Mechanisms of Lymphocytotoxicity Induced by Extracorporeal Photochemotherapy for Cutaneous T Cell Lymphoma

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Abstract
Extracorporeal photochemotherapy is an effective treatment for cutaneous T cell lymphoma but its mode of action is uncertain. The reduction in viability of patients' photoirradiated buffy coat lymphocytes was correlated with a 35% increase in DNA single-strand breaks and marked decreases in cellular ATP and NAD levels (to 58 and 34% of control, respectively) immediately after photoirradiation. Complementary in vitro studies were conducted with normal human peripheral blood lymphocytes using a Therakos ultraviolet A (UVA) light box. UVA light was cytotoxic on its own but was potentiated by 8-methoxypsoralen. 3-aminobenzamide, a poly (ADP-ribose) synthetase inhibitor, mitigated the cytotoxic effect of ultraviolet A light in the presence of 8-methoxypsoralen in lymphocytes and reduced the amount of nucleotide depletion they caused. 10 J/cm² of UVA light in the presence of 300 ng/ml 8-methoxypsoralen increased the poly (ADP-ribose) synthetase activity of peripheral blood lymphocytes. Exposing lymphocytes to deoxycoformycin and deoxyadenosine was found to induce biochemical and physical effects similar to those of photochemotherapy. In summary, we have shown that the lymphocytotoxic effect of extracorporeal photochemotherapy for cutaneous T cell lymphoma is apparently mediated by DNA damage, subsequent poly (ADP-ribose)ylation and adenine nucleotide depletion. It is not known how the DNA damage and resultant biochemical effects relate to the possible immunological mechanism of extracorporeal photochemotherapy; however, it is possible that its effects can be mimicked by other DNA-damaging agents. (J. Clin. Invest. 1990. 86:2080–2085.) Key words: poly (ADP-ribose)ylation • nucleotide depletion • DNA strand breaks • 8-methoxy- psoralen • deoxycoformycin

Introduction
Cutaneous T cell lymphoma (CTCL)¹ is a malignant monoclonal proliferation of T lymphocytes usually of the helper pheno-

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1. Abbreviations used in this paper: CTCL, cutaneous T cell lymphoma; dAdo, deoxyadenosine; dCF, deoxycoformycin; ECPC, extracorporeal photochemotherapy; 8-MOP, 8-methoxypsoralen; % D, percent double-stranded DNA remaining; SSBs, DNA single-strand breaks; 3-ABA, 3-aminobenzamide.

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lymphocytes, complemented by in vitro experiments with PBL using a UVA light source and 8-MOP. We show the potentiation of UVA damage to lymphocyte DNA by 8-MOP and the correlation of cytotoxicity with subsequent adenine nucleotide depletion, apparently secondary to poly (ADP-riboseylation).

Methods

Photopheresis procedure. All patients were treated with the UVAR photopheresis system (Therakos, Inc., West Chester, PA). This machine is the vehicle for the collection of buffy coat lymphocytes and also contains the clear plastic disposable cassette where lymphocytes are exposed to UVA light. At 0 h, the patients ingested 8-MOP tablets at a dose of 0.6 mg/kg. 1.5 h later, the patients were venipunctured and connected to the photopheresis machine. 15 min later when the first buffy coat sample (40 ml) had been collected the UVA source was switched on to irradiate the buffy coat sample. Over the next 1.5–2 h (depending on the rate of venous blood flow) a further five buffy coat samples were collected. In total, there were six cycles of blood withdrawal, centrifugation, and reinfusion. Red cells and all additional plasma were returned to the patient before beginning the next collection cycle. At this stage (3.25–3.75 h after tablet ingestion) the final volume for photoinactivation was reached and the patient was disconnected from the machine. The final volume of 740 ml consisted of 240 ml buffy coat, 300 ml plasma, and 200 ml heparinized normal saline. Photoinactivation continued for another 1.5 h. The hematocrit of the leukocyte-enriched blood was always less than 7% because red cells block leukocyte absorption of UV energy. The mean number of cells reinfused was $3 \times 10^{10}$ (range $2.6-4.9 \times 10^{10}$), which is about 25% of the total circulating numbers of lymphocytes.

Treatment policy and assessment of response. Patients were treated on 2 consecutive d every 4 wk for 6 mo, with treatment frequency modified according to clinical response. Disease progress was monitored by measurement of standardized skin scores (4), skin biopsies, and by regular clinical photographs. Briefly, all skin was graded from 0 to 4, with 4 representing the most severe disease. This was then multiplied by the percentage of total surface area involved so the maximum possible score is 400. Skin score assessments were made before the commencement of therapy and then monthly before each course of ECPC by an experienced dermatologist. Response was defined as a 25% improvement in the skin score, sustainable over a 1-mo period. Complete reassessment of all patients occurred 6 mo after starting treatment.

Processing of buffy coat specimens. Specimens of buffy coat blood were obtained from the first buffy coat specimen and also after UV irradiation was complete, immediately before reinfusion to the patient. About 20 ml of buffy coat blood was underlaid with 5 ml of Ficoll-Hypaque and spun at 450 g for 25 min. The cloudy mononuclear cell fraction was removed and counted. After washing in PBS the lymphocytes were resuspended in RPMI 1640 plus 10% FCS. After this handling, the mean composition of the cells (±SE) was: lymphocytes 74±9%, granulocytes (including band forms) 22±8%, and monocytes 3±1%.

Viability. For viability studies, lymphocytes from the buffy coat samples were suspended in RPMI 1640 plus 10% FCS and incubated at 37°C. Viability was determined by the ability of cells to exclude 0.5% trypan blue. Viability was assessed every 24 to 96 h after photoinactivation and compared with the 24–96-h viabilities of the unirradiated buffy coat lymphocytes that were similarly handled.

Measurement of DNA single-strand breaks. About $1 \times 10^{7}$ lymphocytes were spun to form a pellet and then processed in triplicate samples according to the fluorometric method of Birnboim and Jevcek (7). The percentage of double-stranded DNA remaining (% D) of the first (control) specimen was corrected to 100 and the % D value obtained for the second specimen was expressed as a percentage of this. The increase in DNA single-strand breaks (SSBs) was calculated by subtracting the % D value from 100. In vitro processing of lymphocytes from normal human donors and their exposure to 8-MOP did not affect cellular viability; viability was unimpaired at the time of measuring DNA damage.

Measurement of NAD and ATP content. Approximately $2 \times 10^{7}$ lymphocytes were centrifuged to form a pellet, extracted with ice-cold perchloric acid, and their NAD and ATP content measured by HPLC as previously described (8). The lymphocytes obtained from the unirradiated buffy coat specimen were regarded as having normal (control) NAD and ATP content and the nucleotide content of the lymphocytes from the second specimen was calculated as a percentage of this control value.

Normal human peripheral blood lymphocytes. Fresh human peripheral lymphocytes were obtained by venesection or from normal human volunteers' buffy coat specimens (Victorian Red Cross Blood Bank, Victoria, Australia) which were available within 1 h of leukapheresis. The 40-ml buffy coat specimen was overlaid onto 5–10 ml of Ficoll-Hypaque and spun at 450 g for 25 min. Two washes in PBS were followed by suspension in RPMI 1640 plus 10% FCS and antibiotics.

UV irradiation of lymphocytes. For UV irradiation, cells were suspended in culture medium at about $5 \times 10^{7}$ cells/ml. 8-MOP was freshly made up from stock solutions (1 mg/ml) and was added to cells shielded from the light for at least 15 min before photoinactivation. 5–10-ml aliquots of cell suspension were irradiated in 25-cm² plastic tissue culture flasks (Costar Data Packaging Corp., Cambridge, MA) with a Therakos research light box which contained a Photосette-R UVA light assembly mimicking the therapeutic equipment used for humans. Two banks of six lamps were located behind windows of transparent glass 10 cm apart. The lamps were operated at the maximum power setting of 10 for at least 10 min before experiments. The Commonwealth Scientific and Industrial Research Organization National Measurement Laboratory of Australia measured the ultraviolet irradiance in the center of the lower window as 31 mW/cm² (but only 20 mW/cm² in the periphery). A 32-s exposure therefore was equivalent to 1 J/cm² of energy. The center of the panel was used exclusively for UV irradiation.

Measurement of cellular poly (ADP-ribose) synthetase activity. To measure poly (ADP-ribose) synthetase levels, the method of Berger (9) was used, in which cells were rendered permeable to an exogenously supplied nucleotide and then the incorporation of $[^3H] NAD into acid-precipitable material at 30°C was measured.

HPLC measurement of serum 8-MOP levels. Serum 8-MOP levels were measured according to the HPLC method of Puglisi (10). Blood samples were taken from all patients at hourly intervals during the period of photopheresis, so the peak 8-MOP level and its timing could be determined.

Materials Aqueous liquid scintillant and $[^3H] NAD (3 Ci/mmol) were from Amersham International, Amersham, UK. Deoxycoformycin was provided by Parke-Davis, Morris Plains, NJ, and deoxyadenosine was from Sigma Chemical Co., St. Louis, MO. All other chemicals were from BDH Chemicals Ltd., Kilsyth, Victoria, Australia. Details of the materials for HPLC measurement of nucleotides and the fluorometric assay of DNA SSBs have been described previously (Marks, D. I., and R. M. Fox, submitted for publication).

Results

Clinical response to extracorporeal photochemotherapy. Four patients with CTCL (three males and one female, mean age of 69) were treated with ECPC. Patients received 16–25 courses (mean 19) over a mean period of 25 wk. One patient died 7.5 mo after the commencement of treatment of myocardial infarction and pulmonary embolism. The four patients had significant palliation of their disease. The mean maximum decrease in skin score was 41% (range 29–55%, Table 1).

Effects of extracorporeal photochemotherapy on lymphocyte viability. To assess the effect of the ECPC procedure on the patients' harvested lymphocytes, trypan blue viability was
studied daily for 4 d following the procedure. This was studied in nine patient samples (from nine different treatment cycles) from the four patients. Viability progressively declined over several days, at a rate similar to that reported by Edelson (4). Less than 5% of the treated cells were viable at 96 h (Table I).

**DNA single-strand breaks.** To explore the mechanism of action of the ECPC-induced in vitro lymphocytotoxicity, DNA damage was assessed. Although other forms of DNA damage may have been induced by the UVA/8-MOP, DNA single strand breaks were assayed by the fluorometric technique. DNA damage was assessed in samples of buffy coat lymphocytes after photoirradiation just before reinfusion into the patient. The viability of photoirradiated cells at the time of measuring DNA SSBs was the same as nonphotoirradiated control cells (85%). The control % D value for nonphotoirradiated lymphocytes was corrected to 100% and the treated lymphocyte sample expressed as a percentage of this. The absolute control % D value for the patients' cells (and for normal PBL) was 79 (range 60–97). All samples assessed had a marked increase in DNA SSBs (Table I) but the extent of damage varied considerably among different patients, ranging from 9 to 60%. DNA damage was assessed twice in all except patient 3. There was < 15% variation between readings from samples at different time points; only the mean is stated. There was no correlation between the number of malignant cells and the percentage of DNA SSBs.

**Adenine nucleotide content.** In view of the apparent association between DNA damage and lymphocytotoxicity, the extent of ATP and NAD depletion, possibly secondary to poly(ADP-ribosylation), was investigated. Eight patient samples of immediate postirradiation buffy coat lymphocytes (eight treatment cycles) from four different patients were analyzed. NAD content was measured in three of the four patients. All lymphocyte samples assessed showed depletion of nucleotides. The decline in NAD content was much more marked than the fall in ATP content (Table I). The fall in the nucleotide content of patients' lymphocytes did not correlate with the degree of DNA damage to them. Absolute control NAD and ATP content (in patients' and normal PBL) were 60 (48–74) and 202 (180–220) pmol/10⁶ lymphocytes, respectively.

**Correlation of biochemical parameters with clinical response.** The biochemical and physical damage to buffy coat lymphocytes demonstrated above from CTCL patients has not been previously documented nor correlated with the clinical response of CTCL to ECPC. Four patients with CTCL were evaluable for response (Table I). This number of patients precludes a meaningful statistical analysis of the relationship between response and the parameters of cell injury or death. However, the percentage of DNA SSBs was strongly associated with the decline in cellular viability (simple regression analysis, R² = 0.877, Fig. 1). There was no correlation between the number of cells a patient had reinfused and that patient's clinical response (maximum decrease in skin score) (MDSS, data not shown).

**Serum 8-MOP levels.** The 8-MOP levels of all patients were found to exceed 50 ng/ml during the entire period of photopheresis. The mean peak level was 168 ng/ml and occurred at an average of 2 h after ingestion. Peak serum 8-MOP levels did not correlate with clinical response, in vitro lymphocytotoxicity or with any of the biochemical parameters associated with cell death.

| Table I. Clinical and Biochemical Data about Four Patients with CTCL |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Patient No.     | Initial skin    | Minimum skin    | MDSS            | Mean increase   | 24-hour viability| Mean NAD        |
|                 | score           | score           | %               | DNA SSBs        | %               | content*        | Mean ATP        |
| 1               | 253             | 154             | 39              | 9               | 71              | ND*             | 48              |
| 2               | 345             | 208             | 40              | 32              | 44              | 24              | 66              |
| 3               | 336             | 237             | 29              | 60              | 16              | 33              | 83              |
| 4               | 181             | 81              | 55              | 39              | 53              | 44              | 34              |
| Mean            | 279             | 170             | 41              | 35              | 46              | 34              | 58              |

* The NAD and ATP contents of buffy coat cells were measured immediately after photoirradiation. DNA SSBs were measured immediately after photoirradiation by the fluorometric unwinding method (see Methods). † Not determined.

Samples were taken from patients' buffy coat collections before (control) and after photopheradiation, then placed in RPMI 1640 plus 10% FCS, and trypan blue viability measured for the next 96 h. The results shown represent the mean of nine patient samples from four patients. The mean control cell viability after exposure to 8-MOP and processing was 85%.

**Table II. Effect of Extracorporeal Photochemotherapy on Lymphocyte Viability**

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<tr>
<th>Lymphocyte sample</th>
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<tr>
<td>Control</td>
<td>100</td>
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<tr>
<td>24 h postECPC</td>
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<td>48 h postECPC</td>
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<td>96 h postECPC</td>
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Samples were taken from patients' buffy coat collections before (control) and after photopheradiation, then placed in RPMI 1640 plus 10% FCS, and trypan blue viability measured for the next 96 h. The results shown represent the mean of nine patient samples from four patients. The mean control cell viability after exposure to 8-MOP and processing was 85%.

![Figure 1. Percent increase in DNA single strand breaks vs. percent decrease in cellular (trypan blue) viability. Simple regression analysis yields an R² of 0.877. The intercept for the graph is 82.1 (38.9–125.2, 95% confidence limits) and the gradient is −1.03 (−2.12–0.06).](image-url)
In vitro experiments on the mechanisms of 8-MOP and UVA photochemotherapy-induced cytotoxicity in PBL. Sham photochemotherapy treatment experiments were conducted with peripheral blood lymphocytes from normal blood bank volunteers. These experiments enabled variation of the intensity of UVA light, the concentration of 8-MOP, and the interaction between 8-MOP and UVA light to be studied in an attempt to elucidate the mechanism of lymphocytotoxicity.

Viability. Control experiments revealed that 8-MOP alone (without UV irradiation) was not toxic to PBL: 10–300 ng/ml did not affect corrected viability after 72 h incubation at 37°C. The effect on lymphocyte viability of a wide range of doses of UVA light (0.1–10 J/cm²) was studied: the results for 1 and 2 J/cm² (the dose given to patients) are shown in Fig. 2 A. The addition of 8-MOP clearly potentiated the cytotoxicity caused by UVA light; however, the potentiating effect was dependent on the dose of UVA light. At doses of 3 J/cm² and above there was minimal potentiation (data not shown), while at 1–2 J/cm² the potentiation was marked. 8-MOP concentrations higher than 50 ng/ml did not further increase cytotoxicity in PBL (data not shown).

Many cytotoxic agents that are thought to mediate their cytotoxic effect via poly (ADP-ribose)lation have their cytotoxicity reduced (or completely eliminated) by the addition of specific poly (ADP-ribose) synthetase inhibitors. The addition of 3-aminobenzamide (3-ABA) (2 mM), a dose sufficient to inhibit poly (ADP-ribose) synthetase but which did not affect lymphocyte viability, decreased the cytotoxic effect of exposing PBL to 2 J/cm² UVA and 300 ng/ml 8-MOP. This dose of 8-MOP was chosen to maintain an excess of the drug.

Figure 2. Effects on viability. (A) PBL were exposed to UVA light (1–2 J/cm²) with or without 8-MOP (300 ng/ml) and their corrected viability determined by trypan blue exclusion every 24 h. A, 1 J/cm² UVA; B, 1 J/cm² UVA + 8-MOP; C, 2 J/cm² UVA; D, 2 J/cm² UVA + 8-MOP. Mean of triplicate samples from a representative experiment (with multiple replicate experiments). (B) PBL were exposed to UVA light (2 J/cm²) and 300 ng/ml 8-MOP with or without 2 mM 3-ABA and corrected viability was determined at 24 and 48 h. This is compared with incubating PBL with 2 mM 3-ABA alone. A, 3-ABA alone; B, 2 J/cm² UVA + 8-MOP; C, 2 J/cm² UVA + 8-MOP + 3-ABA. The data represent the results (performed in triplicate) from a typical sample. Multiple replicate experiments show the same pattern. The two figures represent experiments involving PBL from different donors. Control lymphocyte viability was 92%.

viability of the 3-ABA and UVA/8-MOP treated cells (99% at 48 h) was significantly different from the control UVA/8-MOP group (61%, \( P < 0.001 \)) (Fig. 2 B). The protective effect occurred whether the 3-ABA was added before or after irradiation; this excludes the possibility that 3-ABA mitigated the cytotoxic effect merely by decreasing the dose of UVA received by the PBL.

DNA damage. The DNA damage for a range (1–5 J/cm²) of UVA light doses was studied 2 h after irradiation (Fig. 3). The DNA-damaging effects of UVA light on PBL was potentiated by the addition of 8-MOP. However, the extent of this potentiating effect was dependent on the dose of UVA light and the timing of the assay. Edelson (4) reported that the dose of UVA light delivered to his patients’ buffy coat lymphocytes was 1–2 J/cm²; we have shown that in vitro this was the dose where 8-MOP maximally potentiated the DNA damage caused by UVA light. In the sham experiments this dose induced a level of DNA strand breaks similar in extent to those in the patients’ samples. At 1.5 J/cm² 8-MOP increased the number of DNA SSBs by more than twofold. There was far less damage at 24 h (<15% increase in DNA SSBs) than there was at 2 h, presumably reflecting DNA repair (data not shown). 5 J/cm² of UVA light caused only slightly more DNA SSBs than 2 J/cm²; this suggests a ceiling effect.

Nucleotide content. Exposures of PBL to 1–40 J/cm² UVA light alone caused a nonlinear dose-dependent depletion of NAD and ATP 2 h after exposure (Fig. 4 A). Between 1 and 10 J/cm² there was a steep increase in nucleotide depletion with further depletion only occurring at very high doses of UVA light. 8-MOP potentiated the NAD-depleting effects of UVA light alone. At 1 J/cm² UVA light exposure depleted PBL of 11% of their NAD content; adding 8-MOP (300 ng/ml) increased the total NAD depletion to 34% (representative experiment, mean of five observations). At higher UVA doses (2–10 J/cm²) this potentiating effect was not apparent (data not shown). The addition of 8-MOP to 1–2 J/cm² UVA light did not further increase ATP depletion.

2 mM 3-ABA also mitigated the nucleotide depletion caused by 10 J/cm² of UVA light and continuous exposure to 300 ng/ml 8-MOP (Fig. 4 B). This dose of UVA light was chosen because it caused substantial NAD and ATP depletion; therefore, the mitigating effect of 3-ABA could be clearly demonstrated.

Poly (ADP-ribose) synthetase activity. PBL were exposed to 10 J/cm² UVA light in the presence of 300 ng/ml 8-MOP; at 2 h poly (ADP-ribose) synthetase activity rose to 160% of control (three experiments of three to five samples). Significant elevations in enzyme activity were not demonstrable with lower doses of UVA light. 8-MOP alone (300 ng/ml) did not affect...
poly (ADP-ribose) synthetase activity 2 h after exposure. 10 J/cm² UVA light alone raised enzyme activity to 156% of control (two experiments of three samples). Control lymphocyte poly (ADP-ribose) synthetase activity was 1.20 pmol min⁻¹ (10⁶ cells)⁻¹.

Comparison between photochemotherapy and deoxyadenosine lymphocytotoxicity. Deoxycoformycin and deoxyadenosine lymphocytotoxicity has been previously shown to be associated with DNA damage and adenine nucleotide depletion presumably secondary to stimulation of poly (ADP-ribosyl)ation (11). A 4-h exposure of PBL to 1 μM deoxycoformycin (dCF) and 10 μM deoxyadenosine (dAdo) caused 85% cell death in 72 h (Fig. 5 A), comparable to the delayed lymphocytotoxicity caused by ECPC. 2 h after drug exposure there was a 19% increase in DNA SSBs. At this time ATP was 66% of control levels and NAD 76% of control (Fig. 5 B): this is compared to the ex vivo effects of ECPC on lymphocytes from CTCL patients. dCF and dAdo caused comparable ATP depletion but less decrease in NAD.

**Discussion**

This study has documented the relationship between cytotoxicity and the biochemical and physical changes that occur in lymphocytes from patients with CTCL treated ex vivo with photochemotherapy. Photoactivated psoralen-induced DNA damage to cells has been well documented (5) but the way in which this DNA damage resulted in cell death was uncertain. Our studies of the effects of ECPC on buffy coat lymphocytes from patients with CTCL were complemented by in vitro experiments with PBL using a UVA light source that mimicked the effect of photochemotherapy.

Although UVA light caused DNA strand breaks on its own, there was clear evidence that 8-MOP can potentiate the damage caused by UVA light. The degree of this potentiation was highly dependent on the dose of UVA light. At doses of 3 J/cm² and above potentiation was minimal: UVA light alone caused most of the damage. However, when the dose of UVA light was 1 J/cm², the addition of 8-MOP significantly increased the degree of cell kill, the NAD depletion, and the number of DNA SSBs UVA light produced. The maximum potentiation of DNA damage by 8-MOP occurred at 1.5 J/cm². These observations provide the in vitro rationale for using 1–2 J/cm² as the dose of UVA light delivered to buffy coat lymphocytes in the clinical situation. Our four patients had a 41% mean increase in DNA SSBs suggesting that 1–2 J/cm² of UVA light was the dose delivered to them. It is also noteworthy that, even with lethal doses (2 J/cm²) of UVA light and 8-MOP, the DNA damage present at 2 h was much greater than that at 24 h after exposure. It appears that this early DNA damage caused a series of biochemical events that irreversibly damaged the cell and that subsequent DNA repair did not affect this.

This UVA/8-MOP-associated DNA damage is associated with adenine nucleotide depletion. This study also presents some evidence that stimulation of poly (ADP-ribosyl)ation and consequent nucleotide depletion are involved in the lymphocytotoxic effect of photoactivated psoralens. Photoinactivation in the presence of 8-MOP caused significantly elevated levels of poly (ADP-ribosyl) synthetase activity. However, elevated enzyme activity levels were only found at high doses (10 J/cm²) of UVA light. The failure to demonstrate elevated enzyme levels at 1–2 J/cm² may be due to the insensitivity of the assay or may reflect the transient nature of the rise in enzyme activity. The poly (ADP-ribosyl) synthetase inhibitor, 3-ABA, mitigated the cytotoxicity of photoinactivation and reduced the nucleotide depletion it caused. These three pieces of data are regarded as important evidence for the involvement of poly (ADP-ribosyl)ation (11). Poly (ADP-ribosyl)ation accounts for the majority of NAD turnover in the lymphocyte: it would appear to be the most likely cause for the changes in NAD content that we have demonstrated.
A degree of caution is required in the interpretation of the correlations between clinical response and various biochemical parameters. The number of patients was small and although response correlated with ATP depletion it did not correlate with other parameters of cell kill.

In vitro studies in which normal PBL were treated with dCF and dAdo showed many similarities with the effects of ECPC on lymphocytes of patients with CTCL. The rate of cell kill, DNA strand breakage and extent of NAD and ATP depletion were all similar. Carson (12) originally demonstrated that dAdo and dCF caused DNA strand breaks, depleted cells of NAD and ATP, and increased levels of poly (ADP-ribose) synthesis. He found that nicotinamide and 3-ABA protected against these changes. We have shown that photoirradiation is also apparently associated with stimulation of poly (ADP-riboseyl)ation. Although 3-ABA can prevent adenine nucleotide depletion, it appears the biochemical damage to cells is irreversible (13).

It appears that photochemotherapy may have an immunologic mechanism of action. First, the superiority of ECPC to simple leukapheresis is evidence that reinfusion of the damaged (dying) cells is important. Delayed death of the reinfused, damaged lymphocytes may be important; there may need to be sufficient time for them to excite an immunologic response in patients with CTCL. Second, immunocompetent patients respond better than those with impaired immunity (4). Third, there is a reduction in malignant cells in the skin (cells that have not been exposed to ECPC) that suggests they have been eradicated by a mechanism of host origin (4). It has been postulated that the reinfused damaged cells induce the proliferation of lymphocytes bearing antiidiotype receptors that recognize the malignant clone.

We have shown the increased expression of various activation markers by lymphocytes from CTCL patients soon after treatment with ECPC (14). The UVA/8-MOP-induced changes in the reinfused peripheral blood lymphocytes may induce antigenic changes in their cell surface membranes, which might be responsible for this immune stimulation.

The practical implications of the findings in this report are that the extracorporeal treatment of patients'uffy coat with a combination of dAdo plus dCF (or other lymphocytotoxic agents) may achieve similar effects to ECPC. This is of some importance when the potential wider application of ECPC to autoimmune disease is considered. Replacement of ECPC by exposure of the Buffy coat specimens to conventional DNA-damaging agents would be far more convenient. To test this hypothesis further we are conducting studies comparing infusions of photochemotherapy-affected and cytotoxic drug-damaged splenocytes as treatment of an animal autoimmune model of disease.

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