Combination toxicity of etoposide (VP-16) and photosensitisation with a water-soluble aluminium phthalocyanine in K562 human leukaemic cells

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Summary Etoposide (VP-16) is an anti-cancer drug commonly used against several types of tumours and leukemia, either alone or in combination chemotherapy. Photodynamic therapy (PDT) is another, relatively new modality for treatment of various malignancies. The interactions between VP-16 and PDT, using aluminium tetrasulphophthalocyanine as photosensitiser, in K562 human leukaemic cells were investigated. Cell responses to individual and combined drug treatment under different experimental conditions revealed synergistic drug toxicity. The latter was evident from various events of cell response, including supra-additive accumulation of cells in G2/M cell cycle phase and endonucleolytic DNA fragmentation (apoptosis). The involvement of the cellular antioxidant system in the synergistic interactions of photosensitisation and VP-16 is proposed.

Keywords: etoposide; phthalocyanine; photosensitisation; combination therapy; synergy

Etoposide[4′-dimethylleipodophyllotoxin-9-(4,6-O-ethylidene-β-D-glucopyranoside, VP-16] is an important antineoplastic agent used against several tumour types, either alone (Issel et al., 1984), or in combination therapy (Aisner and Lee, 1991). Etoposide is also commonly used for the treatment of acute myelogenous leukaemia (Champlin and Gale, 1987). The cytotoxicity of VP-16 is generally believed to be based on introduction of DNA damage by drug interference with breakage–reunion reaction of DNA–topoisomerase II (formation of DNA–protein cross-links) (Glisson and Ross, 1987; Liu, 1989) and/or induction of direct DNA strand breaks and adducts (van Maanen et al., 1988a; b; Mans et al., 1991). In contrast to the topoisomerase II (topo II) poisoning by VP-16 itself, the direct inactivation of DNA was suggested to be conjugated with oxidation–reduction activation of VP-16 in the cellular environment (Mans et al., 1990, 1992). In particular, cytochrome P450-dependent mono-oxygenases, peroxidases, prostaglandin synthetase, tyrosinase, etc. may be involved in VP-16 metabolic transformation (van Maanen et al., 1987; Haim et al., 1991; Gantchev et al., 1994a). Evidence has been found that peroxidative metabolic products of VP-16, e.g. the ortho-quinone derivative of etoposide, as well as the short-lived intermediates (phenoxy and semi-quinone free radicals) are involved in various oxidative reactions and DNA damage (Mans et al., 1990, 1991; Sinha et al., 1990). Recent studies suggest that the interactions of VP-16 free radicals with intracellular reductants (thiols, ascorbic acid, etc.) might play an essential role in the cytotoxic activity of the drug as well (Mans et al., 1992; Kagan et al., 1994; Yokomizo et al., 1993).

Light activation of photosensitisers that have been accumulated in tumours is the basis of the photodynamic therapy (PDT) of cancer. Cytotoxic action of photosensitisers may involve oxidative damage to different cell constituents, including depletion of the pool of cell antioxidants (free and protein-bound thiols, ascorbic acid, α-tocopherol, etc.) (Buejttner, 1984; Shopova and Gantchev, 1990; Gantchev and van Lier, 1995). Metallo-phthalocyanines (MePc) constitute a class of dyes proposed as second-generation photodynamic agents to supplant Photofrin, a mixture of porphyrin derivatives, currently used in the clinical treatment of various malignancies. Highly water-soluble phthalocya-

nines, such as di- through tetrasulphonated derivatives (MePcSx, y) partially localise in cytoplasm and photosensitise intracellular generation of hydroxylic and organic free radicals (Gantchev et al., 1994b). Among different effects of photosensitised cell damage, tetrasulphonated Al- and ZnPcSx have been shown to weaken cell viability by inactivation of catalase (Gantchev and van Lier, 1995) and inflicting damage to DNA (Hunting et al., 1987; Gantchev et al., 1994c). We have also shown that these phthalocyanines can initiate oxidative transformations of VP-16 in solution via photosensitised generation of VP-16 phenoxyl radical (Gantchev et al., 1994a).

In view of the above properties of phthalocyanines and the interrelation between the rate of VP-16 oxidative transformation, its cytotoxicity, and the activity of intracellular antioxidant systems, we hypothesised that PDT, in conjunction with etoposide, could result in enhanced cytotoxicity. Analysis of different effects of combination therapy with VP-16 and AlPcS, photosensitisation against K562 chronic myelogenous leukaemia cells is the subject of the present work.

Materials and methods

Chemicals

Etoposide (VP-16) was purchased from Sigma (St Louis, MO, USA). The compound was dissolved in dimethyl sulphoxide (DMSO) at 5 mM, aliquoted and stored at −20°C. Further dilutions were made in RPMI medium immediately before use. Although the final DMSO concentrations of 1–2% in cell cultures were not toxic, all control groups received an equivalent amount of DMSO. Aluminium tetrasulphonated phthalocyanine (AlPcS5) was synthesised in our laboratory, purified by high-performance liquid chromatography (HPLC) and dialysis to homogeneity and dissolved in phosphate-buffered saline (PBS) to yield a stock solution of 1.5–2.0 mM, as measured spectrophotometrically \( \text{E}_{\text{max}} = 2.5 \times 10^3 \text{ cm}^{-1} \) in dimethylformamide). All other chemicals used were of the highest available purity.

Cell culture and drug treatments

Human chronic myelogenous leukaemia cells, K562 (ATCC CCL 243), were grown in RPMI-1640 medium, supplemented with glutamine, 10% fetal bovine serum (FBS), 50 µg ml\(^{-1}\) gentamycin and 10 mM Hepes. All experiments were
performed with exponentially growing (asynchronous) cells. Usually 12–14 h before the experiment, the cells were seeded in a complete RPMI medium (3:1 v/v parts fresh to conditioned medium) at cell density of approximately 0.4 x 10^6 cells ml^-1. On the next day the cell density was adjusted by a small volume of fresh medium to give 0.5 x 10^6 cells ml^-1 and cells were exposed to drugs (VP-16 and AlPcS4 alone, or mixed). All experiments were performed in triplicate flasks. In a typical protocol, after incubation with drugs, cells were first washed once with PBS, and a second time with RPMI (without FBS). After resuspension in a complete RPMI medium (conditioned to fresh medium = 1:3 v/v), cells were exposed to broad spectrum red light (λ > 580 nm; I = 10 mW cm^-2). Cell suspensions (12–15 ml) were exposed to red light in ventilated (green-cap Falcon, 75 cm^-2) incubation flasks and were gently shaken during irradiation. Cells were incubated at 5 x 10^5 cells ml^-1, or diluted to 5 x 10^4 cells ml^-1 to obtain a longer exponential phase. Control experiments performed on cells that were first photosensitised in PBS and then resuspended in culture medium did not show any significant difference in cell survival. Dark incubation with AlPcS4 was not cytotoxic. Exposure to red light of cells incubated with VP-16, in the absence of AlPcS4, did not result in altered VP-16 toxicity.

**Cytoxicity assays and drug interaction analysis**

Cell growth inhibition was monitored by means of dye exclusion (staining with 0.4% Trypan blue) and/or by Coulter counter measurements. Under our experimental conditions, the doubling time of control cells was 22–24 h. The growth suppression activity of drugs was compared by estimation of the exponential rate constants using initial applications of growth curves (for 4–60 h after treatment). Clonogenic assay was performed in a standard fashion by plating cells in soft (0.33%) agar in 35 mm Petri dishes (usually 300, 600 and 1200 cells per dish in triplicate).

To analyse the dose–effect relationships in combination treatment of cells with VP-16 and AlPcS4, photosensitisation, two algorithms were used. A simple estimate was performed using the fractional product method (Velereiote and Liu, 1975). This 'multiplicative' model predicts additivity, antagonism and/or synergism of two drugs based on the comparison between the individual drug effect on cell survival, f_i (expected) = (f_iA) x (f_iB) and the experimentally obtained value of cell survival after combination treatment, (f_iAB). The symbol (f_iAB) stands for 'fraction unaffected' and is equal to 1 – (f_iA) x (f_iB), where f_i (‘fraction affected’) refers to the fraction of cells remaining viable at the various concentrations of the two drugs in combination. By definition, (f_iAB) lower, equal to or larger than f_i (expected) determines the border lines of synergistic (supra-additive), additive and antagonistic drug interactions respectively. In a specially designed set of experiments, we also performed a detailed analysis of drug interactions using the median effect principle. Unlike other methods often used to predict drug interactions in biological systems (usually applicable only to mutually exclusive interactions), the median effect principle may be used to analyse both mutually exclusive and mutually non-exclusive interactions (Chou and Talalay, 1983). We used the linearised median effect equation in the form of log [(f_i – 1) – 1] = m log (D) – m log (Dm), where 'm' is the Hill-type coefficient determining the sigmoidality of the dose–effect curve; 'D' is drug(s) dose; and Dm = IC_{50} is the dose equivalent to produce the median effect. The combination index (CI) was calculated at f_i levels of 0.1 intervals, as described by Chou and Talalay (1983). According to this algorithm, CI values < 1.0 indicate synergism; CI > 1.0, antagonism; and CI approximately 1.0, additivity.

**Flow cytometric cell cycle analysis**

After given time periods following drug removal, cells were washed and resuspended in 200 μL PBS containing 5.5 mM glucose and 1 mM EDTA. While vortexing, 1 ml of cold 70% ethanol was added and cells were incubated at 4°C for 30 min. Thereafter, the cells were transferred to PBS–glucose–EDTA buffer containing 0.5% Tween-20 and incubated for 1 h at 4°C to rehydrate. Following a brief exposure to RNAase, DNA was stained with 50 μg ml^-1 propidium iodide. Before cytoflow analysis, cells were filtered through nylon mesh to avoid clumps. The readings were performed on a Hewlett-Packard computer-controlled Becton Dickinson FACScan instrument. DNA chromatograms were analysed by the SOBR software supplied by the manufacturer. In cases in which high levels of cell debris were present, appropriate gate cut-off settings were used.

**Analysis of internucleosomal DNA fragmentation**

The integrity of DNA from drug-treated and control cells was assessed by agarose electrophoresis. After removal of culture medium at different post-treatment times, cells (0.5–2 x 10^6) were resuspended in 150 μl PBS–glucose–EDTA buffer and transferred to Eppendorf microfuge tubes. Immediately after that, 0.5 ml of lysis buffer [0.8% sodium dodecyl sulphate (SDS) and 100 mM EDTA in 50 mM Tris-HCl, pH = 7.4] was added and cells were incubated at room temperature for 20 min. After addition of 125 μl of 5 mM sodium chloride to each tube, samples were gently mixed and kept overnight at 4°C. On the next day, samples were microcentrifuged at 12,000 x g for 30 min at 4°C and the pellets were carefully removed and discarded. Supernatants were further incubated with 0.45 mg ml^-1 proteinase K for 2 h at 50°C. After addition of 50 μl of 5 M sodium chloride and 1 ml isopropanol, samples were left for 1 h at –20°C and microfuged. The pellet was washed with 70% ethanol and dried under vacuum. Resuspended in 20 μl TE buffer, pellets were incubated with 0.8 mg ml^-1 RNAase A at 37°C for 3 h. After mixing with 4 μl of 6 x loading buffer (TE/bromophenol/glycerol), samples were transferred onto 1.2% agarose gel and electrophoresed at 40 V for 4 h. Gels were stained for 40 min with 10 ng ml^-1 ethidium bromide, destained for 30–60 min in distilled water and photographed under UV light. Series of experiments were performed to selected post-treatment time periods and number of cells required to give early and easily detectable DNA fragmentation in combination with selected multiple drug dose equivalents to reveal drug interaction.

**Results**

**Cell growth and loss of clonogenicity**

To determine the toxicity range of individual drugs and their combination effects, several parameters were examined: incubation times of cells with drugs, drug concentrations and the applied light dose for AlPcS4 activation. The cellular response to drug treatment was followed by monitoring the growth rate and cell clonogenic activity. Growth curves shown in Figure 1 exemplify the evolution of immediate drug toxicity with respect to drug incubation time, concentrations and light dose. For individual drugs and their combinations, growth curves show a proliferation lag period, followed either by predominant cell regrowth or cell death. At low levels of drug treatment (Figure 1a and b), and during the initial proliferation lag-times (e.g. at post-treatment times shorter than 24 h), the number of dead cells did not exceed 5–8% of the total, but cell cycle progression was largely inhibited (see below). Figure 2 demonstrates the effect of etoposide concentration on clonogenic activity of K562 cells, either when the drug was applied alone or in combination with AlPcS4, photosensitisation. Under the employed incubation conditions of cells with VP-16 (20–60 min), the IC_{50} concentration of the drug, as determined from cloning experiments, was in the range from 15 to 6 μM, and was markedly decreased after combined treatment (Figure 2). The IC_{50} of AlPcS4 varied with preirradiation incubation time and light dose (D), and, for example, after 4 h preincubation was
Po-PDT and etoposide interactions

TGGantchev et al

Figure 1 Growth of K562 cells incubated for different times with VP-16 (●); AlPcS4 (▲); or both (●) and exposed to red light. (a) 4 μM AlPcS4 and 2 μM VP-16 for 1 h, light dose (D) = 7.2 J cm⁻²; (b) 30 μM AlPcS4 for 5 h and 7.5 μM VP-16 for 1 h, D = 6 J cm⁻²; (c) 20 μM AlPcS4 for 5 h and 10 μM VP-16 for 1 h, D = 9 J cm⁻²; and (d) 40 μM AlPcS4 for 5 h and 30 μM VP-16 for 2 h, D = 9 J cm⁻². Control cells (○). Incubation time starts after drug removal and light exposure. Averaged data from triplicate experiments.

Figure 2 Clonogenic survival after exposure of K562 cells to different concentrations of VP-16 for 1 h (▲); and after combination treatment when cells were preincubated with 4 μM AlPcS4 for 20 h (○); or with 20 μM AlPcS4 for 4 h (●). Light dose in both cases, (D) = 7.2 J cm⁻². Averaged data from triplicate experiments.

about 20 μM (D = 9 J cm⁻²), and increased when cells were exposed to photosensitiser for a shorter time (e.g. 30–35 μM at 2 h). The simple 'multiplicative' model and data from cloning experiments (e.g. shown in Figure 2) largely predict supra-additive (synergistic) combined toxicity [i.e. \( f_A h_2 < f_A \) (expected) = \((f_A) \times (f_B)\). Materials and methods]. A complete analysis of drug interaction, however, was performed using the median effect principle, and is described below.

Cell cycle arrest

It has been shown previously that continuous exposure to low concentrations of VP-16 allows cells of different origin to traverse the cell cycle until a predominant number of them are blocked with the DNA content of G2 phase, but cannot progress to the mitotic stage (Krishan et al., 1975). In the present study, the cells were incubated with VP-16 for given time periods, and, thereafter, the drug was removed. Under conditions of low drug load (<IC₅₀), VP-16 induced transient G₂/M arrest of K562 cells (Figure 3). Cell size analysis performed by means of Coulter counter measurements showed a parallel and transient increase in cell volume, obviously associated with arrest in G₂-phase. Higher doses and longer exposures to the drug, however, resulted in S- and mixed S/G₂-phase arrest (not shown). It is noteworthy that in this case, and after longer post-treatment incubation times...
Figure 3  Effects after exposure to VP-16 and AlPcS₄ on cell cycle distribution of K562 cells. (1b) DNA chromatograms obtained from cells 26 h after treatment with 20 µM AlPcS₄ for 5 h; (1c) 10 µM VP-16 for 1 h; and (1d) combination of 1b and 1c. Light dose \( D = 9 \text{ J cm}^{-2} \). (2b) Chromatograms obtained 20 h after cell treatment with 4 µM AlPcS₄ for 1 h; (2c) 2 µM VP-16 for 1 h; and (2d) combination of 2b and 2c. Light dose 7.2 J cm⁻². (a) and (2a) show the chromatograms of control cells.

(>24 h), cell cycle analysis was more complicated owing to the presence of high amounts of dead cells and a cell population with quadraploid DNA content. In contrast, AlPcS₄ photosensitisation under a wide range of light doses and preillumination incubation times (2–24 h), induced only G₂/M arrest. When cells were treated with doses around the IC₅₀ values of both drugs, the maximum number of G₂/M-arrested cells was reached after 15–20 h post-treatment incubation and did not exceed a value of approximately 60% of the total (Figure 4a). The cell accumulation in G₂/M was accompanied by a depletion, mainly of S-phase population, with very little variation in the G₁-phase content. During combined treatment, the cells were also arrested in G₂/M (Figure 3). The extent of cell response after combined treatment varied and depended on the individual drug effect. Thus, in conditions when individual drugs were relatively more toxic (e.g. at doses approximately equal to the IC₅₀, see also Figure 1b) and the maximum fraction of cells arrested by individual drug treatment was close to 60%, there was no significant increase in the G₂/M population after combined treatment. However, under these conditions, the maximum number of G₂/M-arrested cells after combined treatment was reached at later times and the total proliferation block lasted longer (Figure 4a). It is noteworthy that in these conditions, during the growth lag period and after that, the number of dead cells remained relatively high, as also seen in DNA chromatograms exhibiting a higher percentage of cells with abnormal DNA content (debris and,
perhaps, apoptotic cells). In contrast, when the toxic effects of individual drugs were low (growth curve shown in Figure 1a), combination treatment induced a supra-additive (synergistic) accumulation of cells in G2/M-phase, as also proved by the 'multiplicative' model (Figure 4b). Synergistic increase of the number of cells accumulated in G2/M-phase was reached under various other conditions, but especially when the toxicity of at least one of the drugs was kept low (Figure 4c).

**Internucleosomal DNA degradation**

K562 leukaemic cells are known to be relatively resistant to VP-16-induced internucleosomal DNA fragmentation (apoptosis) (Dubrez et al., 1995). Thus, exposure of cells to VP-16 of concentrations around or higher than IC₅₀ (5 μM and 10 μM for 1 or 2 h) did not reveal any significant DNA fragmentation during post-treatment incubation times up to 36 h, as assessed by agarose gel electrophoresis (Figure 5, lanes 4 and 5). DNA fragmentation was induced when cells were treated with higher VP-16 concentrations (e.g. 5 × IC₅₀), but the pattern was obscured even when DNA was isolated from 2 × 10⁶ or more cells (not shown). In contrast, AlPcS₄ photosensitisation was a much more potent inducer of DNA fragmentation (apoptosis). Internucleosomal DNA damage was readily detected 4–6 h post treatment in DNA samples isolated from 0.5–1.0 × 10⁶ cells treated with drug dose < IC₅₀ (Figure 5, lane 3). This pattern was persistent and progressively augmented during the next 48–72 h of post-treatment incubation. Figure 5 also shows the results from a typical experiment when cells were exposed to two drug concentrations individually, or to their combination. Under the specified conditions (see figure legend), no extensive DNA laddering took place after treatment with either 5 μM AlPcS₄ or 5 and 10 μM VP-16 alone (lanes 2, 4 and 5). Internucleosomal cleavage, however, was well pronounced in all samples when cells were subjected to combined treatment. It is also noteworthy that in every case when DNA internucleosomal fragments were easily seen, they were superimposed on DNA smears. This is probably caused by the occurrence of two parallel processes of DNA degradation when cells were more heavily damaged: unprogrammed (necrosis) and programmed (apoptosis) cell death. The technique of agarose gel electrophoresis, as applied in this study, does not permit a direct quantitative estimate of the levels of internucleosomal damage induced by single and combined treatment. From the selected drug doses and the results shown in Figure 5, however, it is evident that simultaneous employment of VP-16 with photodynamic treatment induces supra-additive increase of DNA fragmentation.

**Median effect – combination index analysis**

The results shown in previous sections indicate that drug combination treatment in most experimental conditions is supra-additive (synergistic). To evaluate modes of drug interactions, we designed experiments suitable for application of combination index analysis. This analysis is based on the median effect principle and is a statistical technique that allows formal evaluation of the nature of interaction between two cytotoxic agents. Cells were incubated with different concentrations of drugs, but at constant molar ratio (VP-16/AlPcS₄ = 1/2). Before irradiation, the cells were exposed to AlPcS₄ for 1.5 h, followed by co-incubation with VP-16 for 0.5 h. Thereafter, the cells were washed and irradiated (D = 7,2 J cm⁻²). Drug toxicity was assessed by their effect on clonogenicity (Figure 6). Drug toxicity was assessed by their effect on clonogenicity and clonogenicity. Figure 6 demonstrates that, under these conditions, the AlPcS₄ phototoxicity was low, as estimated by both growth rate inhibition and loss of clonogenicity. However, growth rate assessment of VP-16 toxicity underestimates the reproductive toxicity of the drug. Median effect plots derived from cloning experiments and linear regression data fit are shown in Figure 7. The dose–effect relationships of the individual drugs are...
strictly linear \( (r \geq 0.999, \text{ lines 1 and 2}) \), while the plot obtained for the mixture of the two drugs tends to concave upward \( (r = 0.99) \) and intersects the plot of the more active drug (Figure 7, line 3). It is, therefore, apparent that the combined drug action is synergistic. This was further confirmed by calculating the combination index values \( (CI) \) for a wide range of \( f_x \) ("fraction affected", Figure 8). Since the plots for the individual drugs are almost parallel (lines 1 and 2 in Figure 7), and the plot for the combined treatment (line 3) intersects the plot of the more active drug, it is likely that the two drugs interact as mutually non-exclusive. The combination index was calculated assuming both possibilities, mutually exclusive and non-exclusive interactions. The plots shown in Figure 8 indicate that these two models predict synergistic toxicity \( (CI \leq 1) \) under conditions in which combined treatment results in \( f_x \geq 0.10 \) or \( f_x \geq 0.15 \) respectively. For a comparison, \( CI \) values obtained from growth inhibition data were also calculated and included in Figure 8. It can be seen that synergism in the latter case is somewhat underestimated, probably due to the underestimation of VP-16 toxicity by this criterion.

Discussion

We have investigated the effects of individual toxicity of etoposide (VP-16) and photodynamic treatment with AlPcS4, as well as the efficacy of drug combination against K562 human leukaemic cells. Similarly, in our previous studies with Namalva Burkitt’s lymphoma cells (Gantchev et al., 1994b), photosensitisation of K562 cells results in division block and, depending on the extent of the treatment, is followed either by cell regrowth or by predominant cell death. Unlike photosensitisation with Photofrin, which induces S-phase cell cycle arrest (Gantchev et al., 1994d), in the present study we show that AlPcS4 photosensitisation arrests cells in G2/M-phase. Under the conditions of relatively mild photodynamic treatment employed, no evidence for immediate membrane disintegration was found. In contrast, most of the toxic effects were delayed and emerged as a function of post-treatment incubation time, as seen, for example, by the time course of the accumulation of G2/M-arrested cells and internucleosomal DNA cleavage. Similar cell growth inhibition effects were observed after short \( (0.5-2 \text{ h}) \) exposure of

Figure 5 Internucleosomal DNA fragmentation in K562 cells 4 h after treatment with drugs. Lanes: (1) control; (2 and 3) individual drug treatment with 5 \( \mu \text{M} \) and 10 \( \mu \text{M} \) AlPcS4 for 4 h; (4 and 5) 5 \( \mu \text{M} \) and 10 \( \mu \text{M} \) VP-16 for 1 h; (6 and 7) combination treatments: 5 \( \mu \text{M} \) AlPcS4 and 5 \( \mu \text{M} \) or 10 \( \mu \text{M} \) VP-16; (8 and 9) 10 \( \mu \text{M} \) AlPcS4 and 5 \( \mu \text{M} \) or 10 \( \mu \text{M} \) VP-16. Lane 10, marker DNA (1, 2 and 3 kb fragments). Photosensitisation light dose, \( (D) = 7.2 \text{ J cm}^{-2} \). DNA was isolated from 0.5 \( \times \text{10}^6 \) cells in each case.

Figure 6 Cell survival (loss of clonogenicity, open symbols) and growth rate inhibition (closed symbols) after photosensitisation with AlPcS4 \( (\bigtriangleup, \bigtriangleup) \), incubation with VP-16 \( (\bigcirc, \bullet) \) and combination treatment \( (\bigcirc, \bigtriangleup) \). Cells were incubated with various concentrations of AlPcS4 for 2 h and with VP-16 for 30 min. Drug concentration ratio VP-16/AlPcS4 = 1/2. Light dose, \( (D) = 7.2 \text{ J cm}^{-2} \). Combination treatment curves are plotted as a function of VP-16 concentration. Growth inhibition was estimated by comparison of growth rate constants. Averaged data from triplicate samples.

Figure 7 (1) Median-effect plots for the cytotoxicity of VP-16; (2) AlPcS4 and (3) their combination. Experimental data from clonogenic survival (Figure 6).
K562 cells to VP-16. The cytotoxicity of AlPcS₄-PDT, however, evolved faster compared with VP-16. In particular, PDT-induced growth and clonogenic inhibition developed at similar rates, while VP-16 displayed a lower level of growth suppression, compared with the more effective clonogenic inhibition (Figure 6). When applied alone, VP-16 did not induce any significant internucleosomal DNA degradation (apoptosis). Apoptotic cell death was an early response to AlPc₄-PDT and was augmented after combined drug treatment (Figure 5). Synergistic cell response to combined drug treatment was attained under various conditions, as clearly demonstrated by growth inhibition rates (Figure 1a and d), loss of clonogenicity (Figure 2) and accumulation of cells in G₂/M cell cycle phase (Figure 3b and c). In particular, synergism (supra-additivity) in cell cycle arrest took place when the fraction of G₂/M cells arrested by single drug treatment did not exceed a value of approximately 60%. Quantitatively, this value corresponds to the normal S-phase fraction of exponentially growing K562 cells and implies that drug treatment exclusively affects cells during the stage of DNA synthesis. After combined drug treatment and during the accumulation of cells in G₂/M-phase, a progressive increase of DNA internucleosomal fragmentation was observed. It is, therefore, likely that when cells are more extensively damaged by combined drug treatment, G₂/M-arrested cells tend to fall out of the cell cycle and die (eventually by apoptosis). The molecular mechanisms that are involved in AlPcS₄-PDT-induced apoptosis were not addressed in this study. However, we suggest that they do not encompass the phospholipases C and A₂, signalling pathway shown to operate during photosensitisation with cytoplasmic membrane-localising (unsubstituted) AlPc (Agarwal et al., 1993). In the latter case, the primary target is the cytoplasmic membrane, and it has been shown that DNA fragmentation evolves within minutes after treatment, in contrast to the slower accumulation of apoptotic cells after AlPcS₄ and/or VP-16 treatment. Water-soluble phthalocyanines, such as AlPcS₄, have been implicated in the induction of DNA strand breaks, DNA-protein cross-links (Hunting et al., 1987; Gantchev et al., 1994c), and impairment of cell organelles, including peroxisomes (Peng et al., 1991). All these oxidative processes may account for AlPcS₄ cytotoxicity and represent primary events in AlPcS₄ - PDT-mediated internucleosomal DNA cleavage.

The cytostatic/cytotoxic action of VP-16 is generally believed to involve formation of DNA-protein cross-links and breaks. However, it is likely that drug toxicity is not limited to topoisomerase II (topo II) poisoning only. Thus, a recent study argued on the lack of correlation between the level of DNA-protein cross-links and cytotoxicity in several leukemic cell lines (Dubrez et al., 1995). Previously, other studies have also indicated that the interaction of VP-16 with topo II and cytotoxicity of the drug can be uncoupled, e.g. drug interaction with topo II might be not intrinsically cytotoxic, and therefore it has been suggested that VP-16 cytotoxic action might involve one, or more, intervening metabolic steps (Kaufmann, 1989; Walker et al., 1991). Oxidoreductive transformations of VP-16 induced by a variety of intracellular enzymatic systems have been shown to constitute a pathway for modulation of drug activity (van Maanen et al., 1987; Mans et al., 1990; Haim et al., 1990). Furthermore, the intermediate species from VP-16 metabolism - VP-16 phenoxyl and semi-quinone radicals, and one of the final products of drug demethylation (VP-16 ortho-quinone) have been implicated as important cytotoxic agents (Mans et al., 1991, 1992; Ritov et al., 1995). It is, however, noteworthy that the intracellular accumulation of these products depends strongly on the presence of cellular antioxidants (e.g. reduced ascorbate and thiols) (Gantchev et al., 1994a; Kagan et al., 1995).

The combination index analysis of AlPcS₄ photosensitisation and VP-16 treatment as performed in the present work largely predicts synergistic toxicity of the two drugs. The experiments throughout this study were performed so that effects, such as cell synchronisation, were avoided and cannot account for the observed synergy. Moreover, the median effect plot of combined toxicity (Figure 7) suggests that the drugs are likely to interact as mutually non-exclusive. Non-exclusivity of drug interaction would imply that AlPcS₄ and VP-16 share a common intracellular target and/or interact chemically. It is not known if AlPcS₄ photosensitisation can directly interfere with the topo II/DNA breakage-reunion process, but it is recognised that photosensitisation (Hunting et al., 1987; Gantchev et al., 1994c) and VP-16 metabolites (van Maanen et al., 1988b; Mans et al., 1990, 1991; Sinha et al., 1990) can instantly induce damage to DNA. Therefore, based on the assumption that metabolic oxidoreductive transformations of VP-16 are important for its cytotoxic action, we suggest that photosensitisation-induced depletion of intracellular reductants facilitates VP-16 metabolism, probably on the level of VP-16 phenoxyl free radical transformations. Generation of the VP-16 phenoxyl radical is the primary step in the enzymatic demethylation of the compound and the interactions of the phenoxyl radical with cell antioxidants prevents etoposide from further transformations and results in recovery of its original form. Also, the VP-16 phenoxyl radical may itself initiate oxidative cell damage (Ritov et al., 1995). A parallel drug interaction mechanism may involve direct AlPcS₄-mediated photooxidation of VP-16 to yield the phenoxyl radical (Gantchev et al., 1994a). Complete elucidation of the role of these different pathways and the involvement of additional processes that may explain the observed synergistic drug toxicity will require further experimental work. It is relevant to note that a recent study showed synergism in cells subjected to gamma radiation and etoposide treatment (Haddock et al., 1995). Since the oxidative processes induced
by PDT and radiotherapy are similar, it is possible that the synergistic interactions with etoposide are based on the same mechanisms. Although the presented results only involve in vitro synergy between photodynamic treatment and etoposide toxicity, our findings may provide alternative perspectives for clinical PDT and etoposide applications.

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