALS AND METHODS

Clinical material

Blood, marrow or lymph nodes were obtained from patients with myeloma. B-cell acute lymphoblastic leukaemia (B-ALL), hairy-cell leukaemia and non-Hodgkin’s...
lymphoma. The number of patients in each diagnostic category, and the source of cells, is shown in Table I. As controls, venous blood was obtained from laboratory workers.

In each instance, the cells were shown to belong to the B-cell lineage by immunofluorescence using a panel of specific antibodies against \( \gamma \), \( \mu \), \( \alpha \), \( \kappa \) and \( \lambda \) chains; standard methods were used to demonstrate cytoplasmic (slg) or surface immunoglobulin (slg) (Preud'homme & Labaume, 1976). These data are also included in Table I. For the patients with myeloma, the immunoglobulin class found in association with the cells correlated with the M protein in sera, except for 2 cases (Table I) where M protein was not present.

Blood and marrow were collected in heparinized syringes; lymph nodes were obtained by biopsy and placed immediately in \( \alpha \)-MEM (Gibco). Cell suspensions were prepared from the latter by mincing the tissue and then teasing small fragments apart with needles. Mononuclear cells were separated from suspensions of blood, marrow or lymph node cells by centrifugation through Ficoll–Hypaque at a density of 1.077 (Boyum, 1968). Cells from patients with myeloma were cultured at this stage in the process without further cell separation. Cells from normal volunteers were depleted of adherent cells by incubation in Petri dishes in \( \alpha \)-MEM, 20% foetal calf serum (FCS, Flow Labs) for 1 h at 37°C. Then, T lymphocytes were removed from the populations as described previously (Minden et al., 1979) by using sheep erythrocytes (SRBC) to form rosettes (Wybran et al., 1973) and removing these rosettes by centrifugation in Ficoll–Hypaque. In some instances, SRBCs treated with 2–5 aminoethylisothiouronium (AET, Sigma) were used (Kaplan et al., 1976). The results with these separation procedures were equivalent and data obtained by this modification will not be presented separately. After T-cell depletion the cell suspensions contained B lymphocytes, non-T, non-B lymphocytes, monocytes and <2% T-cells. The cell suspensions so constituted were washed in \( \alpha \)-MEM and 10% FCS (growth medium) and resuspended.

To obtain T-cell populations, cells that had formed SRBC rosettes were treated with \( \text{NH}_4\text{Cl-tris} \) buffer (Hunt, 1978) washed and resuspended in growth medium.

### Table I.—Clinical material

| Pt. | Diagnosis and clinical status | Tissue studied | M-protein | Immunoglobulin typing, slg or elg (immuno-fluorescence) |
|-----|-------------------------------|---------------|-----------|----------------------------------------------------------|
| 1   | Multiple myeloma              | marrow        | G, \( \lambda \) | 80\% k (lymphoid) |
| 2 a | Multiple myeloma              | marrow        | none      | 10\% A (plasma cells) |
| b   | Multiple myeloma              | marrow        | G, k      | |
| 3 a | Multiple myeloma              | marrow        | G, k      | |
| b   | Multiple myeloma              | marrow        | G, \( \lambda \) | |
| 4   | Multiple myeloma              | marrow        | A, k      | |
| 5   | Multiple myeloma              | marrow        | 2 peaks A | |
| 6   | Multiple myeloma              | marrow        | G, \( \lambda \) | |
| 7   | Multiple myeloma              | marrow        | G, k      | |
| 8   | Plasmacytoma                  | marrow        | none      | |
| 9   | Multiple myeloma              | marrow        | G, \( \lambda \) | |
| 10  | B-cell ALL                    | blood         | --        | M, nd |
| 11  | Hairy-cell leukaemia          | blood         | --        | M, k |
| 12  | American Burkitt’s lymphoma   | blood         | --        | M, k |
| 13  | NHL                           | marrow        | --        | M, k |
| 14  | NHL                           | marrow        | G, nd     | |
| 15  | NHL                           | marrow        | M, k      | |
| 16  | Diffuse histiocytic lymphoma  | lymph node    | --        | M, k |
| 17  | Mixed cellularity lymphoma    | lymph node    | --        | M, k |

* NHL—non-Hodgkin’s lymphoma.

nd—not done.
Culture procedures

Conditioned media were prepared by incubating suspensions of T cells, prepared as described above, with 1% PHA in growth medium at 37°C, for 2 or 3 days in a moist atmosphere with 5% CO₂. Supernatants of such cultures were collected, filter sterilized, and stored at 4°C. Such preparations were termed PHA-T-cell conditioned medium or PHA-TCM. Feeder cells consisted of purified autologous or homologous T lymphocytes irradiated with 20 Gy, using a caesium irradiator. In some experiments, T lymphocytes were treated with mitomycin C (50 μg/ml, Sigma) for 30 min at 37°C washed × 3 and used as feeder cells.

Cells from the B-cell enriched fraction were prepared at concentrations from 5 × 10⁴ to 4 × 10⁵ cells/ml in the presence of 3 × 10⁵/ml irradiated T cells, 20% PHA-TCM, 0-8% methylcellulose and growth medium. After mixing the suspensions with a vortex mixer, 0-1 ml aliquots were dispersed into wells of 96-well microtitre plates (Titertek, Limbro) flat bottomed, 0-6 mm in diameter, at least 4 replicates were plated for each cell concentration. The plates were covered and the covers secured tightly with masking tape. The cultures were then incubated at 37°C for 7 days in a moist atmosphere containing 5% CO₂. Where cytoplasmic immunofluorescence measurements were required, phenol red was omitted from α-MEM.

Assessment of cultures

Colonies containing more than 20 cells were counted on Day 5 and removed from culture for characterization subsequently up to Day 7. Individual colonies were picked using finely drawn Pasteur pipettes containing a small volume of phosphate buffered saline (PBS) diluted 1:1 with distilled water. Cells from single colonies were placed on a coverslip and centrifuged with a table-top centrifuge for 2 min, or a number of colonies were pooled. Cells from single colonies were assessed for morphology using Wright-Giemsa and for myeloperoxidase using the method of Kaplow (1965). Cells from single colonies were also tested for rosette formation with SRBC (Minden et al., 1979). Cells pooled from 50–100 colonies were washed in PBS with 2% BSA and 0-2% sodium azide and then stained for cytoplasmic (cIg) or surface (sIg) immunoglobulin using standard techniques (Preud’homme & Labaume, 1976). For this purpose, labelled antibodies were purchased: fluorescein-goat F(ab)₂ anti-human IgG F(ab)₂ fragment (polyvalent), fluorescein goat IgG anti-human G, A chain and M chain respectively (Cappel Lab.); fluorescein-donkey IgG anti-human kappa chain and fluorescein-goat IgG anti-human lambda chain (Meloy Lab.). Specificity was tested using a panel of multiple myeloma cells or leukaemic B cells that showed clonal staining.

Mitogens.—Phytohaemagglutinin (PHA) (Wellcome H-15) was used at 0-5 or 1%. Pokeweed mitogen (PWM) (Gibco) was used at 1% dilution of stock, soluble protein A (SPA) from Staph. aureus was purchased from Pharmacia and used at 50 μg/ml. 2-Mercaptoethanol (Sigma) 5 × 10⁻³ M was added to cultures in some experiments.

Electron microscopy.—Colonies were picked sequentially between 3 to 8 days of culture, centrifuged through horse serum, diluted 50% with α-MEM, fixed with 3-5% glutaraldehyde in cacodylate buffer and processed by standard techniques for thin-section ultrastructural studies.

RESULTS

Colony formation

When suspensions of B cells, prepared as described in Materials and Methods, are plated under optimal conditions in wells, 2- and 4-cell clusters were seen after 24 h. By the 3rd day there were discrete colonies: these enlarged until Day 7 and became confluent and/or disintegrated. The colonies were of variable size, usually tight with some loosely attached cells at the edge of the colony. They contained from 20–500 cells (Fig. 1a). Cells within the colonies were characterized during the process of growth. By Day 3 they looked like large lymphoblasts as seen in the Wright–Giemsa preparations, but were negative for cIg. From Day 4 to Day 6 cells within colonies were heterogeneous; some continued to look like lymphoblasts while others had the characteristics of immature plasma cells by electron microscopy (Fig. 2) and Wright-Giemsa (Fig. 1b). Cytoplasmic Ig was usually first detected on Day 5 and staining for cIg reached maximum intensity by Day 7; at
this time most, but not all, cells were sIg⁺ (Fig. 1c) and typical plasma cells were seen (Figs 1d and 2b).

This sequence of events was observed for colony formation by cells from both normals and patients with B-cell malignancies; however, cells from marrow, blood or lymph nodes of lymphoma or B-cell leukaemia patients formed colonies that contained cells which remained lymphoid with scant cIg; plasma cells were seen in multiple-myeloma colonies. No peroxidase-positive cells were detected in single or pooled colony staining.

Quantitation

The assay could be used quantitatively, since a linear relationship was observed between cell number plated and colony formation, which was critically dependent upon culture conditions. At high cell densities no colonies were seen in cultures containing methyl-cellulose and growth medium only. The addition of PHA-TCM induced colony formation. But colony formation at the low cell densities required for the quantitative assay was achieved only in cultures with both PHA-TCM and irradiated autologous or homologous T-cell feeders. Fig. 3 shows the effect of adding irradiated feeders to cultures of normal B cell (3a), B-cell leukaemia cells (3b), cells from lymphoma lymph nodes (3c) and myeloma cells (3d). Only in multiple myeloma was a requirement for added feeders not found. Mitomycin C-treated T cells were as effective as irradiated T cells in normal B-cell colony formation.

Fig. 4 demonstrates the importance of PHA-TCM; in this experiment cells were plated with increasing concentrations of PHA-TCM in the presence or absence of irradiated T-cell feeders. It is apparent that optimal plating efficiency was obtained only when both stimulators were present and at concentrations of PHA-TCM > 10%. Not only was the number of colonies increased with increasing stimu-
Fig. 2.—Ultrastructure of B-cell colonies. (a) an immature cell with rough endoplasmic reticulum (Day 4). (b) a characteristic plasma cell (Day 7).
Immunoglobulin distributions in malignant B-cell colonies

Immunoglobulin classes were characterized in the cells of origin of colonies and in the cells from colonies. The data are given in Table II. It is apparent that the same Ig class was found both in freshly obtained cells and in cells from colonies. Further evidence for the origin of malignant cells was obtained in Case 1. Metaphases from fresh marrow and from colonies showed 48 chromosomes.

Fig. 3.—An increasing number of B cells was plated with (○) or without (●) a constant number of irradiated T-cells (3 x 10⁴ per well) and 20% PHA-TCM. (a) includes substitution of PHA-TCM by SPA (△), PHA (□) and PWM (■).
Fig. 4.—Dose–response curve to PHA-TCM. A constant number of cells from B-cell-enriched cell suspensions (3 x 10^4 per well) was plated with (○) or without (●) irradiated T-cells (3 x 10^4 per well) and increasing concentration of PHA-TCM.

**Table II—** B-cell colonies

| No. per 10^4 cells plated | Immunoglobulin typing (Ig or sIg) |
|---------------------------|-----------------------------------|
| Case                     |                                   |
|                           | elg or sIg                        |
|                           | (immunofluorescence)              |
| 1                        | not counted                       |
| 2 a                      | G, λ                              |
| 2 b                      | M, k                              |
| 3 a                      | M, k                              |
| 3 b                      | G, k                              |
| 4                        | G, nd                             |
| 5                        | A, k                              |
| 6                        | k (90% lymphoid)                  |
|                           | A (occasional plasma cells)       |
| 7                        | 56                                |
| 8                        | G, k                              |
| 9                        | G, λ (lymphoid)                   |
| 10                       | M, nd                             |
| 11                       | M, k                              |
| 12                       | M, k                              |
| 13                       | M, k                              |
| 14                       | G, nd                             |
| 15                       | M, k                              |
| 16                       | M, k                              |
| 17                       | M, k                              |
| nd— not done             |                                   |

**Effect of B-cell mitogens and thiols**

The lectins PWM and PHA and soluble protein A (SPA) are known to induce B-cell differentiation and proliferation in the presence of T lymphocytes in suspension cultures (Sakane & Green, 1978; Keightley et al., 1976; Janossy et al., 1977; Siegal & Siegal, 1977). In our colony assay, no colonies were obtained when only PWM, PHA or SPA were added to the plated cells. When added to a mixture of B cells and irradiated T cells, colonies formed only at the highest cell concentration with PHA and SPA; and no colonies were formed with PWM (Fig. 1a). The addition of the lectins and SPA to B cells, irradiated T cells and PHA-TCM did not significantly alter the plating efficiency.

2-Mercaptoethanol, a thiol frequently used to increase the efficiency of lymphoid colony formation (Jones et al., 1979; Hamburger et al., 1979) and required to form murine B-cell colonies (Metcalf et al., 1975) had variable and small effects on B-cell colonies and was sometimes inhibitory (data not shown).

**DISCUSSION**

The significant finding reported in this paper is the development of a quantitative method for obtaining colonies of B lymphocytes from marrow or peripheral blood of both normal and malignant origin. Cells within the colonies were identified as B lymphocytes by morphological, immunofluorescent and electronmicroscopic criteria; stages in B-lymphocyte differentiation could also be identified.

The assay has certain advantages over the procedures previously reported for obtaining B-lymphocyte colonies; no preincubation is required (Radnay et al., 1979); therefore, accounting for cells at the beginning and end of a liquid suspension culture phase is not required. Only a short (7 days) incubation period is needed, thus avoiding both the delay and increased chance of contamination inherent in longer times in culture (Hamburger & Salmon, 1977). The relationship between the number of cells plated and number of colonies formed is linear, with extrapolation through the origin; this is evidence that the culture conditions are indeed
adequate and is the basis for reliable quantitation of the colony progenitors. Finally, the procedure permits a higher plating efficiency (0.25–2.69%) than that obtained with other assays, where the reported values for myeloma are 0.001–0.1% (Hamburger & Salmon, 1977) lymphoma, 0.001–1.1% (Jones et al., 1979) and normal B cells, less than 0.014% (Radnay et al., 1979).

The assay has certain advantages that may permit its use as a tool for dissecting T-lymphocyte function. Two stimulators are required for proliferation of normal and lymphoma B lymphocytes; these are irradiated T lymphocytes (Siegal & Siegal, 1977) and a soluble factor released from T lymphocytes in suspension culture in the presence of PHA. Since colony formation is dependent, quantitatively, on these 2 factors, the method can be used as a guide in the purification of soluble factors and the identification of the active T-lymphocyte population. The observation that myeloma cells do not require the addition of irradiated T cells may provide a clue to a disorder of regulation in this disease.

Since the colonies are cultivated in methylcellulose, it is simple to retrieve cells for further analysis. Our studies have shown that morphological changes occur in cells during colony formation; it is feasible to extend this work using other markers of B-cell differentiation.

Malignant B cells, whether myeloma or lymphoma, produced only a single class of Ig. This observation served to strengthen their identification with the malignant clone in each patient. Moreover, the finding of a single Ig species provides a background against which, in the future, the diversity of Igs produced by colonies of normal B cells may be studied.

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