Acute Lymphoblastic Leukaemia: A Heterogenous Disease

D. G. HAEGERT, J. STUART, J. L. SMITH

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Summary

By using several techniques to detect surface markers on T and B lymphocytes, 11 cases of acute lymphoblastic leukaemia (A.L.L.) were studied. In four cases an insignificant number of markers were detected on the lymphoblast populations. In one case a significant number of blasts formed both sheep red blood cell rosettes and Fc rosettes, suggesting a T-cell origin for the neoplastic cells, and in another case the presence of Fc and C3 receptors on the lymphoblast population indicated a B-cell origin. In a further five cases 14-43% of the blasts had detectable surface immunoglobulin. It is concluded that A.L.L. is a heterogeneous disorder, some cases failing to express surface markers and others having either a T- or a B-lymphocyte origin or both.

Introduction

Lymphocytes derived from human bone marrow (B lymphocytes) express on their surface immunoglobulin (Ig) determinants (Pernis et al., 1971), receptors for fixed IgG (Hallberg et al., 1973), and receptors for activated C3 (Bianco et al., 1973). Thymus-derived lymphocytes (T lymphocytes), on the other hand, have an affinity for non-sensitized sheep red blood cells (R.B.C.) (Lay et al., 1971), with some T cells also expressing Fc and C3 receptors (Dickler et al., 1974). These B- and T-lymphocyte markers have been used by many workers to investigate such neoplasms as chronic lymphatic leukaemia (C.L.L.) (Pfeudhomme and Seligmann, 1972; Dickler et al., 1973; Ross et al., 1973), acute lymphoblastic leukaemia (Seligmann et al., 1972; Berella and Sen, 1973; Kessey et al., 1973), prolymphocytic leukaemia (Catsovsky et al., 1973), and leukaemic reticuloendotheliosis (Catsovsky et al., 1974 a; Haak et al., 1974).

Previously we were unable to show T- and B-lymphocyte markers on A.L.L. lymphoblasts (Collins et al., 1974) though several recent reports suggest that some cases of A.L.L. are T-cell neoplasms (Borella and Sen, 1973; Seligmann et al., 1973; Belpomme et al., 1974; Catsovsky et al., 1974 b). We described a single case of A.L.L. with two distinct neoplastic populations, one of T-cell and one of B-cell origin (Haegert et al., 1974 a), and report here our recent findings in 11 cases of A.L.L. using four rosetting reactions (Haegert et al., 1974 a), presenting evidence that in six cases the neoplastic cells expressed B-lymphocyte surface markers.

Methods

Peripheral Blood Lymphocyte Preparations.—Peripheral blood from 11 patients, aged 5-62 years, with A.L.L. (see table) was collected into heparinized bottles and the erythrocytes were sedimented with 0-6% dextran. The leukocytes-rich supernatants were centrifuged on a mixture of Ficoll and sodium metrizoate10 (Thornby and Bratell, 1970), and the lymphocytes were collected as a band at the interface. The lymphocyte suspensions were washed three times with Hepes-Hanks's balanced salt solution supplemented with 0-2% bovine serum albumin then made up to a final suspension of 2 x 10^6 cells in Hepes-MEM (Eagle's medium) with 0-2% bovine serum albumine. The final suspensions contained less than 5% nonlymphoid cells.

Rosette Tests.—To detect lymphocytes with affinity for non-sensitized sheep R.B.C. a sheep R.B.C. rosetting technique was used. Fc receptors, C3 receptors, and Ig determinants were detected by rosetting reactions (Haegert et al., 1974 a). For scanning purposes

References


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cytocentrifuge preparations of the rosette tests were made then stained with Leishman's stain (Collins et al., 1974; Smith and Haegert, 1974).

Enzymatic Digestion.—Aliquots of the washed lymphocyte suspensions of a single patient (case 11) were treated with Vibrio cholerae neuraminidase (Merck) and Zeller, 1974). The treated cells were then washed immediately and rosetted in all four rosetting experiments. In other experiments from this same patient were treated with trypsin (2.5 g/l for 30 min. at 37°C) (Preud'Homme and Seligmann, 1972) then washed and left overnight in bicarbonate-buffered M.E.M. with 10% heat-inactivated calf serum at 37°C before rosetting in all four test reactions.

Immunofluorescent Staining.—Polyvalent rabbit anti-human Ig antibodies were raised in the laboratory. Fluorescein-labelled goat anti-rabbit Ig was obtained from Nordic Pharmaceuticals, Tilburg, Holland. Membrane fluorescence was performed using the indirect technique. Control preparations using normal rabbit serum were included in all experiments. All preparations were kept on ice then scored under a phase microscope fitted with an ultraviolet epiluminescent lamp.

Results

Peripheral blood lymphocyte preparations from 11 patients with A.L.L. contained variable percentages (8-93%) of blasts. By using a cytocentrifuge technique (Collins et al., 1974; Smith and Haegert, 1974) blast populations could be clearly defined except in the mixed antiglobulin test for surface Ig (M.A.G. reaction) in which prior formaldehyde fixation of the lymphocytes interfered with Leishman staining. In control rosetting reactions with unsensitized ox cells and ox cells coated with IgM anti-ox R.B.C. antibody less than 4% of the blasts formed rosettes. Control preparations for the indirect immunofluorescence experiments gave essentially negative reactions, with less than 1% of the blasts staining.

In four cases significant numbers of surface markers could not be detected on the lymphoblast populations (<5%; see table). In one case (case 3) a significant number (>10%) of lymphoblasts formed sheep R.B.C. rosettes and Fc rosettes, but in no other case did a significant number of blasts form sheep R.B.C. rosettes. In cases 8 and 9 a significant number of blasts had Fc receptors, while in case 9 a significant number of blasts also had C3 receptors. In five cases (cases 5, 6, 7, 8 and 11) the total number of cells with surface Ig determinants (detected either by the M.A.G. reaction or by immunofluorescence) significantly overlapped the number of blasts in the preparations examined, which suggested that 14-43% of the neoplastic cells had surface Ig determinants (see table).

In case 11 treatment of the A.L.L. cells with neuraminidase did not increase the number of cells reacting in the rosette tests. After treatment of an aliquot of lymphoid cells with trypsin followed by overnight incubation there was no change in any of the four surface markers evaluated.

Discussion

We examined 11 cases of A.L.L. for four surface markers using a previously developed cytocentrifuge technique (Collins et al., 1974; Smith and Haegert, 1974) which can yield important information from leukaemic populations containing few or many neoplastic cells, principally because we can characterize cylogically the lymphoid cells forming rosettes. In four cases less than 5% of the lymphoblasts formed sheep, Fc, or C3 rosettes, and there was no evidence of surface Ig on these cells. The meaning of a few blasts forming rosettes in the marker tests is obscure and probably was a result of our technique. Cytocentrifugation of control A.L.L. preparations in the Fc, C3, and M.A.G. rosette tests produced non-specific erythrocyte rosettes around less than 4% of the lymphoblasts.

In one case over 10% of the lymphoblasts formed sheep erythrocyte and Fc rosettes. Though surface Ig was not evaluated and experiments were not performed to see if these markers were present on the same cells the findings suggest a T-cell origin for the blast population. This case seems analogous to those of T-cell lymphomas in mice which express Fc receptors (Greys et al., 1972; Harris et al., 1973). In another case the lymphoblasts formed significant numbers of Fc and C3 rosettes, while the M.A.G. test did not suggest that significant numbers of these cells had surface Ig. Since some normal B cells have readily detectable C3 receptors but not Ig determinants (Ross et al., 1973) these findings suggest a B-lymphocyte origin for the neoplastic cells.

In a further five cases of A.L.L. detection of surface Ig by either the fluorescent antibody or M.A.G. test did suggest that a significant number of the blast cells had surface Ig (the number of cells with surface Ig being too great to be accounted for by the normal population alone). In these cases a few lymphoblasts also rosetted in the Fc and C3 reactions. Since the controls in the M.A.G. test and controls stained with fluorescein-labelled antisera and normal sera in the indirect test gave essentially negative reactions (<5%) the detected Ig seemed not to be cytophilic antibody. Moreover, trypsin digestion (Preud'Homme and Seligmann, 1972) of the lymphoblast population in one case followed by overnight incubation did not decrease the number of Ig-positive cells detected. All this evidence suggests that in this case at least the lymphoblasts expressed surface Ig and were of B-cell origin. Treatment of

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**Table:**

| Case No. | Type of specimen at Presentation (P) or Relapse (R) | Findings at Presentation | % Blasts in Lymphocyte Preparation | T-cell Marker (Surface Ig FR E B.E. B.C. Ro) | Surface Markers | B-cell Markers | Total (% B + T) | % Blasts (% B + T)1
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<tbody>
<tr>
<td>1</td>
<td>P.B. (R.)</td>
<td>16</td>
<td>6.3</td>
<td>50</td>
<td>8</td>
<td>2 (33)</td>
<td>(0)</td>
<td>20</td>
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<tr>
<td>2</td>
<td>P.B. (R.)</td>
<td>15</td>
<td>35</td>
<td>95</td>
<td>71</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>21</td>
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<td>3</td>
<td>R.M.</td>
<td>62</td>
<td>1</td>
<td>2.3</td>
<td>66</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>12</td>
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<tr>
<td>4</td>
<td>P.R.</td>
<td>60</td>
<td>90</td>
<td>6 (%)</td>
<td>0 (0)</td>
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<tr>
<td>5</td>
<td>P.B.</td>
<td>62</td>
<td>7</td>
<td>52</td>
<td>88</td>
<td>0 (0)</td>
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<tr>
<td>6</td>
<td>P.B.</td>
<td>25</td>
<td>71</td>
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<td>7</td>
<td>P.B.</td>
<td>2</td>
<td>14</td>
<td>18</td>
<td>78</td>
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<tr>
<td>8</td>
<td>P.B.</td>
<td>6</td>
<td>17</td>
<td>66</td>
<td>89</td>
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<tr>
<td>9</td>
<td>P.B.</td>
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<td>43</td>
<td>50</td>
<td>93</td>
<td>0 (0)</td>
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*Values in parentheses represent total percentage of cells, both neoplastic and non-neoplastic, reacting with given indicator system. Values not in parentheses represent percentage of neoplastic cells reacting with given indicator system.*

†Surface Ig determinants were recognized by indirect immunofluorescence.

‡% Sum of percentage of total cells, neoplastic and non-neoplastic, with T-lymphocyte marker and highest percentage of cells with any B-cell marker.

§% Sum of percentage of blasts in preparation and total percentage of cells recognized by both T and B markers.

N.D. = Not done

Conversion: 1 ml to Traditional Units—W.B.C. 1 x 109/l = 1000/mm3.
these lymphoblasts with neuraminidase (Chapel, 1973; Mehrishi and Zeiller, 1974) did not increase the number of cells with markers and failed to show masked determinants.

It is noteworthy that four out of the five patients with detectable Ig on the surface of their lymphoblasts were investigated at presentation before treatment. Further studies are planned to investigate if these lymphoblasts with surface Ig persist during the course of the disease. The findings in one patient (case 11) suggest that they do. Apart from this observation there was no correlation between the presence or absence of markers with the prognostic assessment at diagnosis, based on leucocyte count, percentage blasts, and presence or absence of mediastinal enlargement on chest x-ray picture.

Nevertheless, important information can be obtained by characterizing surface markers on A.LL lymphoblasts. Such an approach may be clinically important in the early classification of cases of A.LL since there is some evidence that cases of T-cell origin have an exceptionally poor prognosis (Borella and Sen, 1973; Catovsky et al., 1974 b). The important observation that some A.LL lymphoblasts express B-cell markers, together with other recent findings (Haegert et al., 1974 b), suggest that A.LL is a disease of heterogeneous origin divisible into at least four groups—those without surface markers, those with T- or B-cell markers, and those cases expressing combined T- and B-cell determinants (Haegert et al., 1974 b).

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 References


Thymus transplantation in patients with DiGeorge's syndrome (August et al., 1968; Cleveland et al., 1968) has been shown to produce rapid and dramatic reconstitution of cell-mediated immunity. Temporary reconstitution of cell-mediated immunity (Levy et al., 1971) but not of humoral immunity (Amman et al., 1973) has been achieved in various immunodeficiency diseases. We report the successful grafting of a fetal thymus into a 6-year-old boy with T-lymphocyte deficiency proved on three occasions before the operation.

Case Report

A 6-year-old Caucasian boy was admitted to hospital with a history of repeated respiratory infections. He was regularly off school three days out of four and was almost continuously on antibiotics. He was an adopted child and had a half-brother aged 7 years, also adopted by the same family. They had the same mother but different fathers.

The patient had normal facies and none of the features of DiGeorge's syndrome. He was apparently well until the age of 1 year, when he began to have febrish illnesses with coughs, diagnosed as bronchitis, which responded to antibiotics. Various antibiotics had been prescribed over a relatively short period. His doctor had tried on one or two occasions to leave him without antibiotics but he became rapidly worse and eventually had to be given antibiotics. He showed no eczema and sneezing was not a feature though wheezing was common. At one stage he developed severe allergy to streptomycin. He received quadruple immunization, oral polio vaccine, and, later, measles vaccination. His mumps and chickenpox were no worse illnesses than those of his brother and he adopted had not had measles. He had frequent attacks of pneumonia and clearly had a much higher tendency to infections than did his brother.

Investigations

Methods and Materials

Lymphocyte transformation to phytohaemagglutinin (PHA) was done according to the standardized method of Yamamura (1973) and to candida immunogen by the standardized method of

MEDICAL MEMORANDA

Successful Thymus Graft for T-cell Deficiency in a 6-year-old Boy

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