Melphalan is widely used as a preparative agent in patients with multiple myeloma (MM) undergoing autologous stem cell transplantation (SCT). Although disease relapse is the major cause of death after a melphalan-conditioned autograft, the mechanism remains unclear. Melphalan produces a number of DNA adducts with the DNA interstrand crosslink (ICL) considered to be the critical cytotoxic lesion. By using a modification of the single-cell gel electrophoresis (Comet) assay, we have measured formation and repair of DNA ICL in plasma cells from melphalan-naive and melphalan-treated patients (ie, those who have relapsed after a melphalan-conditioned autologous SCT or oral melphalan therapy). Similar levels of dose-dependent DNA interstand crosslinking were observed in cells from both melphalan-naive and -treated patients. However, marked differences in ICL repair were observed: cells from naive patients showed no repair, whereas those from treated patients exhibited between 42% and 100% repair at 40 hours. In vitro sensitivity to melphalan in plasma cells was found to correlate with ICL repair. These findings suggest that ICL repair may be an important mechanism by which melphalan resistance emerges after autologous SCT or oral therapy. This mechanism may have implications for MM patients undergoing melphalan therapy.

(Blood. 2002;100:224-229)
tumor xenograft models were extended to tumor biopsies from patients undergoing antibody-directed enzyme prodrug therapy phase 1 clinical trials, and in vivo formation of ICL has also been followed in lymphocytes taken from patients receiving ifosfamide therapy.

In the present study we have used the Comet assay to examine formation and repair of melphalan-induced ICL in plasma cells from patients with MM following ex vivo treatment with melphalan. Two patient populations were studied: patients who had not received melphalan therapy (melphalan naive) and those treated with melphalan. The latter population includes patients who had relapsed following a high-dose melphalan-conditioned autograft and those receiving oral melphalan therapy.

Patients and methods

Patients

Twelve patients with MM were studied. Six patients were studied at diagnosis or after initial treatment with VAD and had never received melphalan. Six patients had relapsed following a melphalan-conditioned (200 mg/m²) autograft. One patient had received oral melphalan (7 mg/m²) autograft. One patient had received oral melphalan (7 mg/m²) without conditioning. Two patient populations were studied: patients who had not received melphalan therapy (melphalan naive) and those receiving oral melphalan therapy.

Drug treatment of plasma cells

Plasma cells were isolated from bone marrow aspirates with the use of standard Ficoll-Hypaque centrifugation. All samples studied contained in excess of 80% plasma cells. Formation of melphalan-induced ICL was measured at concentrations ranging from 1 to 100 μM. Plasma cells (2.5 × 10⁴ cells/mL) in RPMI 1640 medium containing 10% fetal calf serum and 2 mM glutamine (Autogen Bioclear, Calne, United Kingdom) were incubated with melphalan for 1 hour at 37°C. The drug was removed by centrifugation at 200g for 5 minutes, removing the supernatant and resuspending in drug-free media. Cells were incubated for a further 16 hours at 37°C to allow maximum formation of ICLs (t = 16 hours). Repair of ICLs was studied at 50 μM melphalan, a further 24 hours following maximum ICL formation (t = 40 hours).

Determination of DNA interstrand crosslinking

The details of the single cell gel electrophoresis (Comet) assay to measure ICLs are described in detail elsewhere. All procedures were carried out on ice and in subdued lighting. All chemicals used were obtained from Sigma Chemical (Poole, United Kingdom) unless otherwise stated. Immediately before analysis, cells were irradiated (10 Gy) to deliver a fixed number of random DNA strand breaks. After embedding cells in 1% agarose on a precoated microscope slide, the cells were lysed for 1 hour in lysis buffer (100 mM disodium EDTA, 2.5 M NaCl, 10 mM Tris-HCl, pH 10.5) containing 1% Triton X-100 added immediately before analysis and then washed every 15 minutes in distilled water for 1 hour. Slides were then incubated in alkali buffer (50 mM NaOH, 1 mM disodium EDTA, pH 12.5) for 45 minutes, followed by electrophoresis in the same buffer for 25 minutes at 18 V (0.6 V/cm), 250 mA. The slides were finally rinsed in neutralizing buffer (0.5 M Tris-HCl, pH 7.5) then saline.

After drying, the slides were stained with propidium iodide (2.5 μg/mL) for 30 minutes then rinsed in distilled water. Images were visualized with the use of a NIKON inverted microscope with high-pressure mercury light source (NIKON UK Limited, Kingston Upon Thames, United Kingdom), 510 to 560 nm excitation filter, and 590 nm barrier filter at ×20 magnification. Images were captured by using an on-line charge-coupled device (CCD) camera and analyzed by using Komet Analysis software (Kinetic Imaging, Liverpool, United Kingdom). For each duplicate slide 25 cells were analyzed. The tail moment for each image was calculated by using the Komet Analysis software as the product of the percentage DNA in the comet tail and the distance between the means of the head and tail distributions, based on the definition of Olive et al. Crosslinking was expressed as the percentage decrease in tail moment compared with irradiated controls calculated by the following formula: percentage of decrease in tail moment = [1 – (TMdi–TMcu/TMci)] × 100; where TMdi equals tail moment of drug-treated irradiated sample, TMcu equals tail moment of untreated unirradiated control, and TMci equals tail moment of untreated irradiated control.

Repair at time = 40 hours following exposure to 50 μM melphalan and subsequent irradiation was calculated as a percentage by the following formula: percentage of repair = (PD16–PDcu/PDc0) × 100; where PD16 = percentage decrease in tail moment at 16 hours and PDcu = percentage decrease in tail moment at 40 hours.

In vitro cytotoxicity

The methyl-thiazole-tetrazolium (MTT) cytotoxicity assay was carried out as previously described. Plasma cells (1.5 × 10⁶ cells/mL) were exposed to varying concentrations of melphalan in 96-well plates (200 μL/well) and incubated at 37°C, 5% CO₂ for 72 hours. Then 50 μL 3 mg/mL MTT (Sigma Chemical) was added to each well and incubated for an additional 4 hours. After centrifugation the supernatant was removed and the formazan crystals were dissolved in 200 μL dimethylsulphoxide (VWR International Limited, Poole, United Kingdom) and 25 μL Sorenson glycine buffer. The OD 570 nm was then measured and the dose giving 50% inhibition of cell growth (IC₅₀) was calculated.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age, y</th>
<th>Sex</th>
<th>Stage at diagnosis</th>
<th>Paraprotein type</th>
<th>B₉M (g/L)</th>
<th>Prior therapy</th>
<th>Autograft/no.</th>
<th>Time for relapse after autograft</th>
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<tr>
<td>1</td>
<td>46</td>
<td>M</td>
<td>IIIA</td>
<td>κ free light chain</td>
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<td>M</td>
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<td>BJκ</td>
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<td>VAD × 3</td>
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<td>3</td>
<td>66</td>
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<td>IIIA</td>
<td>IgGκ</td>
<td>2.5</td>
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<td>N/A</td>
</tr>
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<td>4</td>
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<td>IgGκ</td>
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<td>5</td>
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<td>F</td>
<td>II</td>
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<td>7</td>
<td>59</td>
<td>F</td>
<td>I</td>
<td>IgGκ</td>
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<td>IIIB</td>
<td>IgGκ</td>
<td>4.4</td>
<td>VAMP × 6</td>
<td>Yes/2</td>
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<td>10</td>
<td>55</td>
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<td>IIIA</td>
<td>NS</td>
<td>1.8</td>
<td>VAD × 5</td>
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<td>43 months</td>
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<tr>
<td>4</td>
<td>56</td>
<td>M</td>
<td>IIIA</td>
<td>IgGκ</td>
<td>*</td>
<td>VAD × 4</td>
<td>Yes/2</td>
<td>11 months</td>
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<tr>
<td>11</td>
<td>41</td>
<td>F</td>
<td>IIIA</td>
<td>IgAκ</td>
<td>3.0</td>
<td>CVAMP × 6</td>
<td>Yes/1</td>
<td>7 months</td>
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<tr>
<td>12</td>
<td>70</td>
<td>M</td>
<td>IIIA</td>
<td>IgAκ</td>
<td>3.5</td>
<td>M + P</td>
<td>No</td>
<td>N/A</td>
</tr>
</tbody>
</table>

B₂M, beta-2-microglobulin; N/A, not applicable; BJ, Bence-Jones; VAD, vincristine + adriamycin + dexamethasone; IgGκ, immunoglobulin Gκ; MGUS, monoclonal gammapathy of unknown significance; CVAMP, cyclophosphamide + vincristine + adriamycin + methylprednisolone; VAMP, vincristine.

*Not performed as sample taken prior to second assay.
Results

DNA ICL formation in plasma cells

DNA ICL formation in isolated plasma cells from melphalan naive and treated patients was detected by using the modified Comet assay. Cells were treated with melphalan for 1 hour followed by 16 hours drug-free after incubation. This time has previously been shown to be the peak time of crosslink formation for melphalan in cell lines and was confirmed for myeloma cells in the present study (data not shown).

Examples of typical Comet images are shown in Figure 1. In control nondrug treated, unirradiated plasma cells no DNA damage was detected, and the high-molecular-weight supercoiled DNA remained intact as shown in Figure 1A. Following irradiation of cells with 10 Gy to introduce a fixed level of random DNA single strand breaks, the resulting shorter fragments of DNA migrated from the bulk of the DNA during electrophoresis to produce the typical comet images (Figure 1B). The extent of DNA damage was quantitated by image analysis to produce a tail moment, defined as the product of the percentage DNA in the comet tail and the distance between the means of the head and tail distributions, based on the definition of Olive et al. Following ex vivo treatment with 50 μM melphalan, no drug-induced single strand breakage was detected in unirradiated cells (Figure 1C), and these cells showed similar profiles to the nondrug-treated controls (Figure 1A). When the drug-treated cells were irradiated (Figure 1D), comet tails were visible but with decreased length and intensity compared with irradiated controls. The comet heads were larger and of greater intensity compared with the nondrug treated irradiated control cells (Figure 1B) because of the retention of DNA by the melphalan-induced ICLs. The decrease in comet tail moment compared with nondrug-treated irradiated controls was used to quantitate the level of DNA interstrand crosslinking.

The sensitivity of the Comet assay to detect melphalan-induced ICLs in plasma cells from a single patient is illustrated in Figure 2. A linear decrease in tail moment was observed with increasing doses of melphalan, and ICLs could be easily detected at levels as low as 1 μM. At each dose point the mean tail moment from 50 cells was quantitated, and variability was small, indicating that formation of melphalan-induced ICLs was homogeneous throughout the plasma cell population.

Comparison of DNA ICL formation in plasma cells from melphalan-naive and melphalan-treated patients

ICL formation was quantitated in plasma cells from 6 patients who were melphalan-naive and 6 patients previously treated with melphalan as shown in Figure 3. Data were expressed as the percentage decrease in tail moment as compared with nondrug-treated controls from the same patient. ICL formation was dose dependent with all patients displaying more than 70% crosslinking at 100 μM (Figure 3A,B) regardless of previous melphalan exposure. No significant differences in ICL formation were seen between the 2 populations (Figure 3C), indicating that intracellular melphalan levels were similar in cells from each patient population, allowing an equal level of DNA alkylation.

Comparison of DNA ICL repair in plasma cells from melphalan-naive and melphalan-treated patients

Repair of melphalan-induced ICLs was studied in both patient populations 24 hours following the 16 hours after incubation required for peak of ICL formation (ie, 40 hours following initial drug exposure). Figure 4 shows representative comet images. Samples were taken from a melphalan-naive (Myeloma 1) and a patient who relapsed following a melphalan-conditioned autograft (Myeloma 2). Both patients showed a similar reduction in comet tail length because of the presence of melphalan-induced ICLs (t = 16 hours) when compared with nondrug-treated irradiated control subjects (no melphalan). However, no repair of melphalan-induced ICLs at 40 hours was observed in plasma cells from the naive patient, and the reduction in comet tail persisted. In contrast, significant repair was observed in the relapsed patient, and the comet tail returns to a similar level as the nondrug-treated irradiated control subject.

The repair data from all patient samples is summarized in Table 2. No repair was seen in any of the 6 naive patients and was regardless of previous exposure to the VAD regime. In contrast all patients who had received prior melphalan therapy, both oral and high dose, displayed significant repair at 40 hours, ranging from 42% to 100%. ICL repair was homogeneous throughout all the plasma cell populations. In some patients it was possible to detect ICL repair in sequential samples taken some months later. For
example, patient 9 demonstrated 54% repair at initial sampling (Table 2) compared with 58% repair 13 months later, whereas patient 12 demonstrated 48% repair (Table 2) compared with 50% 4 months later. In one patient, patient 4, samples were taken at initial diagnosis and at relapse following a melphalan-conditioned autograft (Table 2). No repair of melphalan-induced ICLs was observed prior to melphalan therapy. In contrast, 56% repair was seen in plasma cells taken at relapse after melphalan exposure.

The persistence of ICL in the naive patient population should result in increased sensitivity to melphalan. Indeed, when there were sufficient cell numbers to perform the MTT cytotoxicity assay, the in vitro sensitivity to melphalan in plasma cells was found to correlate with ICL repair (Table 2).

**Discussion**

The Comet assay has been used to detect both the formation and repair of melphalan-induced ICLs in plasma cells from 12 patients with MM who were either melphalan naive or melphalan treated. The formation of melphalan-induced ICLs was found to be similar in both patient populations and was unaffected by previous melphalan exposure. This finding suggests that upstream resistance mechanisms such as drug transport and detoxification, although likely to be associated with VAD resistance, particularly the acquisition of the multidrug resistance phenotype, do not play a significant role in the development of melphalan resistance in MM. Such mechanisms have previously been found to show no correlation with resistance to nitrogen mustards in chronic lymphocytic leukemia (CLL).

We have shown a marked difference in the ability to repair melphalan-induced ICLs between plasma cells from melphalan-naive patients and melphalan-treated patients. Increased repair is independent of any previous treatment with VAD/CVAMP (cyclophosphamide, vincristine, adriamycin, methylprednisolone) and can be detected in both patients who have relapsed after a melphalan-conditioned autograft and those receiving oral melphalan therapy. The analysis of patient 4 at both initial presentation and relapse following a melphalan-conditioned autograft clearly illustrates the development of this mechanism after melphalan exposure. The ultimate aim is to study all patients at presentation and relapse; however, it must be noted that time to relapse following an autograft is approximately 3 years. The role of DNA repair in clinical resistance to agents such as the nitrogen mustards is becoming increasingly important. Sequential sampling from melphalan-treated patients indicates that similar levels of repair can be detected between intervals of several months prior to further therapy, indicating a stable phenotype. For example, at initial sampling patient 9 demonstrated 54% repair compared with 58% repair observed 13 months later. Because the Comet assay analyzes single cells, heterogeneity of response can be determined. The level

**Table 2. Repair of melphalan-induced interstrand crosslinks in plasma cells and in vitro cytotoxicity to melphalan**

<table>
<thead>
<tr>
<th>Patient</th>
<th>Melphalan</th>
<th>% Repair at 40 h</th>
<th>IC50 (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Naive</td>
<td>0</td>
<td>ND</td>
</tr>
<tr>
<td>2</td>
<td>Naive</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>3</td>
<td>Naive</td>
<td>0</td>
<td>ND</td>
</tr>
<tr>
<td>4</td>
<td>Naive</td>
<td>0</td>
<td>ND</td>
</tr>
<tr>
<td>5</td>
<td>Naive</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>6</td>
<td>Naive</td>
<td>0</td>
<td>ND</td>
</tr>
<tr>
<td>7</td>
<td>High dose</td>
<td>100</td>
<td>ND</td>
</tr>
<tr>
<td>8</td>
<td>High dose</td>
<td>72</td>
<td>55</td>
</tr>
<tr>
<td>9</td>
<td>High dose</td>
<td>54</td>
<td>ND</td>
</tr>
<tr>
<td>10</td>
<td>High dose</td>
<td>42</td>
<td>23</td>
</tr>
<tr>
<td>11</td>
<td>High dose</td>
<td>65</td>
<td>ND</td>
</tr>
<tr>
<td>12</td>
<td>Oral</td>
<td>48</td>
<td>52</td>
</tr>
</tbody>
</table>

IC50, dose of melphalan giving 50% inhibition of cell growth; ND indicates not determined.
of repair was, however, found to be equivalent within an individual cell sample, suggesting that the underlying mechanism is homogeneous throughout the plasma cell population following melphalan therapy.

Resistance to melphalan and other nitrogen mustards has been studied in CLL and was found not to be the result of a drug transport defect or enhanced metabolism. However, significantly increased repair of melphalan-induced ICL was observed ex vivo in lymphocytes taken from patients with CLL known to be resistant to nitrogen mustards compared with untreated patients. A number of repair mechanisms have been studied in CLL in an attempt to elucidate the mechanism by which this increased removal of ICL occurs. No significant differences in the expression and activity of enzymes involved in base excision repair and nucleotide excision repair were seen in lymphocytes isolated from untreated patients and patients with CLL resistant to nitrogen mustard. However, a clear linear correlation between in vitro cytotoxicity to chlorambucil and DNA protein kinase activity and localization of DNA repair proteins suggests a possible role for recombination repair processes.

Bifunctional alkylating agents such as melphalan are capable of producing a number of DNA adducts and, although the ICL is considered to be the cytotoxic lesion, only constitutively a small proportion of the total adducts formed, the majority formed being monoadducts. Apurinic sites can result from monoalkyla-
tions that can be converted to DNA strand breaks under alkaline conditions and that could potentially interfere with the detection of ICLs by the Comet assay. However, no evidence of strand break formation was detected in any of the patients’ unirradiated, melphalan-treated cells. Indeed, studies with mono-melphalan, which produces equivalent levels of monoalkylation to melphalan but is unable to form ICLs, have not revealed any significant formation of single strand breaks under the conditions used in the Comet assay (data not shown). Drug-induced apoptosis after melphalan treatment is possible particularly following the 40 hours after incubation. The Comet assay can be used to detect apoptosis within a single cell population. Because of the severe fragmentation of DNA during the apoptotic process, almost the entire DNA within a cell migrates to the tail and the tail characteristically appears “detached” from the head. In the cell populations analyzed in the present study however, the number of apoptotic cells was small and did not interfere with the analysis of ICLs.

We have clearly shown that repair of melphalan-induced ICLs may be the major mechanism for the development of melphalan resistance in MM. These data suggest that the role of melphalan as a preparative agent in MM may require further assessment particularly in patients who have previously received melphalan as a conditioning regime. This finding may be of relevance in patients undergoing a second transplantation as part of a tandem autograft protocol or as treatment of relapsed disease. The data also suggest that the Comet assay may be of value in both the detection of emergence of a resistant phenotype and the prediction of response to melphalan therapy in MM. The complex molecular mechanism by which human cells repair ICL is becoming established.

The present study suggests that pharmacologic manipulation of this mechanism may be of value in optimizing the effect of repeated courses of melphalan chemotherapy.

References


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Repair of DNA interstrand crosslinks as a mechanism of clinical resistance to melphalan in multiple myeloma

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