Effective removal of SCLC cells from human bone marrow. Use of four monoclonal antibodies and immunomagnetic beads

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Summary
High dose chemotherapy with autologous bone marrow transplantation (ABMT) has shown promise in several types of cancer. There is, however, a risk of transfusing contaminating tumour cells with the bone marrow cells, e.g. in patients with small cell lung carcinoma (SCLC). To eliminate SCLC cells from normal human bone marrow, four monoclonal antibodies reactive with SCLC cells were used with immunomagnetic beads in model experiments. With two cycles of immunomagnetic elimination the individual antibodies removed 2.5–4.4 log of H-146 tumour cells from a single cell suspension, as assessed in a highly reproducible soft agar assay. Different combinations of two antibodies were only marginally more effective than the individual MABs, whereas 5–6 log removal was obtained with a combination of all four antibodies. The method was equally effective when the tumour cells were mixed with bone marrow cells at a ratio of 1:10. The immunomagnetic procedure did not significantly affect the survival of normal progenitor cells, assessed in CFU-GM and CFU-GEMM assays. The results indicate that the procedure safely and effectively can be used to eliminate tumour cells from the bone marrow in conjunction with ABMT in patients with SCLC.

Most patients with small cell lung cancer (SCLC) have disseminated disease at time of diagnosis. Although SCLC is initially very sensitive to chemotherapy, the responses are of short duration and relapse of the disease occurs rapidly in 90–95% of the cases. For limited disease the 5-year survival rate is only about 5–10% (Albain et al., 1991), and the 2-year survival rate for extensive disease is less than 6% (Albain et al., 1991).

To improve the situation, autologous bone marrow transplantation (ABMT) combined with high dose chemotherapy has been tried (Humblet et al., 1987; Symann et al., 1989; Souhami et al., 1989; Williams et al., 1989; Marangolo et al., 1989; Nomura et al., 1990; Lazarus et al., 1990; Gomm et al., 1991). This treatment has increased the number of patients with complete remissions (Humblet et al., 1987; Souhami et al., 1989) and improved relapse free-survival (Humblet et al., 1987; Symann et al., 1989), but, unfortunately, the long time survival rate has not increased significantly. One contributing factor may be infiltration of tumour cells in the transfused bone marrow. It has become increasingly clear that tumour cells are often present in the marrow of SCLC patients, even if cytomorphological and histomorphological analysis remains negative (Stahel et al., 1985; Berendsen et al., 1988; Canon et al., 1988; Trillet et al., 1989; Beiske et al., 1992). In order to prevent giving tumour cells back to the patient, it is logical to purge the bone marrow for tumour cells before reinfusion. The usefulness of immunological, physical and chemical procedures have been examined for this purpose (Gordon et al., 1984; Mabry et al., 1985; Okabe et al., 1985; Benard et al., 1988; Combaret et al., 1988; Vredenburgh & Ball, 1990; Humblet et al., 1989; Meagher et al., 1989; Elias et al., 1990).

In this study the purging efficacy of immunomagnetic beads used in combination with four different monoclonal antibodies (MABs), previously selected from a panel of 17 MABs (Myklebust et al., 1991), was tested in preclinical model experiments. The optimal procedure involved the use of a mixture of all four MABs and two purging cycles, resulting in as high as 6 log of tumour cell depletion with an acceptable recovery of normal bone marrow progenitor cells.

Materials and methods

Bone marrow
Human bone marrow aspirates were obtained from healthy volunteers or from patients at the Norwegian Radium Hospital with non-SCLC malignancies that were free of tumour cells in their marrow. All samples were obtained with informed consent from the donors. Ten ml of bone marrow were layered upon Lymphoprep (Nycomed, Oslo, Norway) and the mononuclear cell fraction was obtained by centrifugation at 1200 r.p.m. for 30 min. The mononuclear cells were washed once in phosphate buffered saline (PBS) before being used in the experiments.

Cell line
The H-146 human SCLC cell line (Carney et al., 1985), kindly provided by Dr Adi F. Gazdar, National Cancer Institute, Bethesda, MD, USA, was used in the experiments. The cells were grown as suspension cultures in RPMI 1640 medium (Gibco, Paisley, UK) supplemented with 10% foetal calf serum (FCS).

Monoclonal antibodies
MOC-1 (Leij et al., 1985) and MOC-31 (Leij et al., 1986) antibodies were gifts from Dr L. de Leij, Groningen, The Netherlands. MOC-1 (IgG1) binds to the neural cell adhesion molecule (CD56/NCAM) (Moolenaar et al., 1990), and MOC-31 (IgG2a) to an epithelial antigen (cluster 2). NfLu10 (also denoted TFS-2), an IgG2b antibody recognising a 39 kDa antigen (Okabe et al., 1984), was obtained from NeoRx Corporation, Seattle, WA, USA. MLuCl1 (earlier denoted M0V15), kindly provided by Dr Sylvie Menard, Milan, Italy, is an IgG1 antibody that binds to a saccharide epitope carried by neural glycolipids, glycoproteins and mucins (Ripamonti et al., 1987).

Immunomagnetic beads
The immunomagnetic beads used, Dynabeads® M-450 SAM IgG ST (Dynal, Oslo, Norway), are uniform, magnetic, polystyrene beads with sheep anti mouse IgG (SAM) antibodies covalently bound to their surface. SAM bind all mouse IgG subclasses.

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Mechanically dispersed tumour cells (1 x 10^7), mononuclear bone marrow cells (2 x 10^6), or mixes of both (total 5 x 10^7) were incubated in plastic tubes for 30 min at 4°C with one or more MAbS (10 μg of each) in 1 ml of RPMI 1640, supplemented with 10% FCS, 100 units of penicillin and 100 μl of streptomycin (Gibco) per ml medium. The cells were washed twice in PBS with 1% FCS before adding the immunomagnetic beads suspended in 0.5 ml of medium and then incubated at 4°C for 30 min. In the experiments with the cell line the ratio of beads to tumour cells was 50:1, and in the control experiments to test the effect on bone marrow progenitor cells the ratio of beads to nucleated bone marrow cells was also 50:1. To ensure proper contact between cells and MAbS/beads, during the incubations, the tubes were rolled on a mixer (Coulter Electronics Ltd., UK). Tumour cells were removed from the suspension by placing a flat cobalt samarium magnet to the wall of the tubes. After 1 min the suspension with remaining cells was aspirated, centrifugated at 1000 r.p.m. for 5 min, and the cells resuspended in 1 ml of medium. Remaining clonogenic tumour cells and bone marrow cells in the suspension were measured by colony-forming assays as described below. In some experiments, a repeat cycle of immunomagnetic separation was performed.

Colonial-forming assays for tumour and bone marrow progenitor cells

The number of colony forming tumour cells was assessed in a soft agar assay (Courtenay & Mills, 1978), as previously described for malignant B cells (Kvalheim et al., 1987). Briefly, soft agar cultures were set up in triplicates in 10 ml tubes by adding 0.2 ml August rat blood diluted 1 to 8, 0.2 ml of appropriately diluted bone marrow/tumour cell suspensions, and 0.6 ml of 0.5% agar (Difco Laboratories, Detroit, Michigan, USA). The tubes were incubated at 37°C in 5% CO₂, 5% CO₂, and 90% N₂, and after 21 days of incubation colonies of more than 50 cells were counted using a Zeiss stereo microscope.

The clonogenic progenitor cells remaining in the bone marrow after immunobead separation were assessed in a modified version (Kvalheim et al., 1987) of the method described by Burgess et al. (1977). Mononuclear BM cells were suspended at a concentration of 2 x 10^7 cells ml⁻¹ in MACOY's 5A medium (Gibco) containing 0.3% agar, 15% FCS, 20 ng of GM-CSF (kindly provided by Schering Plough, NJ, USA) and antibiotics. Triplicate 1 ml aliquots were cultured in 35 mm plastic dishes at 37°C in 5% CO₂, 5% CO₂, and 90% N₂. After 14 days of incubation colonies of more than 40 cells were counted.

In the GEMM assay (Messner et al., 1982), 2 x 10^8 mononuclear cells were suspended in 1 ml iscove's modified Dulbecco's medium (Gibco) containing 0.8% methylcellulose, 10% FCS, 10% leucocyte conditioned medium, 30% human plasma, 5 x 10⁻⁵ M 2-mercaptoethanol and 1 unit of erythro-

### Table 1

| Individual MAb | One cycle of elimination | Two cycles of elimination | Combination of two MAb | One cycle of elimination | Two cycles of elimination |
|----------------|--------------------------|---------------------------|------------------------|--------------------------|--------------------------|
| MOC-1          | 3.7                      | 4.4                       | MOC-1                  | 3.0                      | 3.6                      |
| MOC-31         | 3.7                      | 2.5                       | NrLu10                 | 3.7                      | 3.7                      |
| NrLu10         | 3.6                      | 3.9                       | MLuCl                  | 3.6                      | 3.8                      |
| MLuCl          | 2.6                      | 4.2                       |                        |                          |                          |

H-146 cells were incubated for 30 min at 4°C with MAbS (10 μg ml⁻¹ of each) and then for 30 min with SAM-M-450 Dynabeads at a ratio beads to tumour cells of 50:1. *Calculated from the number of colonies counted, taking the plating efficiency into account, and determined as the logarithm of the number of cells depleted by the treatment. **Mean of the results obtained in 2-3 independent soft agar experiments, each performed in triplicate.

**Results**

**Cloning efficiency of SCLC cells**

When different numbers of H-146 cells were seeded out in soft agar the number of colonies formed was proportional to the number of cells plated (Figure 1). This relationship was close to linear down to ten cells plated per tube, and the plating efficiency calculated from the slope of the curve was approximately 40% in three independent experiments. Similar curves were established in each model experiment, and the curves were used to calculate the tumour cell depletion obtained by the treatment, taking the plating efficiency in each experiment into account. Efficacy of the treatment is given as the logarithm of the number of tumour cells removed.

**Tumour cell removal**

**Efficacy of immunobeads used with single and combinations of two MAbS**

Of the individual MAbS, MOC-1 and NrLu10 were the most effective in the immunomagnetic procedure, resulting in 3.7 log and 3.6 log tumour cell removal, respectively (Table I). The corresponding value for MLuCl was 2.6 log, compared to 2.1 log for MOC-31. When the procedure was performed twice, a significant increase in efficacy was obtained for MOC-1 (4.4 log depletion) and for MLuCl (4.2 log depletion), whereas the values with the other two MAbS increased only slightly (Table I).

Since it is known that tumour-associated antigens commonly are heterogeneously expressed on tumour cells, we wanted to test different combinations of two MAbS. As can be seen from Table I, the combination of MOC-1 and NrLu10 is more effective than the individual MAbS, and the combination of MOC-1 and MLuCl is more effective than the combination of MOC-1 and NrLu10.
be seen from Table I, the different combinations gave log tumour cell depletions in the range of 3.0–4.7. In most cases the combinations were not more effective than the most effective single antibody, except in some experiments involving MOC-31 and MLuCl. The results were clearly improved by performing the elimination procedure twice. The overall best results were obtained with combinations of either MOC-31 and MLuCl or NrLu10 and MOC-31, resulting in 4.6–4.7 log tumour cell depletion.

**Combinations of three and four MAbs** In the next set of experiments, combinations of three (MOC-1 + MOC-31 + NrLu10) and four (MOC-1 + MOC-31 + NrLu10 + MLuCl) antibodies were tested. As can be seen from the results presented in Table II, neither of the combinations was more effective than the most active single antibodies (Table I) when only one cycle of immunomagnetic removal was used. With two cycles, however, the tumour cell depletion with a combination of three MABs increased to 4.7 log, a comparable efficacy to that of the most effective 2-MAB combination. This value was further increased to 5.1 log when an additional incubation with the antibodies was introduced before the second cycle of immunomagnetic separation (Table II). The most effective treatment was obtained with two elimination cycles involving the combination of four antibodies. Thus, the mean log tumour cell depletion obtained in three different experiments was 5.6 log, ranging from 5.3 to at least 6 logs. In this case, the additional incubation with the MABs did not improve the results.

With the encouraging results obtained, the combinations of three and four MABs were also tested in the more clinically relevant situation with H-146 tumour cells added to fresh human bone marrow at a ratio of 1:10. In repeat experiments, a mean log tumour cell depletion of 4.8 was obtained with a mixture of three MABs (Table III). In one of the experiments, only 10 out of 1 x 10^6 tumour cells plated remained in the bone marrow after purging, as calculated from the number of colonies formed in soft agar. Even more encouraging, with the combination of four MABs no colony formation in soft agar was observed with a tumour cell/bone marrow mixture that contained 1 x 10^6 SCLC cells before purging. Since the same result was obtained in three independent experiments, it can be concluded that with the 4 MAB combination at least 6 log tumour cell depletion from the bone marrow can be achieved, a result that is even better than when the procedure was performed without the presence of bone marrow cells.

**Survival of progenitor bone marrow cells** The effect of the immunomagnetic procedure on the survival of normal bone marrow progenitor cells was assessed (Table IV). With single antibodies, the fraction of surviving CFU-GM differed with the antibodies from 80 to 49%, the lowest value was seen in experiments with MLuCl. With three and four MAB combinations, the fraction of surviving CFU-GM was 68% and 52%, respectively. The combinations were also tested in the CFU-GEMM assay, where it was found that 70–80% of the progenitor cells had survived the immunomagnetic treatment. The results indicate that the immunomagnetic purging procedure involving combinations of MABs can be used safely for purging purposes in patients.

**Discussion**

A magnetic procedure was developed for removing SCLC cells from human bone marrow. We have demonstrated that by using a combination of four antibodies and immunomagnetic sheep anti-mouse IgG beads, a 6-log tumour cell separation could be achieved, as determined by a reproducible clonogenic soft agar assay. The procedure reduced the survival of normal bone marrow progenitor cells to only a limited extent, showing that the procedure can be safely used in a clinical setting.

Heterogeneity of tumour cell surface antigen expression suggests that a cocktail of MABs may be necessary to achieve effective removal of tumour cells. This was confirmed in the present work, as the best results were obtained with a combination of all four MABs. The MABs were selected because of their ability to bind to small cell lung cancer cells, and not to normal cells in bone marrow and peripheral blood (Myklebust et al., 1991). Also, the usefulness of the same antibodies for detecting bone marrow metastases in small cell lung cancer patients has recently been demonstrated (Beiske et al., 1992). We have found that the NrLu10 and MOC-31 antibodies bind to the same antigen, but to different epitopes (unpublished data). However, they differ in their ability to

### Table II

| MAbs                  | Number of experiments | Cycle of Elimination | Log tumour cell depletion (mean) | Log tumour cell depletion (range) |
|-----------------------|-----------------------|----------------------|----------------------------------|----------------------------------|
| MOC-1 + MOC-31 + NrLu10 | 3                     | 1                    | 3.8                              | 3.4–4.1                          |
| MOC-1 + MOC-31 + NrLu10 + MLuCl | 3                     | 1                    | 4.7                              | 4.4–5.1                          |

**Number of experiments:** A, 1; B, 2; C, 3; D, 1

**Treatment and calculations as in Table I.**

### Table III

| MAbs                  | Number of experiments | Tumour cells plated (10^6) | Tumour cells remaining (mean) | Log tumour cell depletion (mean) | Log tumour cell depletion (range) |
|-----------------------|-----------------------|----------------------------|-----------------------------|----------------------------------|----------------------------------|
| MOC-1 + MOC-31 + NrLu10 | 2                     | 10^6                      | 3                           | 4.8                              | 4.5–5.0                          |
| MOC-1 + MOC-31 + NrLu10 + MLuCl | 3                     | 10^6                      | 0                           | >6.0                             | >6.0                             |

H-146 cells mixed with bone marrow cells (ratio 1:10) were incubated with the MABs (10 μg ml^-1 of each) and then with SAM-M-450 Dynabeads at a ratio tumour cells to beads of 1:50. Two purging cycles were used. Calculations as in Table I.
detect contaminating SCLC cells in fresh bone marrow aspirates (Beiske et al., 1992), and as suggested by the present results, both colonies should be included in a cocktail recommended for clinical use, together with MOC-1 and MLuCl that recognise distinct antigens (Moolenaar et al., 1990; Ripamonti et al., 1987).

To limit the need for fresh human bone marrow, the removal experiments with individual and combinations of two MABs were performed on a suspension of H-146 cells only, as previous experience has shown that such experiments closely predict results obtained when tumour cells are mixed with normal bone marrow cells. In accordance with this, the present procedure proved to be equally effective when H-146 tumour cells were mixed with bone marrow cells at a ratio of 1:10. Thus, the optimal combinations of four MABs resulted in a mean log tumour cell removal of 5.6 in the absence, and >6.0 in the presence of bone marrow cells. In experiments with MOC-31 antibody no difference in log tumour cell removal was found between 1:10 and 1:100 mixes of SCLC and marrow cells (not shown). However, limited access to human bone marrow and the high efficacy of the purging method prevented further testing at a low tumour to bone marrow cell ratio. The validity of calculating log tumour cell removal is dependent on a close to linear relationship between the number of tumour cells plated and the number of soft agar colonies formed. This requirement was met with the H-146 cell line, which showed a high and reproducible plating efficiency in the clonogenic assay.

Several reports have described methods to eliminate SCLC cells from bone marrow (Gordon et al., 1984; Mabry et al., 1985; Okabe et al., 1985; Benard et al., 1988; Vredenburgh & Ball, 1990; Humbert et al., 1989; Meagher et al., 1989; Elias et al., 1990). Vredenburgh & Ball (1990) obtained 4–5 log separation using immunomagnetic beads and three different antibodies, administered at a relatively high concentration, whereas the results of Elias et al. (1990) were less satisfactory (2.4–2.6 log). In both cases, two types of immunobeads had to be used because their tumour-associated primary antibodies were of both IgG and IgM isotypes. The results reported for other depletion procedures show varying degrees of efficacy and toxicity to normal bone marrow cells (Gordon et al., 1984; Mabry et al., 1985; Okabe et al., 1985; Benard et al., 1988; Meagher et al., 1989), the best results (4–5 log depletion) obtained by Humbert et al. (1989). The latter investigators combined immunological and pharmacological methods, and their regime was relatively toxic to bone marrow progenitor cells.

Altogether, the results in the present work were better than any of those previously reported, probably reflecting both the efficacy of immunomagnetic procedures (Kiesel et al., 1987; Kvalheim et al., 1988) and the properties of the antibodies used. In agreement with the experience of Vredenburgh & Ball (1990), the best results were obtained with two purging cycles at a 50:1 ratio of beads to tumour cells. The procedure had only minor effects on cell survival in the bone marrow progenitor cell assays. Some bone marrow specimens used for these studies were obtained from patients with other types of malignancies but without bone marrow involvement. In a few of these cases, the proliferative capacity of the progenitor cells was relatively low. This did not influence the results, except that with MLuCl used alone an unpredictable variation in the number of surviving CFU-GMs was observed in some experiments.

High dose chemotherapy followed by ABMT in SCLC patients has been used by several investigators (Humblet et al., 1987; Symann et al., 1989; Souhami et al., 1989; Williams et al., 1989; Marangolo et al., 1989; Nomura et al., 1990; Lazarus et al., 1990; Gott et al., 1991), but no improvement in survival, compared to patients treated with standard chemotherapy, has been reported. Several factors may have contributed to the reported limited benefit of ABMT, including a less than optimal patient selection with regards to stage of the disease, age, response to induction therapy, and timing and intensity of the ablative treatment. Moreover, with the development of sensitive diagnostic techniques it has become increasingly clear that a high number of SCLC patients have contaminating tumour cells in their bone marrow (Stahel et al., 1985; Berendsen et al., 1988; Canon et al., 1988; Tillet et al., 1989; Beiske et al., 1992). It should also be noted that the detection of tumour cells in bone marrow is based on studies of small volume aspirates, whereas about 11 of bone marrow is harvested for ABMT. Thus, the risk of tumour cell contamination of the transfused marrow may be higher than indicated by studying bone marrow aspirates.

One argument against ABMT in SCLC has been that several patients have relapsed locally in the lungs. It cannot be excluded, however, that these relapses are associated with i.v. transfection of contaminated bone marrow cells. The reinused cells will meet the first capillary bed in the lungs, and it is well known that interaction between tumour cells and host tissues favours tumour growth in orthotopic sites, in this case represented by the lungs. Furthermore, the effect of tumour cell trapping in the lungs is also exemplified by the finding of Glorieux et al. (1986) that a patient with neuroblastoma and another with Burkitt's lymphoma developed diffuse carcinomatosis in the lungs after ABMT.

In cancer types where patients undergoing ABMT are at risk for having contaminating tumour cells in the bone marrow, it seems logical that the marrow should be purged before transfusion. Although the value of tumour cell purging has not been definitely proven, recent evidence indicates that effective removal of lymphoma cells from the marrow has a significant effect on the clinical outcome (Gribben et al., 1991). We have previously developed highly effective immunomagnetic methods which are currently in routine clinical use in patients with B and T-cell malignancies (Kvalheim et al., 1988; Wang et al., 1992). The similar method described here for purging SCLC has been shown to be equally efficacious in tumour cell depletion from bone marrow, and the method is recommended for use in conjunction with ABMT in selected groups of SCLC patients.
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