NATURAL CYTOTOXICITY OF HAEMOPOIETIC CELL POPULATIONS AGAINST MURINE LYMPHOID TUMOURS

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Summary.—Homozygous nude and normal mice of 3 strains, BALB/c, CBA and C57BL, were used as sources of nucleated haemopoietic “natural killer” (NK) cells. These killer cells could lyse a wide range of syngeneic and allogeneic lymphoid tumour cell lines in vitro, and it was found that cell suspensions from nude mice were always significantly more active than those from normal mice, and that the most active effector population was a polymorph-enriched peritoneal-exudate cell suspension. Eosinophils did not appear to be involved in the phenomenon, and mononuclear peritoneal-exudate cell suspensions were actually highly inhibitory. Three non-lymphoid tumours, a carcinoma, a fibrosarcoma and a mastocytoma, were totally resistant to in vitro lysis. Although all susceptible tumour cell lines were C-type virus-associated, not all of these tumours were killed by all strain sources of spleen cells, indicating a specificity of killing.

Reports of the in vitro lysis of tumour cells by haemopoietic cells from unimmunized animals and humans have appeared sporadically in the literature. Cell-mediated cytotoxicity, as assayed by the lysis of tumour cell lines, has been reported in a number of different systems using human blood leucocytes (Takasugi, Mickey and Terasaki, 1973; Petranyi et al., 1974), mouse spleen cells (Herberman et al., 1973; Greenberg and Playfair, 1974) and rat spleen cells (Nunn et al., 1973; Holterman, Klein and Casale, 1973) from normal donors. In 1975 reports from several laboratories clearly demonstrated the killing of various mouse lymphoma lines by spleen-cell suspensions from non-immune animals. Thus, Zarling, Nowinski and Bach (1975) drew attention to the fact that $^{51}$Cr release from a $^{51}$Cr-labelled AKR leukaemic cell line resulted from exposure to spleen cells from low, but not high, leukaemia strains of mice. Kiessling, Klein and Wizgell (1975a) demonstrated that murine spleen cells killed particular oncogenic-virus associated tumours in vitro, but did not lyse all such tumour cell lines, and this “specificity” of killing was subsequently confirmed by other workers (Herberman, Nunn and Lavrin, 1975a; Sendo et al., 1975).

Current interest has focused on three aspects of this phenomenon: the identity of the “natural killer” (NK) cell, the genetic control of the phenomenon and the nature of the antigenic determinants recognized by the NK cells. To date, the NK cell has been characterized mainly in terms of negatives. Thus, it does not express markers of mature T or B cells (Kiessling et al., 1975b; Herberman et al., 1975b) nor the properties of “classical” adherent macrophages (Kiessling et al., 1975b; Herberman et al., 1975b). It does not manifest antibody-dependent cell-mediated cytotoxicity in the standard antibody-coated chicken-red-cell assay (Kiessling et al., 1975b; 1976; Herberman.

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et al., 1975b) although recent studies with a more sensitive system have suggested that the NK cell may express Fe receptors (Herberman et al., 1977) which further emphasizes the need more strictly to compare these cells with K-cell function. To date, its positive qualities are a declining presence with increasing mouse age (Kiessling et al., 1975a, b) a specific tissue distribution (Kiessling et al., 1975a) an increased incidence in nude mice (Kiessling et al., 1975b) and a lability in tissue culture at 37°C (Herberman et al., 1975b).

Recent studies have identified a specific cell-surface alloantigen on the NK cell (Glimcher, Shen and Cantor, 1977) and may provide an approach to its eventual isolation and precise characterization. Studies in a human system have indicated that peripheral human blood leucocytes with natural cytotoxic activity have complement receptors and are not T cells or B cells (Pross and Jondal, 1975); however, the murine natural killer cells lack complement receptors (Kiessling et al., 1976).

Studies of the genetic control of this phenomenon have indicated that different mouse strains can manifest either high or low levels of "natural" cytotoxicity against particular tumour cell lines when their spleen cells are tested in vitro (Zarling et al., 1975). More detailed analysis with congenic mouse lines and backcross analysis has indicated that the phenomenon is under polygenic control, at least one gene of which is linked to the major histocompatibility complex (MHC) (Petranyi, Kiessling and Klein, 1975; Kiessling et al., 1975c), although our own studies described here do not demonstrate any evidence of MHC involvement in the responses studied.

This NK-cell phenomenon was first defined with a Rauscher-virus-induced leukaemia (Herberman et al., 1973) and the possible specificity of NK cells for virally determined tumour-associated antigens (TAA) has been widely investigated. It is unlikely that histocompatibility antigens themselves are involved in the antigenic site, as has been suggested for TAAs in other systems (Germain, Dorf and Benacerraf, 1975; Schrader and Edelman, 1976) since 51Cr-labelled tumour target cells allogeneic to the NK cells are readily lysed (Zarling et al., 1975; Kiessling et al., 1975a) and unlabelled allogeneic tumour cells can also successfully compete in cellular competitive-inhibition assays with 51Cr-labelled tumour cells (Zarling et al., 1975; Kiessling et al., 1975a). Since normal lymphoid cells do not inhibit in such assays it has been inferred that normal tissue antigens are not involved (Zarling et al., 1975). The viruses so far implicated belong to both the murine leukaemia virus (MuLV) and the Moloney sarcoma virus (MSV) groups (Herberman et al., 1973; Zarling et al., 1975; Herberman et al., 1975a); however, no specific virion-or viral-cell-surface-associated antigens have yet been definitely designated as the determinants for NK cells. Comparison of the specificity of NK cells and cytotoxic T (Tc) cells from mice immune to MSV-induced tumours has indicated that different antigenic determinants are recognized by these different cell types (Herberman et al., 1976).

In this paper we report our investigations into the nature and specificity of this phenomenon, with particular reference to a possible role for polymorphonuclear leucocytes in the in vitro lysis of 51Cr-labelled tumour cells.

MATERIALS AND METHODS

Tissue Culture Medium (DMEF).—Dulbecco's modified Eagles Medium (CSL, Melbourne, Australia) supplemented with 10% foetal calf serum (CSL, Melbourne, Australia) was used.

Mice.—Inbred mice obtained from the Hall Institute specific-pathogen-free colonies were: BALB/c An Bradley WEHI, BALB/c nu (N8), C57BL/6J WEHI, CBA/CaH WEHI, CBA.nu (N10) and (BALB/c × C57BL) F1 hybrids. Strains from the Hall Institute conventional colonies were: C57BL/6.nu (N10) (obtained from Dr I. Lefkovits, Basel, Switzerland) B10.D2/n Sn and B10.BR/Sg Sn
(obtained from Jackson Laboratories, U.S.A. in 1974) and BALB/c H-2b, BALB/c H-2k and BALB/c H-2k (all 3 obtained in 1975 from Dr F. Lilly, New York, U.S.A.). All mice were 5–8-week-old males, unless otherwise indicated.

Tumour cells.—The tumour cell lines were propagated in continuous tissue culture as previously described (Horibata and Harris, 1970) and are listed in Table I. Their origins, with the exception of WEHI-265, have all been described in recent publications from this laboratory (Chism, Burton and Warner, 1976; Burton and Warner, 1977). WEHI-265 is a BALB/c tumour of the granulocyte-macrophage cell lineage and was induced by Abelson virus (Warner, N. L., Harris, A. W., Gutman, G., Metcalf, D., Burgess, A. W., Warr, G. W. and Haustein, D., manuscript in preparation).

Cell suspensions.—Spleen-cell suspensions were prepared by teasing and mincing spleens from mice killed by cervical dislocation through an 80 gauge stainless-steel sieve into ice-cold DMEM. After thoroughly pipetting the suspensions, it was layered with FCS for 10 min in ice, to remove cell clumps and large debris. The supernatant was then transferred to another tube, layered again with FCS, and centrifuged at 1500 rev/min for 5 min, to pellet the cells and remove fine debris. The cells were then resuspended in 5 ml of 0.17M ammonium chloride, prewarmed to 37°C, to lyse the red cells. One min later a further 5 ml of ice-cold DMEM was added and the cells pelleted by a further light centrifugation. Finally, the cells were resuspended in ice-cold DMEM and stored on ice, while their viability was determined by an eosin-dye-exclusion count in a haemocytometer. The cell suspension was then adjusted as necessary for the assay. This technique is a modification of that described by Shortman, Williams and Adams (1972) and usually results in a nucleated erythrocyte-free spleen-cell suspension that is at least 80% viable.

Lymph-node suspensions were prepared from pooled mesenteric and peripheral lymph nodes in the same manner as for the spleen-cell suspensions, except for the omission of the ammonium chloride treatment.

Thymus-cell suspensions were prepared by treating thymus lobes in the same manner as lymph node cell suspensions.

Marrow-cell suspensions were prepared by aspirating the femoral marrow with a 21-g needle and suspending the cells in ice-cold DMEM by vigorous pipetting. The cell suspension was then treated in an identical manner to the spleen-cell suspension.

Proteose-Peptone-Broth Peritoneal-Exudate Cells: PPB-PEC.—These suspensions were prepared by injecting the appropriate mice i.p. with 1 ml of 10% proteose peptone broth (PPB) (Difco Laboratories, Detroit, Michigan) 3–4 days prior to harvest. The mice were killed by cervical dislocation and 3–5 ml of ice-cold 0.02M phosphate-buffered saline (PBS) was injected into the peritoneal cavity. This was then aspirated with a pipette, and the cells pelleted by light centrifugation. The pellet was resuspended in 1–2 ml of ice-cold DMEM and a viability count made. This procedure, which is based on a previously described technique (Katz and Unanue, 1973) produces a cell suspension that is composed of at least 85% large mononuclear cells.

Polyvinylpyrrolidone Peritoneal-exudate Cells: PVP-PEC.—These suspensions were prepared by injecting 2 doses i.p. of 1 ml of 15% polyvinylpyrrolidone (PVP) (British Drug Houses, Melbourne, Australia, Stock 29579) in PBS 15 h and 2 h prior to harvest, in the appropriate mice. The cell suspension was then harvested and treated in the same way as for the PPB-PEC. The use of PVP in this way (Lord, 1975) produces a cell suspension which is usually at least 80% poly-morphonuclear leucocytes (polymorphs). Considerable variation has been found between various batches of PVP in their ability to induce a primarily granulocytic exudate with cytotoxic activity, so that individual batches of PVP must be screened for this ability.

Peripheral blood leucocytes.—These were prepared according to a modification of the method of Davidson and Parish (1975). Briefly, the mice were exsanguinated by eye bleeding into tubes containing 0.5 ml of 0.8% sodium citrate solution as anti-coagulant. The blood was then divided into 5 ml aliquots, 20 μl of 25% sodium azide added (final concentration 0.1%), and the blood warmed to 20°C. It was then layered over 4–0 ml of Isopaque/Ficoll (prepared as described by Davidson and Parish, 1975) in U-bottomed glass siliconized centrifuge tubes. The tubes were placed in a centrifuge at room temperature and spun at 2000 g for 15 min, with rapid acceleration to 2000 g in 20 sec. After centrifugation the supernatant above the Ficoll/
Isopaque interface was discarded and the white-cell layer at the interface, together with the separating medium above the red cell/dead cell pellet, collected. The cells were then washed twice in DMEF, a viable cell count performed by eosin-dye exclusion, and the suspension adjusted for the assay.

$^{51}$Cr-release assay.—The techniques for $^{51}$Cr-labelling tumour cells have been previously described in detail (Burton, Thompson and Warner, 1975) and are, therefore, only outlined here. In general, tumour cells were incubated for 30 min in DMEF, at $37^\circ$C in a humidified $10\%$ CO$_2$ incubator, at a maximal cell concentration of $10^7$/ml and $^{51}$Cr (sodium chromate, sp. act. 50–200 $\mu$Ci mol, CEA, Gif-Sur-Yvette, France) concentration of 100 $\mu$Ci/ml. The cells were then washed $\times$3 in warm DMEF, counted in a haemocytometer, adjusted to $25 \times 10^4$/ml, and stored at room temperature until used.

The details of the $^{51}$Cr-release assay have also been reported in detail (Burton et al., 1975; Chism et al., 1976). Briefly, 2-fold serial dilutions of the haemopoietic nucleated cytotoxic cells (CL) were made to give final CL/$^{51}$Cr-labelled tumour cell (CL/T) ratios in the assay over the range 200/1–6.25/1. The assay was performed as 4 replicates of each CL/T ratio in microtitre trays (Microtest 11 Tissue Culture Plate, Falcon Plastics, Oxnard, California, U.S.A.) and the $^{51}$Cr-labelled tumour cells were added to each well at a constant number of $25 \times 10^3$ cells. The total assay volume was 200 $\mu$l, and the trays were then incubated for 4 h at $37^\circ$C, and for 1 h at $45^\circ$C, in humidified $10\%$ CO$_2$ incubators. At the end of this period 100 $\mu$l of the supernatant was removed from each well and counted in a Beckmann Biogamma scintillation counter. The cytotoxicity was calculated as:

% Specific lysis =

$$\text{Test count} - \text{Background count} \times \frac{100}{\text{Maximal count} - \text{Background count}}$$

where the Background count is the $^{51}$Cr release from $25 \times 10^3$ labelled tumour cells incubated alone, and the Maximal count that of the same number of tumour cells lysed with Zaponin (Coulter Electronics Ltd., Dunstable, Beds., U.K.).

As discussed in a current publication (Burton and Warner, 1977) a level of specific lysis of a particular tumour can be considered significant when it is at least as high as the mean background, as listed in Table I. These figures remain remarkably constant for each cell line as long as cells in the log phase of their growth are used. This convention corresponds to a level of statistical significance of at least 1% ($P < 0.01$, Student's t-test). Furthermore, a difference between 2 levels of specific lysis is considered significant in this assay if the higher is 1.5 times the lower. This corresponds to a level of statistical significance of 1% or better (Student's t-test).

**Ascaris suum-infected mice.**—In order to prepare leucocyte suspensions that were enriched for eosinophils, BALB/c male mice were infected orally with embryonated *Ascaris suum* eggs. The mice were killed 11 days later and spleen and blood leucocyte suspensions prepared as described before. This technique results in an eosinophilia of at least 20% in the peripheral blood (Mitchell et al., 1976).

### Table I.—Tumour Cell Lines used as Targets for Natural Cytotoxic Cells

| Tumour | Strain of origin | Type            | H-2 type | % Background $^{51}$Cr release* |
|--------|-----------------|-----------------|----------|-------------------------------|
| WEHI-7  | BALB/c          | T lymphoma      | d        | 9                             |
| WEHI-22 | BALB/c          | T lymphoma      | d        | 10                            |
| ABE-8   | BALB/c          | B lymphoma      | d        | 15                            |
| WEHI-265| BALB/c          | Myeloid leukaemia| d        | 20                            |
| WEHI-164| BALB/c          | Fibrosarcoma    | d        | 15                            |
| MPC-11  | BALB/c          | Plasmacytoma    | d        | 22                            |
| EMT-6   | BALB/c          | Carcinoma       | d        | 10                            |
| HPC-10  | NZB             | Plasmacytoma    | d        | 15                            |
| P815    | DBA/2           | Mastocytoma     | d        | 17                            |
| EL4     | C56BL           | T lymphoma      | b        | 14                            |
| Cl.18   | C3H             | Plasmacytoma    | k        | 20                            |
| RILQ    | CBA             | T Lymphoma      | k        | 19                            |

* Mean spontaneous $^{51}$Cr release by the various tumour lines under the assay conditions, calculated as:

$$\frac{\text{Background count}}{\text{Maximal count}} \times 100$$

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Differential cell counts.—Smears were made of various effector-cell preparations, and were stained with Giemsa stain.

RESULTS

Variability of cytotoxicity in the one system.

The NK activity of spleen cells from BALB/c. nu mice for the T lymphoma WEHI-7 is represented in Fig. 1, with CL/T ratios of 200/1–12.5/1. There was a wide day-to-day variation in the cytotoxicity measured over 20 experiments. However, for any particular experiment a CL/T curve of the form illustrated for a representative sample of experiments was always found. CL/T ratios of 200/1 and 12.5/1 gave low levels of cytotoxicity, and the peak level was almost invariably at 100/1 or 50/1. Although these are not distinguished in the figure a number of experiments were performed using age-matched 5–8-week-old male and female BALB/c nudes. There was no significant difference in the cytotoxicity, as measured by % specific lysis of 51Cr-labelled WEHI-7, between the 2 sexes.

Variation in cytotoxicity with CL source.

There was a wide variation in cytotoxicity when nucleated cells from various tissue sources were tested on the same target at the same time (Table II). As can be seen, the BALB/c. nu spleen was the most potent source of CL, while the BALB/c thymus was totally without reactivity. Of particular note is the low reactivity of marrow. If the NK cell is a marrow-derived cell, either its activity is suppressed or it must undergo some maturation process after leaving the marrow. The relative inactivity of PPB-PEC, a rich source of macrophages, argues against this cell type being the CL in this system.

Strain variation in cytotoxicity to one tumour target.

Some evidence of genetic control of this effect was obtained (Table III), although

Table III.—Comparison of the Cytotoxicity of Spleen CL from Various Mouse Strains for 51Cr WEHI-7 Tumour Cells

| Strain | No. expts. | Mean % specific lysis* |
|--------|------------|------------------------|
| BALB/c. nu | 20 | 40 |
| BALB/c | 8 | 22 |
| CBA. nu | 6 | 36 |
| CBA | 3 | 10 |
| C57BL. nu | 3 | 19 |
| C57BL | 3 | 11 |
| BALB/c. H-2b | 2 | 29 |
| BALB/c. H-2k | 2 | 27 |
| BALB/c. H-2g | 2 | 13 |
| B10. D2 | 3 | 13 |
| B10. Br | 3 | 7 |
| B10. A | 2 | 15 |

* All at CL/T ratios of 50/1.

The precise genetic basis remains obscure. Spleen CL from nude mice were always significantly more active than those from normal mice in the 3 strains compared, indicating complete lack of involvement of T lymphocytes in this phenomenon. There was also some suggestion that an H-2-linked gene is important, as in both groups of H-2-congenic normal mice a difference in cytotoxicity between strains was seen. Thus, the BALB/c. H-2b, and H-2k spleen CL showed higher cytotoxic activity than CL from the H-2k congenic strain, whereas in the C57BL congenic strains, the B10.A(H-2b) spleen CL gave significantly higher lysis than the B10.Br (H-2k) strain.

Comparisons of such congenic strains
would, at face value, appear to indicate the role of an H-2-linked gene in controlling levels of cytotoxic activity. However, no correlation with a particular H-2 allele was seen across the different strains, in that whereas BALB/c, H-2k was quite active, B10.Br and CBA were not. Although this could indicate that the effect is due to a gene only loosely linked to the H-2 region, the variations observed even within the one strain (Fig. 1) indicate that some caution might be made in assuming a direct genetic immune response control. Environmental influences may appreciably affect the cellular composition of lymphoid organs as a result of antigenic confrontations, and these may be due in part to H-2-linked Ir gene effects. Thus, the apparent influence of the MHC region in the present phenomenon may be quite indirect.

**Lysis of H-2 incompatible targets by NK cells**

Spleen CL from 3 different H-2 normal and nude mouse strains readily lysed both H-2-compatible and incompatible 51Cr-labelled tumour cells (Table IV). The highest levels of specific lysis were in fact seen across H-2 differences, i.e. CBA.nu (H-2k) caused 45 ± 1% specific lysis of 51Cr WEHI-7 (H-2d) and 39 ± 3% specific lysis of 51Cr EL4 (H-2b). However, there was no evidence that the lysis of syngeneic tumour cells was either consistently less or greater than that of the allogeneic tumour cells. It is also to be noted that in most cases the degree of lysis of a particular tumour by spleen cells from nude mice of a particular strain, is greater than that observed from the normal mice of that strain. Whether this is due simply to an increased proportion of NK cells, by virtue of the absence of the T-cell population, or due to removal of a suppressor cell population, has not

**Table IV.**—**NK-cell Activity of Spleen CL Directed against H-2-Compatible and H-2-Incompatible Tumour Cells**

| Tumour       | BALB/c | CBA | C57BL |
|--------------|--------|-----|-------|
|              | Nude   | Normal | Nude | Normal | Nude | Normal |
| WEHI-7       | 34±1   | 14±1   | 45±1 | 17±4   | 26±1 | 23±1   |
| (H-2d)       |        |        |      |        |      |        |
| HPC-10       | 13±1   | 9±2    | 13±1 | 11±2   | 17±2 | 9±1    |
| (H-2h)       |        |        |      |        |      |        |
| EL4          | 22±2   | 1±1    | 39±3 | 24±3   | 27±3 | 0      |
| (H-2b)       |        |        |      |        |      |        |
| CL.18        | 32±2   | 12±1   | 34±2 | 6±2    | 15±2 | 7±3    |
| (H-2k)       |        |        |      |        |      |        |

* All values were obtained with CL/T ratios of 50/1.
been determined. However, in all instances, various CL/T ratios have been used, and the differences indicated between the respective nude and normal spleen-cell preparations are observed at all levels of the CL/T lysis curve.

The range of tumour-cell lysis by NK cells

The lysis of a number of \(^{51}\text{Cr}\)-labelled tumour cells, all lymphomas or leukaemias, was observed with spleen CL from 3 different strains (Table V). It is again apparent that there is a strain and tumour variation in the results obtained, but that incompatibilities in the major histocompatibility complex (MHC) do not restrict lysis. Of particular interest was the observation that certain combinations, even with nude-mouse-derived spleen CL, showed no lysis. Since each preparation of spleen cells contained good cytotoxic activity on at least some of the tumours, and as all tumours were capable of being lysed by at least one of the CL preparations, it must be concluded that lysis, where observed, is not due to the action of a non-specific toxic cell or substance in the preparation. The existence of the negative combinations is more compatible with the concept that lysis is due to a specific recognition, albeit of undefined nature. It is to be stressed that although some variability in levels of lysis have been observed in combinations where lysis is indicated (e.g., CBA.nu cells with EL-4 19\% lysis, as contrasted to Table IV, 39\% lysis) the combinations indicated by a 0 have consistently failed to involve any lysis on many repeat experiments.

Resistance of certain tumours to lysis by natural CL

Most of the evidence for the specificity of the NK-cell phenomenon has come from comparisons between the lysis of different tumour types. This point is illustrated both by ranking the various lymphomas in order of their susceptibility to NK cell killing (WEHI-7, EL4, ABE8, RILQ and WEHI-22; Tables IV and V) and by the demonstration that some tumour lines are resistant to NK cell lysis (e.g., EMT-6, WEHI-164 and P815; Table VI). These latter 3 tumours are all easily lysed in other systems (P185 in the \textit{in vitro} allograft reaction (Burton et al., 1975) and all 3 in the \textit{in vitro} oncofoetal reaction (Chism et al., 1976)) also indicating that the specificities involved in the NK reaction are probably not of oncofoetal type. However, they are totally resistant to lysis by spleen CL from the most active strains. These tumours are spontaneous or carcinogen-induced, and are thus probably \textit{non}-viral in origin.

The nature of the effector cell

As reviewed herein, the nature of the haemopoietic cell mediating the NK-cell phenomenon is not known. Experiments were conducted which confirmed the findings from other laboratories, referred to earlier (i.e. the NK cells lack T- or B-cell markers and are not adherent to plastic). Since the readily identifiable lymphocyte types, adherent macrophages and mononuclear peritoneal-exudate cells, seemed to have been excluded as candidates for the NK cell, the possibility that polymorphonuclear leucocytes were involved was considered. Therefore, experiments were conducted with neutrophil-enriched populations of peritoneal-exudate cells (PVP-PEC) and with specifically eosino-
phil enriched populations, namely spleen and blood leucocytes from *Ascaris suum*-infected mice.

It was found that the PVP-PEC were significantly more cytotoxic than spleen cells, and very much more active than PPB-PEC (Fig. 2) and that this applied equally well with cells from nude or normal BALB/c mice. The proportion of polymorphs (mainly neutrophils) in the PVP-PEC ranged from 70–90%. The specificity of the PVP-PEC was similar to that of nude spleen CL, indicating that the same effector cell (i.e. the NK cell) was probably involved. Thus PVP-PEC only lysed *in vitro* lymphoid and leukaemia cell lines, and spontaneous and carcinogen-induced tumours were not killed. Furthermore, these PVP-PEC lost their cytotoxic activity after overnight incubation in tissue-culture medium, a procedure which also killed most of the polymorphs.

These results suggested that a polymorphonuclear leucocyte cell type may play a role in this phenomenon. This interpretation, however, does not appear compatible with the data which indicated that marrow-derived cells had only marginal NK-cell activity. One possible explanation for the failure of these cells to mediate lysis was that other mononuclear cell types present in the marrow might actually inhibit their activity in the 51Cr-release assay. Accordingly, a mix assay was performed with peritoneal-exudate cells. In this experiment (Fig. 3A) and 4 other similar ones, higher cytotoxicity of BALB/c.nu PVP-PEC was verified by comparing PVP-PEC with PPB-PEC over the same range of CL/T ratios on 51Cr WEHI-7. The PPB-PEC were then mixed at varying CL/T ratios (32/1–1/1) with PVP-PEC, which were held at a constant CL/T ratio of 8/1. It was found that the PPB-PEC were inhibitory to the cytotoxicity of the PVP-PEC over the whole range of CL/T ratios of PPB-PEC used (Fig. 3B). The effect diminished as the number of PPB-PEC mixed with the PVP-PEC diminished, but the level of lysis of 51Cr WEHI-7 mediated by the PVP-PEC at a CL/T ratio of 8/1 was still

| Spleen-cell source | No. expts. | EMT-6 | No. expts. | WEHI-164 | No. expts. | P815 |
|--------------------|------------|-------|------------|----------|------------|------|
| BALB/c.nu          | 5          | 0     | 4          | 2 ± 1    | 2          | 1 ± 1|
| BALB/c             | 4          | 0     | 2          | 2 ± 1    | 2          | 0    |
| CBA.nu             | 1          | 0     | 1          | 0        | 1          | 0    |
| CBA                | 2          | 0     | 1          | 0        | 2          | 0    |
| C57BL.nu           | 1          | 0     | 1          | 3 ± 1    | 1          | 0    |
| C57BL              | 2          | 0     | 2          | 3 ± 1    | 2          | 0    |

**Table VI.**—Tumour Cells not Lysed *in vitro* by NK Cells

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**Fig. 2.**—Comparison of PVP-PEC (□——□) nucleated spleen cells (▲——▲) and PPB-PEC (●——●) from BALB/c.nu mice as effectors of natural cytotoxicity.
less than the control, even when the PPB-PPEC were present at their lowest CL/T ratio (1/1).

These results suggest that there is a subpopulation of highly active NK cells in various tissues, whose activity can be suppressed by another cell type, perhaps macrophage in nature. Thus, the low activity of marrow, a rich source of polymorphs, may be explained by this suppression phenomenon rather than by involving a functional immaturity of the effector cell in marrow.

The possible role of Eosinophils in the NK-cell phenomenon

Since these results suggested that polymorphonuclear leucocytes might play a part in the NK-cell phenomenon, experiments with eosinophil polymorphonuclear-leucocyte-enriched suspensions were performed. These suspensions were obtained from the blood and spleens of Ascaris suum-infected mice. There was no significant increase in the lysis of 51Cr WEHI-7 when the cell suspensions from infected and non-infected BALB/c normal mice were compared. Table VII shows the details for 1 of 4 such experiments. As can be seen from the differential cell counts the Ascaris suum-infected mice had a marked peripheral-blood eosinophilia, and a significant eosinophil count in the spleen, and yet showed no increase in cytotoxic activity. Similar results were obtained in 3 other experiments, including 2 with cells from nude mice.

**Table VII.—Comparison of Lysis of WEHI-7 Tumour Cells by CL from Spleen and Blood of Ascaris-infected and Uninfected BALB/c Mice**

| Cell source | Ascaris* infection | Eosinophils | Mean % specific lysis |
|-------------|--------------------|-------------|----------------------|
| Spleen      | 0                  | 16 ± 0      |                      |
| Blood       | 0.5                | 6 ± 0       |                      |
|             | 33                 | 4 ± 1       |                      |

* Mice infected with Ascaris suum 11 days previously.
† Differential count of Giemsa-stained smears.

**DISCUSSION**

These results confirm the findings of others, that nucleated haemopoietic cells from the spleens of various strains of non-immune inbred mice are capable of in vitro lysis of a variety of tumour cell lines. The reported higher response of nude (athymic) mice than of their normal counterparts (Kiessling et al., 1975a, b), has been confirmed and extended by examining spleen cells from 3 nude mouse
strains, BALB/c, CBA and C57BL, for cytotoxic activity on a variety of tumour lines. Some evidence for genetic factors controlling the phenomenon was obtained, but the basis of this is not clear at present, and a consistent MHC-linked effect was not observed. Further work is in progress with a wider range of H-2 and allotype congenic normal and nude mice to further define possible genetic factors controlling the activity of the NK cell. The inference that might be made from these studies at present would be that multiple specificities may be involved in NK recognition, and that at least some of these may be under genetic control, but that several different such controls are operating.

An attempt was made to identify the NK cell type by comparing the cell sources of the NK cells. The most cytotoxic cells were the polymorph-enriched peritoneal-exudate cells (PVP-PEC) from mice treated with i.p. polyvinylpyrrolidone, which were 70–90% polymorphonuclear leucocytes. Further experiments demonstrated that eosinophils were not likely to be involved, although “non-specific” in vitro cell destruction by eosinophil-enriched peripheral-blood leucocyte preparations has been reported (O'Toole, 1973). Although PVP-PEC cell suspensions are 10–30% mononuclear, the comparison of these with mononuclear-enriched peritoneal-exudate cell suspensions (PPB-PEC) harvested from mice treated i.p. with proteose peptone broth, suggested that mononuclear cells were not the effectors. Furthermore, it was shown that the PPB-PEC were actually inhibitory when mixed with PVP-PEC.

A similar type of in vitro inhibition of tumour-cell lysis by mononuclear PEC, harvested 4–5 days after i.p. injection of 1 ml of 10% thioglycollate, has been reported (Fernbach, Kirchner and Herberman, 1976). In that instance, the cytotoxic effectors were not NK cells but C57BL Tc induced in vitro to BALB/c alloantigens, and the tumour target was P815. In the PVP-PEC and PPB-PEC mixing experiment reported herein (Fig. 3B) there was a dramatic reduction in specific lysis once the mononuclear PEC exceeded 10% of the effector cells (which corresponds to PPB-PEC/target ratios of 1/1 and higher in Fig. 3B). This finding suggests that a balance might exist in the various cell sources between inhibitory and cytotoxic cells, so the final lytic activity need not directly reflect the major histological type present. Such a balance has also been suggested by Jolley, Boyle and Ormerod, (1976) who found that PEC monolayers were capable of both effecting and inhibiting in vitro antibody-dependent cell-mediated lysis of lymphoma cells. Although their monolayers were predominantly macrophages, they speculated that a non-phagocytic minor subpopulation might be the effector population and the macrophages the inhibitors.

There have been many reports of in vitro target cell destruction by polymorphs and polymorph components in both the microcytotoxicity assay (Takasugi and Klein, 1970) and in 51Cr-release assays of the type described here (Lundgren, Zukowski and Moller, 1968; Takasugi et al., 1972, 1975; Edelson and Cohn, 1973; Clark and Klebanoff, 1975). However, in all these reports, target cell destruction was non-specific, and it is only recently that studies on antibody-dependent cell-mediated cytotoxicity (ADCC) have indicated that the polymorphonuclear leucocyte is, like the lymphocyte (Lamon et al., 1975) and the macrophage (Zembala, Ptak and Hanczakowska, 1973) capable of specific tumour cell cytotoxicity (Gale and Zigelboim, 1974). This appears to be mediated via a highly cytophilic anti-tumour antibody and the Fc receptor of the polymorph (Gale and Zigelboim, 1975). Thus, this more recent work provides evidence for a mechanism of specific polymorph mediated tumour-cell killing. As was reviewed earlier, there are studies which suggested that the NK cell does not lyse by an ADCC mechanism, although the possible presence of the Fc receptor on NK cells (Herberman et al., 1977) reopens this question. In view of the distinctive Ig-
class requirements for Fe-receptor binding by different cell types (Warner, 1974; Dickler, 1976) it is possible that a highly cytoplasmic antibody of a particular subclass bound to a specific cytotoxic hematopoietic cell subpopulation might be involved in determining the specificity of NK killing. This natural antibody in turn may represent a natural anti-lymphotropic virus antibody, and further studies on both aspects of cytophilic antibody and possible viral specificity are required. Against this view, however, is our observation that anti-light-chain pretreatment of the NK cell population prior to cytotoxic assay does not suppress the cytotoxic activity.

However, since PVP-PEC, although enriched for polymorphs, are a very heterogeneous source of NK cells, the results reported herein do not provide direct evidence that the NK cell is a polymorph. It may well be that the 20–30% of non-polymorphonuclear leucocytes in the suspension contain an enriched NK-cell subpopulation. These findings do, however, suggest that the whole range of nucleated cell types present in particular suspensions which show NK-cell activity in vitro should be considered in the search for the identity of the effector cell.

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