Anti-tumour efficacy of mouse spleen cells separated with *Dolichos biflorus* lectin (DBA) in experimental pulmonary metastases of B16 melanoma cells

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Summary Anti-tumour effector cells were generated through 4 days culture of normal C57BL/6 splenocytes in a medium containing concanavalin A supernatant and then fractionated with *Dolichos biflorus* lectin (DBA) into DBA+ (agglutinable with DBA) and DBA− (non-agglutinable with DBA) cells. The DBA+ cells, fused intravenously into mice together with B16 melanoma cells, or adoptively transferred into mice 3 days after the injection of B16 cells, caused a marked decrease in the number of lung nodules, while the DBA− cells exerted no effect. On the other hand, the DBA+ cells exhibited higher cytolytic activity in *vitro* than the DBA− cells in short-term 3Cr-release assays. Then, we analysed the mechanism of the strong anti-tumour activity of DBA+ cells in *vivo*. We found that DBA+ cells showed higher response to recombinant interleukin-2 (rIL-2) than DBA− cells and proliferated very well with a small amount of IL-2. In addition, DBA+ cells adhered more strongly to lung endothelial cells than DBA− cells in response to rIL-1 or rTNF. Furthermore, DBA− cells produced larger amounts of macrophage activating factor (MAF) including IFN-γ when cultured with B16 melanoma. Taken together, our results show that DBA+ cells are effective in reducing experimental pulmonary metastases not only by the direct lytic activity but also by the indirect killing activity through the activated macrophage.

In previous studies, we fractionated splenocytes of X5563 tumour-bearing mice, which had been treated with the supernatant of mouse splenocytes cultured with concanavalin A (Con A sup), with *Dolichos biflorus* lectin (DBA) into DBA+ (agglutinable with DBA) and DBA− (non-agglutinable with DBA) cells and demonstrated that DBA+ cells showed strong antitumour activity against X5563 in *vivo* when injected into mice together with human recombinant interleukin-2 (rIL-2), despite the fact that DBA− cells showed stronger cytolytic activity in *vitro* than DBA− cells (Okada et al., 1986).

Recent studies have shown that murine splenocytes or human peripheral blood lymphocytes, on incubation in *vitro* in the presence of IL-2, acquire the ability to lyse a variety of fresh syngeneic murine and autologous human tumour cells in short-term 3Cr-release assays. These lymphokine-activated killer (LAK) cells are distinct from natural killer (NK) cells or classical cytotoxic T-cells, and are lytic toward allogenic tumour target cells from murine and human sources as well (Grimm et al., 1982, 1983; Grimm & Rosenberg, 1983; Rosenstein et al., 1984).

The adoptive transfer of syngeneic LAK cells in conjunction with the systemic administration of rIL-2 reduced the number of pulmonary and hepatic metastases from murine tumours of different histological types (Lafreniere & Rosenberg, 1985a, b; Mule et al., 1984, 1985, 1986a; Moshe et al., 1986).

In the present study, we examined the ability of Con A sup-cultured cells, separated with the use of DBA, to reduce the number of experimental pulmonary metastases in a murine model. Then we investigated the mechanism of anti-tumour activity of DBA− cells in *vivo*, in terms of the following characteristics of the effector cells: (1) the proliferation rate in response to rIL-2; (2) the lymphokine producing activity; (3) the binding activity to lung endothelial cells under inflammatory conditions.

Materials and methods

Mice
Female C57BL/6, BALB/c and DBA/2 mice (7–12 weeks old) were used in these experiments. The mice were obtained from Charles River Japan Inc. (Kanagawa, Japan).

Tumours
The B16 melanoma tumour line used was of C57BL/6 origin. B16 tumour cells (1 × 10⁶) were injected into the tail veins of C57BL/6 mice. After 21–23 days, the animals were killed, their lungs harvested and then the pulmonary nodules counted. Complete enumeration of the experimental metastases was possible because they comprised black nodules on the surface of the lungs. This tumour line was maintained in RPMI-1640 medium supplemented with 10% heat-inactivated fetal calf serum (FCS; MA Bioproducts, Walkersville, MD, USA), 20 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid (HEPES), 2 mM glutamine (Wako Pure Chemical Co., Tokyo, Japan), penicillin (100 U ml⁻¹; Sigma Chemical Co., St Louis, MO, USA) and streptomycin (100 μg ml⁻¹; Sigma Chemical Co.). Cultured cells were harvested by treatment with Ca²⁺/Mg²⁺-free Hank's balanced salt solution containing 0.05% trypsin and 0.02% disodium EDTA at 37°C for 5 min. RPMI-1640 medium containing 10% FCS was added to stop the reaction and then the cells were washed twice with RPMI-1640 medium. Cells were resuspended in the appropriate buffer at the appropriate concentration and used for inoculation.

Con A sup
Spleen cells from female BALB/c mice (8–12 weeks old) were treated with modified Gey's BSS, to lyse erythrocytes, and then cultured at 37°C under 5% CO₂ in air for 22–24 h in RPMI-1640 medium supplemented with 5 × 10⁻⁵ M 2-mercaptoethanol (Sigma), 4 mM glutamine, 1 mM sodium pyruvate, 20 mM HEPES (pH 7.2), 1% non-essential amino acids (GIBCO), penicillin (100 U ml⁻¹) and streptomycin.

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(100 μg ml−1) (complete medium) containing Con A (2.5 μg ml−1) and 2.5% FCS. Then the supernatant was filtered through a membrane filter (pore size, 0.2 μm; Toyo Roshi, Tokyo, Japan), and stored at −20°C until use. The supernatant (Con A sup) thus prepared was generally used for the cell cultures.

Assay for IL-2 activity

The IL-2 activity present in the Con A sup prepared by the method described above was determined in a standard microassay method based upon the IL-2 dependent proliferation of CTLL-2 cells (Gillis et al., 1978), which were cultured in flat-bottomed microplate wells in complete medium (200 μl). Each well contained 10⁴ CTLL cells together with log 2 dilution (ranging from 3% to 50% by volume) of the Con A sup. As a positive control, rIL-2 (Takeda Chemical Industries Ltd, Osaka, Japan) was added to the cultures. The activity of the rIL-2 preparation was defined as 1 U per 27 ng by the supplier. After a 24 h incubation (37°C in a humidified atmosphere of 5% CO₂ in air) the microplate wells were pulsed with 0.5 μCi of ³H-TdR (15 Ci mmol⁻¹; New England Nuclear, Boston, MA, USA). Cultures were harvested 6 h later onto glass fibre filter strips and ³H-TdR incorporation was determined with a liquid scintillation counter (LCS-700, Aloka, Tokyo, Japan).

Generation of cytotoxic effector cells

The spleens were removed from female C57BL/6 mice and then teased into a single cell suspension in RPMI-1640 medium. The cells were then centrifuged and the pellet resuspended in modified Gey’s BSS for 3–4 min at room temperature, to lyse erythrocytes. The cells were then washed three times and cultured at 37°C under 5% CO₂ in air in 24-well plates (no. 3424; Costar, Cambridge, MA, USA) at a concentration of 2 × 10⁶ cells ml⁻¹ with 10% Con A sup. On day 3, the culture supernatant was removed, and the cells were harvested, washed and then recultured in the same medium containing 10% Con A sup. Two days later, the cells were harvested and separated with the DBA lectin (EY Laboratory, San Mateo, CA, USA) before i.v. injection or in vitro cytotoxicity assays.

Separation of cytotoxic effector cells through the use of the DBA

The Con A-sup-cultured cytotoxic effector cells were separated through rosetting with DBA-coupled sheep red blood cells (SRBC) followed by Ficoll-Utografain density gradient centrifugation as previously described (Okada et al., 1986). The DBA⁺ (rosette-forming cells with DBA-SRBC) and DBA⁻ (non-rosette-forming cells with DBA-SRBC) cells were treated with modified Gey’s BSS, to lyse SRBC, and then washed with a solution of 50 mM N-acetyl-D-galactosamine, a haptenic sugar for the DBA lectin.

³⁵Cr-release of cytotoxicity assay

The 4⁰ chromium release assay was used in this study. Varying numbers of effector cells (100 μl) and 10⁶ viable ³⁵Cr-labelled tumour target cells were plated in the wells of 96-well round bottomed microplates (Sumitomo, Osaka, Japan) in a total volume of 200 μl. The plates were centrifuged at 100 g for 1 min and then incubated at 37°C under 5% CO₂ in air for 4 h. Then the plates were centrifuged again at 250 g for 5 min, and the supernatants (100 μl) were harvested and subjected to radioactivity counting with a gamma-counter (ARC-500 auto well gamma system; Aloka, Tokyo, Japan). The spontaneous release was determined by incubating ³⁵Cr-labelled target cells alone, and the maximum release was determined after lysing the target cells in 0.4 N NaOH. The percentage of specific ³⁵Cr release was calculated according to the following formula:

\[
\% \text{ of specific } ^{35}\text{Cr release} = \frac{\text{experimental release (c.p.m.)} - \text{spontaneous release (c.p.m.)}}{\text{maximum release (c.p.m.)} - \text{spontaneous release (c.p.m.)}} \times 100
\]

The spontaneous release was less than 10% B16 of the maximum release in all experiments.

Adaptive immunotherapy models

C57BL/6 mice were given i.v. injections of 1 × 10⁶ melanoma cells in 200 μl of RPMI-1640 medium. Cytotoxic effector cells were suspended at 4–5 × 10⁶ cells in 200 μl of RPMI-1640 medium and then injected into the tail vein together with the tumour or 3 days after the tumour cell injection. On days 21–23 after the tumour cell injection, the mice were killed for enumeration of metastatic pulmonary nodules. The nodules were counted in a blind fashion, i.e. without knowledge of the treatment. Lungs with metastases too numerous to count upon autopsy were assigned an arbitrary value of 280, since we were able to count reliably only up to about 280 metastases per lung.

Phenotyping of cytotoxic effector cells

DBA-separated or unseparated cells (1 × 10⁶) were preincubated on ice for 1 h (first incubation) with 200 μl of monoclonal biotinyl anti-Lyt1.2 (final dilution, 1/50) or monoclonal biotinyl anti-Lyt2.2 (final dilution, 1/50) (Becton Dickinson, Mountainview, CA, USA) in PBS containing 0.1% Na₃N₃ and 0.1% bovine serum albumin (PBS-Na₃N₃-BSA), and then they were washed twice and treated with 200 μl of fluorescein isothiocyanate (FITC)-conjugated avidin (final dilution, 1/100) (Vector Laboratories, Burlingame, CA, USA) on ice for 30 min (second incubation). When staining cells for analysis of L3T4 antigens, the first incubation was performed with monoclonal anti-L3T4 (final dilution, 1/10; Becton) and the second with FITC-conjugated anti-rat kappa (final dilution, 1/50; Becton).

When staining cells for determination of the affinity of the DBA lectin, the first incubation was performed with biotinyl DBA (final dilution, 15 μg ml⁻¹; Vector) for 30 min, and the second with FITC-conjugated avidin (final dilution 1/100; Vector Lab.).

In the case of staining cells for analysis of asialo GM₁ (AGM), the first incubation was performed with anti-AGM (final dilution, 1/250; Wako Chemical Co., Osaka, Japan) and the second with FITC-goat anti-rabbit Ig (IgA + IgG + IgM) (final dilution, 1/100; EY Laboratory, San Mateo, CA, USA) for the second. The expression of NK1.1 on effector cells was examined by staining them with the supernatant of anti-NK1.1-producing PK136 hybridoma (Koo & Peppard, 1984) (final dilution, 1/100; Cappel, West Chester, PA, USA).

We examined the expression of T cell receptor on effector cells by staining them with an anti-T₁ monoclonal antibody 145–2C11 (Leo et al., 1987) (final dilution, 1/20) for the first incubation, and with FITC-goat anti-hamster IgG (final dilution, 1/100; EY Laboratory, San Mateo, CA, USA) for the second. The expression of NK1.1 on effector cells was examined by staining them with the supernatant of anti-NK1.1-producing PK136 hybridoma (Koo & Peppard, 1984) (final dilution, 1/100) for the first incubation, and with FITC goat anti-mouse IgG (Fc fragment specific, final dilution, 1/50; Cappel) for the second.

Staining of cells for analysis for Thy1.2 was performed by first incubating the cells on ice for 1 h with FITC-conjugated anti-mouse Thy1.2 (final dilution, 1/100; Bio-Yeda, Rehovot, Israel), and then each lot of cells was analysed using a cell sorter (FCS-1; Japan Spectroscopic Co. Ltd, Tokyo, Japan) after washing three times with PBS-Na₃N₃-BSA.

Binding to endothelial cells (EC)

EC monolayer adhesion assay (Damle et al., 1987; Cavender et al., 1987a, b) was performed as follows. Briefly, to a monolayer of rat lung endothelial cells established by Nakajima et al. (1987) in a 96-well flat bottomed plate was added various doses of rIL-1 or rTNFα (200 μl per well). After 4 h incubation (37°C, 5% CO₂ in air), the plate was washed
twice, and 51Cr-labelled DBA-separated cells (4 x 10⁷ per 200 μl per well) were added. After 1 h incubation (37°C, 5% CO₂) the plate was washed three times with a warm medium to remove non-adherent cells, and 1% Triton X-100 (200 μl per well) was added. Then the plate was incubated for 10 min at room temperature. The supernatant (100 μl per well) was harvested and assayed for radioactivity with a gamma-counter (ARC-500 auto well gamma system; Aloka, Tokyo, Japan). The percentage of the effector cells which bound to EC was calculated with the following formula:

\[
\% \text{T-EC binding} = \frac{\text{c.p.m. 0.1 ml lyse}}{\text{c.p.m. in original lymphocyte suspension}} \times 100
\]

**Macrophage activating factor (MAF) sample**

DBA-separated cells (2 x 10⁶ ml⁻¹) were cultured with the B16 melanoma (1 x 10⁶ ml⁻¹) in a 24-well plate (Costar no. 3424, Cambridge, MA, USA) in complete medium including rIL-2 (0.2 U ml⁻¹). The supernatant was harvested after 3 days, and stored at −20°C until assaying for MAF activity.

**Assay for macrophage activating factor activity for cytotoxicity (MAF-C activity)**

MAF-C activity was assayed essentially by the method of Higuchi et al. (1987). Briefly, peritoneal exudate cells (PEC) were obtained from DBA/2 mice 3 days after intraperitoneal injection of proteose peptone. These PEC (1 x 10⁷ per 100 μl per well) were plated in a 96-well microplate (Sumitomo, Osaka, Japan) in RPMI-1640 medium including 10% FCS. After 24 h incubation (37°C, 5% CO₂ in air), the supernatant was removed, and the MAF sample was added. When the effect of anti-IFN-γ was examined, anti-IFN-γ (1/100 rabbit anti-mouse IFN-γ having activity of neutralising 10⁶ U IFN ml⁻¹; Toray Industries Inc., Tokyo, Japan) was added to the sample. The plate was then incubated for 24 h (37°C, 5% CO₂ in air) before the addition of 1H-TdR prelabelled P815 cells (10⁵ cells per 200 μl per well). After 2 days incubation (37°C, 5% CO₂ in air), the supernatant (100 μl per well) was harvested and 1H-TdR release was determined with a scintillation counter. Control release was determined by incubating 1H-TdR-labelleed target cells alone, and the total scintillation was determined with target cells dissolved with 1% sodium dodecyl sulphate. The percentage of MAF-C activity was determined with the following formula:

\[
\text{MAF-C activity} (%) = \frac{\text{release in test sample} - \text{release in control sample}}{\text{total count} - \text{release in control sample}} \times 100
\]

**Results**

**In vitro cytolytic activity of DBA-separated effector cells**

Since we previously demonstrated that allogeneic CTL (Yamazaki et al., 1983) and Con A sup-cultured splenocytes from X5563 tumour-bearing C3H/HeN mice (Okada et al., 1986) could be enriched in the cell fraction having affinity for DBA (DBA⁺) cells, when examined in vitro against X5563 cells in a 4 h 51Cr release assay, we investigated whether or not Con A sup-cultured splenocytes from normal mice were enriched in the DBA⁺ fraction in the present study.

To induce cytotoxic effector cells, we used normal C57BL/6 mouse spleen cells expanded in vitro in a Con A sup-containing medium. The IL-2 activity present in 10% (v/v) Con A was about 0.2 U ml⁻¹ when determined in a standard microassay using the IL-2 dependent CTL line, CTLL-2. Splenocytes were cultured for 4 days in the presence of 10% Con A sup (v/v) and then separated into DBA⁺ and DBA⁻ fractions by means of rosette formation using DBA coupled SRBC. The distribution into the DBA⁺ and DBA⁻ fractions was 29.6 ± 2.4% and 70.4 ± 2.4%, respectively (mean ± s.e. for seven experiments). To confirm that DBA⁺ cells were highly enriched in the DBA⁺ fraction, separated and unseparated cells were incubated with the biotinyl-DBA followed by incubation with FITC-avidin, and then the stained cells (10⁵ cells per sample) were analysed by flow cytometry. As shown in Table 1, DBA⁺ and DBA⁻ cells were highly enriched in the DBA⁺ and DBA⁻ fractions, respectively.

The cytolytic activity of these cells was determined by the standard 4-h 51Cr-release assay. As shown in Figure 1a, the DBA⁺ cells showed stronger cytolytic activity against the B16 melanoma than the DBA⁻ cells in vitro (E/T = 25; Student’s t test, P < 0.01). We also assessed the cytolytic activity of Con A sup-cultured splenocytes from B16 tumour-bearing C57BL/6 mice against B16 melanoma cells in vitro. Viable B16 tumour cells (1 x 10⁶ per mouse) were inoculated subcutaneously into the backs of C57BL/6 mice. Seven days later, splenocytes were prepared from these mice and cultured for 4 days in the same manner. They were separated with the DBA and then separated cells were tested for lytic activity against B16 cells. As shown in Figure 1b the DBA⁺ cells also exhibited stronger cytolytic activity than the DBA⁻ cells in this case (E/T = 25; Student’s t test, P < 0.01).

We then examined in vitro cytolytic activity of Con A sup-cultured splenocytes from normal mice against NK sensitive target cells, YAC-1 and NK resistant target cells, P815 and EL-4. As shown in Figure 2, the DBA⁺ cells also showed significantly stronger cytolytic activity than the DBA⁻ cells in these cases (E/T = 15; Student’s t test, P < 0.001).

**In vivo anti-tumour activity of DBA-separated cells**

To evaluate the anti-tumour efficacy of the DBA-separated cells in vivo, DBA-separated cells (4 x 10⁷) were intravenously injected into C57BL/6 mice together with B16 cells (1 x 10⁵) injected (Winn assay). Figure 3a shows the number of experimental pulmonary metastases determined for each group in a blind fashion. The infusion of DBA⁻ cells caused a marked decrease in the number of lung metastases detectable on day 21, compared with that in mice given DBA⁺ cells (Student’s t test, P < 0.01), or unseparated cells (Student’s t test, P < 0.01) or without treatment (Student’s t test, P < 0.001). Since DBA⁻ cells showed significantly stronger anti-tumour effect than DBA⁺ cells in the Winn assays, then we investigated the effect of adoptively transferred DBA-separated cells on B16 tumour nodules in lungs that had already undergone micro-metastasis. B16 cells (1 x 10⁵) were intravenously injected into mice, and 3 days later, DBA-separated cells (5 x 10⁴) were i.v. transferred. On day 23 after the tumour cells injection, the number of experimental pulmonary metastases was determined. Figure 3b shows that the adoptive immunotherapy for established 3-day pulmonary B16 metastases with DBA⁻ cells resulted in a significant reduction in the number of pulmonary metastatic nodules compared with the transfer of unseparated or DBA⁺ cells (Student’s t test, P < 0.001). Entirely similar results to those in Figure 3b were obtained in three independent experiments.

It was demonstrated by Mazumder and Rosenberg (1984) that injection of LAK cells (1 x 10⁹) could significantly decrease the number of experimental pulmonary B16 melanoma metastases in mice injected with 2 x 10⁵ B16 cells.

**Table 1 Surface phenotypes of DBA-separated cells**

| Cells                  | DBA Thy1.2 2C11 L3T4 Lyt1.2 Lyst2.2 AGM NK1.1 |
|------------------------|-----------------------------------------------|
| DBA⁺                   | 90.4 98.3 63.6 40.4 41.5 92.6 79.1 9.1       |
| DBA⁻                   | 11.5 96.6 72.5 66.0 36.3 86.3 90.4 12.2       |
| Unseparated            | 27.2 96.1 67.8 n.t. 40.6 91.7 n.t. 10.8       |

Values are % positive cells. *DBA-separated cells were analysed for the expression of surface markers by the flow cytometry. n.t., not tested.
Figure 1 Cytolytic activity of DBA-separated cells against B16 melanoma cells. Spleen cells from normal C57BL/6 mice were separated with the DBA lectin after culturing for 4 days in the presence of 10% (v/v) Con A sup. a. The cytolytic activity against B16 cells of DBA⁺ (○), DBA⁻ (●) and unseparated (□) cells was examined by means of the 4 h ⁵¹Cr-release assay. Points, means for triplicate experiments; bars, s.e. b. Cytolytic activity of DBA-separated spleen cells from B16-bearing mice cultured with Con A sup. B16 cells (10⁶) were subcutaneously inoculated into the backs of C57BL/6 mice, and 7 days later, the spleen cells were cultured for 4 days in the presence of 10% (v/v) Con A sup. These cells were separated into DBA⁺ (○), DBA⁻ (●) and unseparated (□) cells, and then examined by means of the 4 h ⁵¹Cr-release assay. Points, means for triplicate experiments; bars, s.e.

Figure 2 Cytolytic activity of DBA-separated cells against B16 melanoma (a), YAC-1 (b), P815 cells (c) and EL-4 cells (d). The effector cells are the same as Figure 1a. The cytolytic activity against each target of DBA⁺ (○), DBA⁻ (●) and unseparated (□) cells was examined by the 4 h ⁵¹Cr-release assay. Points, means for triplicate experiments; bars, s.e.

3 days before the injection of LAK cells, but in the present study, a much lower number of DBA⁻ effector cells was found to be sufficient to decrease the number of lung metastases. On the other hand, DBA⁺ or unseparated cells were ineffective on adoptive transfer under our experimental conditions.

We obtained the same results for another mouse experimental pulmonary metastasis model; DBA-separated BALB/c effector cells against NL-17 cells (M. Takano, T. Okada and T. Osawa, unpublished results) which are a highly metastatic cell line established from colon adenocarcinoma 26 cells (Tsuruo et al., 1983).

Phenotypes of DBA-separated cells

We then determined the surface phenotypes of the DBA-separated cells. As shown in Table I, most of the DBA⁺, DBA⁻ and unseparated cells were highly Thy1.2⁺. Although both separated populations (DBA⁺ and DBA⁻) were found to express the L3T4, Lyt1.2 and Lyt2.2 antigens (Table I and Figure 4), the DBA⁺ fraction contained more L3T4 positive cells than the DBA⁻ fraction, and the DBA⁻ fraction consisted of the cells which are a little more positive in Lyt1 and Lyt2 antigens than the DBA⁻ fraction. Both DBA⁺ and DBA⁻ cells were AGM1⁺, and these cell populations contained NK1.1⁺ cells. As for the T cell receptor, both DBA⁺ and DBA⁻ cells were 2C11⁺.

Growth rate of DBA-separated cells in the presence of IL-2

To examine the reason why DBA⁻ cells, despite their low cytolytic activity in vitro, are markedly effective in vivo, we
Binding to endothelial cells of blood vessel under inflammatory condition

Infiltration of leukocytes into inflammation sites, such as tumour metastasis sites, is related to their ability to bind to endothelial cells of blood vessels under inflammatory condition. The binding is well known to be regulated by cytokines such as IL-1 and IFN (Cavender et al., 1987a,b). In vitro study showed that the pretreatment of endothelial cells with rIL-1 or rTNF augments their adhesion to lymphocytes or IL-2 cultured lymphocytes (Damle et al., 1987; Cavender et al., 1987a,b), and the lymphocyte-binding reached a plateau at the concentration of 1 U ml⁻¹ of IL-1 (Pober et al., 1986) or TNF (Cavender et al., 1987b). We tested the rate of the adhesion of the DBA-separated cells to the endothelial cells of blood vessels in rat lung after treatment of the endothelial cells with various doses (0–5 U ml⁻¹) of rIL-1 or rTNF. Since mouse lung endothelial cells were not available, rat lung endothelial cells kindly donated by Dr. M. Nakajima (MD Anderson Hospital, Houston, TX, USA) were used in this study. As shown in Figure 6, rIL-1 or rTNF increased the adhesion of DBA⁻ cells to endothelial cells while these lymphokines had no effect on the adhesion of DBA⁺ cells to endothelial cells. At the site of the tumour metastases, the inflammatory lymphokines such as IL-1 or TNF may be secreted. These results indicate that DBA⁻ cells have an ability to attach to endothelial cells under the influence of IL-1 or TNF, and they can infiltrate advantageously into the tumour metastasis sites. However, similar studies on mouse lung endothelial cells are still necessary, because there is a possibility that cell surface adhesion molecules are different between mouse and rat endothelial cells.

Indirect anti-tumour effect through the elaboration of lymphokines

Furthermore, since there is a possibility that DBA⁻ cells act indirectly through activation of other types of cells by secretion of certain lymphokines, experiments were performed to find out whether DBA-separated cells secrete the macrophage activating factor (MAF). DBA-separated or unseparated cells were cultured with the B16 melanoma in the presence of rIL-2 for several days. Then the supernatants were tested for MAF activity. MAF activity was estimated by the induction of cytolytic activity of proteose peptone-induced macrophages (Mφ) against P815 cells. The culture medium used in these experiments contained polymyxin B to block the effect of contaminating LPS, if any. As shown in Figure 7, DBA⁻ cells produced a large amount of MAF when cultured with the B16 melanoma in the presence of rIL-2 for 3 days. On the other hand, DBA⁺ or unseparated cells produced only a detectable level of MAF. In the absence of the B16 melanoma, neither DBA⁺ nor DBA⁻ cells released MAF, and no MAF was produced from B16 cells alone under these culture conditions. If rIL-2 was not added to the culture medium, all the effector cells died in 2 days and no MAF was secreted. These results suggested that DBA⁻ cells were stimulated by the B16 melanoma in the presence of rIL-2 and secreted MAF in the culture medium. The production of MAF was detectable on day 2 and reached a peak on day 3. After 4 or 5 days, the viability of effect cell decreased and no MAF activity was detected.

We then examined characteristics of the MAF activity in DBA⁻ supernatant. As shown in Figure 8, neither LPS nor murine rIFN-γ showed MAF activity by themselves, but showed MAF activity synergistically. This effect was blocked by anti-IFN-γ antibody. When anti-IFN-γ was added to DBA⁻ sup, MAF activity was almost completely diminished. These results show that MAF activity in DBA⁻ cells is mainly mediated by IFN-γ, and suggest the possibility that DBA⁻ cells secrete IFN-γ in vivo and the activated Mφ thus induced may also contribute to the anti-tumour activity. However, direct determinations of IFN-γ in the supernatants of DBA-separated and unseparated cell populations are necessary to verify this assumption. Furthermore, cytotoxic activity of
alveolar macrophages isolated from DBA- cells treated mice should be examined to confirm the participation of alveolar macrophages in the suppression of experimental pulmonary metastasis of tumour cells in DBA- cells treated mice. These are the next targets of our research.

Discussion

In this study, the DBA- cells separated from Con A-supercultured mouse splenocytes were found to show high inhibitory activity toward experimental metastasis when injected together with B16 tumour cells (Figure 3a) and also when adoptively transferred 3 days later after inoculation of the tumour cells (Figure 3b). Furthermore, the cell number that was sufficient for reducing the number of metastatic lung nodules was markedly less than that employed for the therapy involving LAK cells by other investigators. On the other hand, DBA+ cells showed stronger cytolytic activity in vitro than DBA- cells (Figure 1), but they showed much weaker activity in vivo (Figure 3).

The dissociation of the cells responsible for the in vitro cytolytic effect and the in vivo anti-tumour effect, respectively, was also demonstrated by other investigators. Mule et al. (1986b) reported that, in combination with the systemic administration of rIL-2, LAK cells which had been treated with anti-Thy1 and complement, and which were not cytolytic in vitro assays, remained as effective as untreated cytolytic LAK cells upon adoptive transfer in reducing the number of established pulmonary metastases. Shu et al. (1987) also showed that although spleen cells, which had been sensitised in vitro with viable tumour cells and IL-2, from normal C57BL/6 mice did not exhibit anti-tumour activity in vivo, they displayed non-specific LAK-like cytotoxic activity in vitro. They therefore concluded that it is unlikely that the in vitro cytotoxic activity was directly related to the in vivo anti-tumour effect.

We investigated some properties of DBA-separated cells in order to examine the reasons why DBA- cells, despite their low cytolytic activity in vitro, are much more effective than DBA+ cells in vivo. For the proliferation response to IL-2, DBA- cells were demonstrated to be much more responsive to rIL-2 than DBA+ cells (Figure 5). Many investigators
tried to attack tumour cells with LAK or CTL in combination with the injection of rIL-2 (Chou & Shu, 1987; Chou et al., 1988). Since IL-2 is liable to be metabolised and excreted quickly in vivo, administration of a relatively large amount of IL-2 is necessary to ensure the in vivo effect of the factor and this may result in serious side-effects. In this context, the fact that DBA- cells can proliferate with less IL-2 than DBA+ cells may also explain the greater anti-tumour efficacy of DBA- cells in vivo, because only a little amount of endogenous IL-2 is actually available in vivo.

At the site of the inflammation, such as the tissue where tumour cells are growing, the inflammatory lymphokines are released. So we examined the adhesion rate of DBA-separated cells to rat lung endothelial cells (EC) in the presence of the inflammatory lymphokines such as IL-1 and TNF. The binding of DBA- cells to rat lung EC was augmented by the pretreatment of EC with rIL-1 or rTNF in dose-dependent manner (Figure 6). However, the binding of DBA- cells to EC was not enhanced by this treatment. These results suggest that DBA- cells effectively bind to the lung EC in situ of tumour tissue where the inflammatory lymphokines are released, then infiltrate into the tumour site and attack the tumour cells.

Since the anti-tumour activity of activated Mφ in vivo is well known (Higuchi et al., 1988), we tested the DBA-separated cells for their ability to generate Mφ activating factor (MAF). It was found that DBA- cells secreted Mφ activating factor when cultured with B16 cells (Figure 7), and this MAF activity mainly depended upon IFN-γ (Figure 8). Although DBA+ cells could also lyse B16 cells and proliferated in the presence of IL-2 in vitro, the supernatant of DBA+ cells showed only a detectable level of MAF-activity. These results suggest that DBA- cells can induce activated Mφ in vivo and in addition to the direct cytotoxicity of DBA- cells, the activated Mφ thus induced may contribute greatly to the anti-tumour activity of DBA- cells in vivo. Considering the fact that the unseparated cells include about 70% of DBA- cells, but the supernatant of the unseparated cells shows very low level of the Mφ activating activity, there is a possibility that DBA- cells produce an inhibitor of Mφ activation.

There is another possibility that DBA- cells can more
Figure 8 Effect of anti-IFN-γ on MAF activity of DBA− sup. Murine rIFN-γ (20 U ml−1), LPS (0.1 µg ml−1) was added to the assay medium of MAF activity. Anti-IFN-γ (having activity of neutralising 10 U IFN) was added to DBA− sup (3 days culture supernatant of DBA− cells with B16 melanoma in the medium containing 0.2 U ml−1 rIL-2) and the MAF activity was assayed and % cytolysis was calculated by the same method as in Figure 7.

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