Mitochondria-targeting drugs arsenic trioxide and lonidamine bypass the resistance of TPA-differentiated leukemic cells to apoptosis

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Exposure of U937 human leukemic cells to the phorbol ester 12-O-tetradecanoylphorbol 13-acetate (TPA) induces their differentiation into monocyte/macrophage-like cells. This terminal differentiation is associated with a resistant phenotype to apoptosis induced by the topoisomerase II inhibitor etoposide. The inhibition occurs upstream of the mitochondrial release of cytochrome c and the activation of procaspase-2, -3, -6, -7, -8, and -9. By using cell-free systems, it was demonstrated that the mitochondrial pathway to cell death involves mitochondrial membrane depolarization, cytochrome c release and cytosolic activation of procaspases by cytochrome c/dATP remains functional in TPA-differentiated U937 cells. Accordingly, 2 drugs recently shown to target the mitochondria, namely lonidamine and arsenic trioxide, bypass the resistance of TPA-differentiated U937 cells to classical anticancer drugs. Cell death induced by the 2 compounds is associated with mitochondrial membrane depolarization, release of cytochrome c and Smac/Diablo from the mitochondria, activation of caspases, poly(ADP-ribose) polymerase cleavage and internucleosomal DNA fragmentation. Moreover, the decreased glutathione content associated with the differentiation process amplifies the ability of arsenic trioxide to activate the mitochondrial pathway to cell death. Similar results were obtained by comparing undifferentiated and TPA-differentiated human HL60 leukemic cells. These data demonstrate that mitochondria-targeting agents bypass the resistance to classical anticancer drugs induced by TPA-mediated leukemic cell differentiation. (Blood. 2001;97:3931-3940)

Introduction

Chemotherapeutic drugs can kill cultured leukemic cell lines by inducing apoptosis. This mode of cell death may be clinically relevant because apoptotic blast cells have been detected in the peripheral blood of patients with acute leukemia who received chemotherapeutic drugs. Hallmarks of apoptosis in leukemic cells include characteristic morphologic changes such as cell shrinkage, membrane blebbing, and cell fragmentation into apoptotic bodies, internucleosomal DNA fragmentation, and limited proteolytic cleavage of selective intracellular proteins such as the nuclear enzyme poly(ADP-ribose)polymerase (PARP). This cell death phenotype is induced by activation of a constitutively expressed apoptotic machinery that involves a family of cysteine proteases known as caspases, organized in a branched proteolytic cascade. These enzymes are synthesized as inactive proenzymes (procaspases) whose proteolysis at internal aspartate residues generates large and small subunits. Heterodimerization of these subunits is required to form the active enzyme (caspase) that cleaves intracellular substrates, either a downstream procaspase or other cellular proteins.

Activation of the caspase cascade in response to specific damage induced by chemotherapeutic drugs is the consequence of a disruption of the mitochondrial membrane barrier function. Although the exact mechanism remains controversial, mitochondrial changes lead to the release of soluble apoptogenic proteins such as cytochrome c, apoptosis-inducing factor (AIF), and Smac/Diablo from the mitochondrial intermembrane space to the cytosol. Once in the cytosol, cytochrome c, in the presence of adenosine triphosphate (ATP), induces conformational changes of an adaptor molecule designated Apaf-1 that recruits and activates the caspase-9 molecules. In turn, caspase-9 activates the downstream caspase cascade that involves caspase-3 and other effector enzymes. Smac/Diablo was recently shown to suppress the inhibition of caspases by proteins known as inhibitor of apoptosis proteins (IAPs), thereby increasing their sensitivity to the cytochrome c/ATP pathway of activation. Once activated, effector caspases cleave a limited set of essential cellular proteins, leading to cell dismantling. Mitochondrial function during apoptosis is controlled by the Bcl-2 family of proteins that could prevent the opening of the permeability transition pore or stabilize the barrier function of the outer mitochondrial membrane and prevent the release of apoptogenic molecules.

How the specific damage induced by chemotherapeutic drugs leads to mitochondrial changes and activation of the caspase cascade under the control of Bcl-2–related proteins remains poorly understood. We have shown previously that differentiation of human myeloid leukemia cell lines along a macrophagic pathway induced by exposure to the phorbol ester 12-O-tetradecanoylphorbol 13-acetate (TPA) correlated with a resistance phenotype to apoptosis induced by a series of anticancer drugs with distinct primary intracellular targets. This inhibition occurs downstream of specific damage, measured as cleavage complexes,
induced by topoisomerase poisons. In addition, TPA-differentiated cells are resistant to apoptosis induced by exposure to Fas-ligand, a specific pathway that was proposed as an amplifier system in drug-induced cell death. A resistance to drug-induced and TRAIL-mediated apoptosis was also reported in vitamin D$_3$- and dimethyl sulfoxide (DMSO)-differentiated HL60 cells, respectively.

Identification of the molecular mechanisms that modulate cell sensitivity to chemotherapeutic drugs in differentiated cells could help to define strategies combining differentiating and cytotoxic agents for treating acute leukemias. The present study further explores the level of inhibition of drug-induced apoptosis in TPA-differentiated cells. By using cell-free systems, we demonstrate that the pathway involving mitochondria and the downstream caspase cascade remains functional in terminally differentiated cells. Based on this observation, we tested the ability of 2 chemotherapeutic drugs recently shown to target the mitochondria, namely the indazole-3-carboxylic acid derivative lonidamine and the metalloid arsenic trioxide, to induce apoptosis in these cells.

DNA fragmentation

DNA fragmentation was quantified by using a previously reported filter elution assay. Briefly, exponentially growing cells were incubated with 0.02 μCi/mL [3H]-thymidine and cultured at 37°C for 2 days. Then, cells were chased in isotope-free medium overnight, resuspended in fresh medium, and treated with different agents before or after TPA treatment. Approximately 1.0 × 10$^6$ treated or untreated labeled cells were loaded onto a protein-absorbing filter (polyvinylidene difluoride filters, 0.65 μm pore size, 25 mm diameter; Durapore membrane, Millipore, Saint-Quentin en Yvelines, France). Cells were then washed once with ice-cold phosphate-buffered saline (PBS) and subsequently lysed in 0.2% sodium sarkosyl, 2 M NaCl, 0.04 M EDTA, pH 10.0. Filters were washed with 0.02 M EDTA, pH 10.0. DNA was depurinated by incubation of filters in 1 M HCl at 65°C for 45 minutes, then released from the filters with 0.4 M NaOH for 45 minutes at room temperature. Radioactivity was counted by liquid scintillation spectrometry in each fraction (wash, lysis, EDTA wash, and filter). DNA fragmentation was measured as the fraction of disintegrations per minute (dpm) in the lysis fraction plus EDTA wash relative to the total intracellular dpm. For analyzing DNA fragmentation by agarose gel electrophoresis, cellular DNA was extracted by a previously described salting-out procedure, and electrophoresis was performed in 1.8% agarose gel in Tris-borate–EDTA buffer (pH 8.0) at 20 V for 14 hours. After electrophoresis, DNA was visualized by ethidium bromide staining.

Western blot analysis

After treatment, whole cell lysates were prepared by lysing the cells in boiling buffer (1% SDS, 1 mM Na-β-mercaptoethanol, 100 mM Tris pH 7.4) in the presence of protease inhibitors (0.1 mM phenylmethanesulfonyl fluoride (PMSF, 2.5 μg/mL pepstatin, 10 μg/mL aprotinin, 5 μg/mL leupeptin). Viscosity of the samples was reduced by several passages through a 27-gauge needle. Protein concentration was measured by the Bio-Rad DC protein assay kit (Ivy sur Seine, France). Thirty micrograms protein was incubated in loading buffer (125 mM Tris-HCl, pH 6.8, 10% β-mercaptoethanol, 4.6% SDS, 20% glycerol, and 0.003% bromophenol blue), separated by sodium dodecyl sulfate–polyacrylamide gel (SDS-PAGE), and electroblotted to polyvinylidene difluoride membrane (Bio-Rad). After blocking nonspecific binding sites overnight by 5% nonfat milk in TBBS (0.1% PBS-Tween 20), the membrane was incubated for 2 hours at room temperature with primary antibody. After 2 washes in TBBS, the membrane was incubated with horseradish peroxidase–conjugated goat antimouse or antirabbit antibodies (Jackson ImmunoResearch Laboratories, West Grove, PA) for 30 minutes at room temperature, then washed twice in TBBS. Immuno blotting was revealed using enhanced chemiluminescence detection kit (Amersham) by autoradiography.

Cell fractionation

Cytoxic fractions were prepared by resuspending the cells in ice-cold buffer A (250 mM sucrose, 20 mM HEPES, 10 mM KCl, 1.5 mM MgCl$_2$, 1 mM EDTA, 1 mM EGTA, 1 mM dithiothreitol, 17 μg/mL PMSF, 8 μg/mL aprotinin, 2 μg/mL leupeptin, [pH 7.4]) before passing them through an ice-cold cell homogenizer. Unlysed cells and nuclei were pelleted by a 10-minute, 750g spin. This step was repeated twice. The supernatant was centrifuged at 10 000g for 25 minutes, and the resultant supernatant was further centrifuged at 100 000g for 1 hour. The supernatant (cytosolic fraction) was frozen at ~80°C. Nuclei-free, mitochondria-free cytosolic extracts (cell-free extracts) were generated as previously described. Briefly, cells (1–2 × 10$^9$) were pelleted and washed twice with PBS, pH 7.2, followed by a single wash with 4 mL ice-cold cell extract buffer (20 mM HEPES, 10 mM KCl, 1.5 mM MgCl$_2$, 1 mM EDTA, 1 mM EGTA, 1 mM dithiothreitol, 100 μM PMSF, 2 μg/mL aprotinin, 10 μg/mL leupeptin [pH 7.4]). Two volumes ice-cold extract buffer were added to 1 vol packed cell pellet before the cellular suspension was transferred to a 2-mL Dounce homogenizer. Cells were allowed to swell under the hypotonic condition for 20 minutes, then were disrupted with 30 strokes of a B-type. Lysis was confirmed with the trypsin blue dye exclusion test before centrifugation of lysates at 16 000g for 15 minutes at 4°C. Caspase activation by the cytochrome c/dATP combination was tested in these clarified supernatants.

Materials and methods

Drugs and chemical reagents

Etoposide (VP-16) and TPA were obtained from Sigma-Aldrich Laboratories (St Quentin Fallavier, France). Lonidamine was kindly provided by Dr M. F. Poupon (Institut Curie, Paris, France). Stock solutions were prepared by diluting these reagents in DMSO and storing them at −20°C. Further dilutions were made in culture medium just before use. The final concentration of DMSO in culture medium never exceeded 0.1% (vol/vol), which was nontoxic to the cells. Arsenic trioxide was kindly provided by Dr G. Q. Chen (Shanghai Institute of Hematology, China). [2-14C]-Thymidine (56 mCi/mM) was obtained from Amersham (Les Ulis, France). All other chemicals were of reagent grade and were purchased from local sources.

Antibodies

The tested mouse monoclonal antibodies (mAbs) included antihuman HSC70 from Santa Cruz Biotechnology (Santa Cruz, CA), cytochrome c, caspase-2, caspase-6, and caspase-8 from Pharmingen (San Diego, CA), caspase-3 and caspase-7 from Transduction Laboratories (Lexington, KY), and cytochrome oxidase subunit IV (COX IV) from Molecular Probes (Eugene, OR). We also used the polyclonal rabbit antibodies raised against human caspase-3-p17 (kindly provided by Donald W. Nicholson, MerkFrost, Toronto, ON, Canada), caspase-9 (Pharmingen), caspase-7 (StressGen, Victoria, BC, Canada), PARP (Boehringer-Mannheim, Germany), and Smac/Diablo (kindly provided by Xiaodong Wang, Howard Hughes Medical Institute, Dallas, TX).

Cell culture and differentiation

The human leukemic cell lines U937 and HL60 (American Type Culture Collection, Rockville, MD) were grown in suspension in RPMI 1640 medium with glutamax-I (Gibco BRL, Life Technologies, Cergy Pontoise, France) supplemented with 10% (vol/vol) fetal bovine serum (Bios Whitaker, Fontenay-sous-Bois, France) in an atmosphere of 95% air and 5% CO$_2$ at 37°C. Cell viability was determined by using a trypan blue dye exclusion assay. To ensure exponential growth, cells were resuspended at a density of 0.5 × 10$^6$ cells/mL in fresh medium 24 hours before each treatment. To induce differentiation, cells were culture in the presence of 20 μM all-trans retinoic acid (t-RA) and dimethyl sulfoxide (DMSO)–differentiated HL60 cells, respectively.

DNA fragmentation

DNA fragmentation was quantified by using a previously reported filter elution assay. Briefly, exponentially growing cells were incubated with 0.02 μCi/mL [3H]-thymidine and cultured at 37°C for 2 days. Then, cells

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Purification of mitochondria

Purified mitochondria were isolated as previously described. Briefly, cells (1 × 10^6) were resuspended in buffer H (300 mM sucrose, 5 mM N-tris(hydroxymethyl)methyl-2-amino-ethanesulfonic acid, 200 mM EGTA, pH 6.9), homogenized in a Potter-Thomas homogenizer, then centrifuged for 10 minutes at 760 g. The supernatant was recovered while the pellet was resuspended in H buffer and centrifuged to recover the supernatant as above. Both supernatants were mixed and centrifuged for 10 minutes at 8740 g before the mitochondria pellet was resuspended in buffer H.

Activation of cell-free apoptosis

Cell-free extracts (10 µL, 5-10 mg/mL protein) were incubated at 37°C with 5 µM horse heart cytochrome c (Sigma) and 1 mM dATP (Pharmacia). Then caspase activation was determined by Western blot analysis. For cell-free reaction activated by actrylsode, purified mitochondria were exposed to 5 mM atractyloside for 30 minutes and pelleted by a 10 000g spin for 10 minutes before cytochrome c release was determined by Western blot in both the supernatant and the mitochondria pellet.

Analysis of mitochondrial membrane potential

Mitochondrial membrane depolarization was analyzed using the DePsipher kit (R&D Systems, Abington, United Kingdom) according to the manufacturer’s instructions. Briefly, 10^6 cells were incubated for 20 minutes at 37°C in the presence of 5 µM/µL DepSipher (R&D Systems) solution and then washed 2 times in PBS. In some cases, cells were treated for 30 minutes with the uncoupling agent carbonyl cyanide m-chlorophenyldihydrazone (100 µM; Sigma-Aldrich). Analysis was performed by the use of a FACScan cytometer (Becton Dickinson, Le Pont de Claix, France).

Quantification of intracellular glutathione and reactive oxygen species content by flow cytometry

The level of cellular glutathione (GSH) was determined by flow cytometry after staining with monochlorobimane (Molecular Probes) as previously described. Briefly, 200 µM monochlorobimane [(syn-(CICH2, CH3)-1,5-diazabicyclo-[3.3.0]-octa-3,6-dione-2,8-dione)] was added to cell suspensions for 30 minutes at 37°C. After 2 washes in PBS, cells were resuspended in PBS and analyzed on a Bio-Rad flow cytometer (Hercules, CA). For reactive oxygen species (ROS) measurements, cells were incubated for 15 minutes at 37°C in the presence of 6.6 µM dihydro-ethidium (Sigma-Aldrich) and analyzed by flow cytometry.

Results

TPA-induced differentiation of U937 human leukemic cells prevents etoposide-induced apoptosis

U937 leukemic cells were induced to differentiate along a macrophagic pathway on exposure to 20 nM TPA. After a 48- to 72-hour exposure to TPA, more than 90% cells demonstrate a differentiated phenotype including adhesion to the culture flask and increased expression of the glycoprotein CD11b at the cell surface. On continuous exposure to 50 µM etoposide, most of the parental U937 cells undergo apoptosis in 6 hours, as demonstrated by staining nuclear chromat in with Hoechst 33342, whereas TPA-differentiated cells strongly resist drug-induced cell death (Figure 1).

To link the resistance phenotype to the differentiation process, we exposed the cells for 4.5 hours to 50 µM etoposide at various times after the beginning of TPA treatment (20 nM) (Figure 2). Apoptosis was identified by Hoechst staining of nuclear chromat in (Figure 2A) and quantified by measuring nuclear DNA fragmentation (Figure 2B). Exposure to TPA provoked a time-dependent decrease of etoposide-induced morphologic changes and DNA fragmentation that were almost completely inhibited 48 to 72 hours after the beginning of TPA exposure. Accordingly, by using agarose gel electrophoresis, the characteristic nucleosome-sized ladder fragmentation of nuclear DNA observed in parental undifferentiated cells exposed for 4.5 hours to the drug was not detected in TPA-differentiated cells treated in the same conditions (Figure 2B, insert). Etoposide treatment of parental cells also induced a decrease of procaspase-2, -3, -6, -7, -8, and -9 expression, as determined by Western blot analysis (Figure 2C). By using antibodies reacting with the cleavage fragments that characterize active enzymes, decreased expression of the 48-kd procaspase-2, the 32-kd procaspase-3, the 35-kd procaspase-7, and the 48-kd procaspase-9 correlated with the appearance of a p35, a p19/p17, a p10, and a p37 active fragment, respectively. The 30-kd fragment identified with the anti-caspase-9 antibody might correspond to the previously described caspase-9b isoform. The significance of the 18-kd band remains unknown. All these etoposide-induced events were progressively inhibited as U937 cells progressed toward the differentiation phenotype on TPA exposure. This inhibition became identifiable between 6 and 12 hours after the beginning of TPA exposure, depending on the experiment. In addition, TPA-induced differentiation was associated with a progressive inhibition of the 116-kd DNA repair enzyme PARP cleavage into an 85-kd N-terminal fragment (Figure 2C).

Caspases remain able to be activated in cell-free extracts from TPA-induced differentiated cells

Exposure of undifferentiated U937 cells to 50 µM etoposide for 4.5 hours triggered the appearance of cytochrome c in the cytosol. This event was prevented by TPA-induced differentiation (Figure 3A). To determine whether the cytochrome c–mediated activation of the caspase cascade was influenced by the differentiation process, we used a previously described cell-free system. Exposure of cell-free extracts from undifferentiated U937 cells to 5 µM cytochrome c and 1 mM dATP induced a time-dependent decrease of the expression of procaspase-2, -3, -6, -7, -8, and -9, suggesting their cleavage into active fragments. Accordingly, immunoblot experiments detected the time-dependent appearance of the 35-kd active fragment of caspase-2, the 19- and 17-kd fragments of caspase-3, the 11-kd fragment of caspase-6, the 30-kd fragment of caspase-7, and the 37-kd fragment of caspase-9 (Figure 3B). In this time-course, procaspase-8 expression remained unchanged. TPA-mediated differentiation of U937 cells induced some changes in the
subcellular localization of procaspases, mainly the disappearance of procaspase-2 from the cytosol (O.S. et al, manuscript in preparation). The other procaspases remained sensitive to activation by the cytochrome c–dATP combination with kinetics similar to that observed in cytosolic extracts from undifferentiated cells (Figure 3B). Thus, TPA-induced differentiation does not significantly influence procaspase activation by cytochrome c/dATP.

Mitochondria from TPA-induced differentiated U937 cells remain reactive

The mechanisms that account for the mitochondrial release of cytochrome c are still controversial. One of the proposed mechanisms involves a decrease of the mitochondrial membrane potential ∆ψm.5 To determine whether TPA-induced differentiation was associated with a decreased reactivity of mitochondria to uncoupling agents, we exposed undifferentiated and TPA-differentiated U937 cells for 30 minutes to 100 μM of the uncoupling agent carbonyl cyanide m-chlorophenylhydrazone (mCICCP). This treatment induced a switch from the mitochondrial aggregate (FL2-H) to the cytosolic monomer (FL1-H) that was similar in mitochondria from both undifferentiated and TPA-differentiated cells indicating mitochondrial depolarization (Figure 4A). A mechanism proposed to account for the mitochondrial release of cytochrome c was the opening of the megachannel that can be induced by exposing purified mitochondria to atractyloside (5 mM, 30 minutes). This compound induced the release of cytochrome c from mitochondria of undifferentiated and differentiated U937 cells with a similar efficacy (Figure 4B). In the presence of cell-free extracts from undifferentiated U937 cells, the effect of atractyloside on purified mitochondria resulted in cleavage of the DEVD-AFC substrate, suggesting caspase activation (data not shown). Together these results suggested that the mitochondrial pathway to cell death may remain functional in TPA-differentiated U937 cells.

Mitochondria-targeting agent lonidamine induced apoptosis of TPA-differentiated cells

The previous observations suggested that drugs shown to directly target the mitochondria might overcome the resistance of TPA-differentiated cells to etoposide and various other classical antican- cer drugs.18 We first tested the mitochondria-targeting drug lonidamine.27 The ability of this compound to induce apoptosis of TPA-differentiated cells was only slightly delayed in TPA-differentiated compared to undifferentiated U937 cells, as demonstrated by quantifying DNA fragmentation (Figure 5A) and by counting the percentage of cells demonstrating nuclear chromatin condensation after a 24- to 48-hour exposure to 200 μM lonidamine (Table 1). Lonidamine-induced apoptotic DNA fragmentation and chromatin condensation were completely inhibited by stable overexpression of Bcl-2 in U937 cells (Figure 5A; Table 1). In undifferentiated and TPA-differentiated cells, lonidamine induced mitochondrial depolarization (Figure 5C), cytochrome c release in the cytosol (Figure 5B), procaspase-3 cleavage in its active form.
fragments, and PARP cleavage (Figure 5D). All these events were prevented by Bcl-2 overexpression. The recently described Smac/Diablo protein was only slightly released in the cytosol on lonidamine exposure, an effect that was not prevented by Bcl-2 overexpression (Figure 5B).

TPA-induced differentiation sensitizes U937 cells to arsenic trioxide–induced apoptosis

As$_2$O$_3$ is another member of this novel class of chemotherapeutic agents that induce apoptosis, at least in part, by acting on the mitochondria. As$_2$O$_3$ induced apoptosis of undifferentiated U937 cells in a dose- and time-dependent manner (Figure 6A; Table 1). TPA-induced differentiation did not prevent, and even sensitized, U937 cells to As$_2$O$_3$-induced apoptosis. The differentiation process also increased the ability of As$_2$O$_3$ to induce mitochondrial polarization (Figure 6C), cytochrome c release from the mitochondria to the cytosol (Figure 6B), procaspase-3 cleavage in its active fragments and PARP cleavage (Figure 6D). All these events were prevented by Bcl-2 overexpression in U937 cells (Figure 6A-D; Table 1). Smac/Diablo protein was released in the cytosol on As$_2$O$_3$ treatment more efficiently than on lonidamine exposure. Again, this effect was not prevented by Bcl-2 overexpression (Figure 6B). Because As$_2$O$_3$-induced apoptosis has been shown to be modulated by the cellular GSH redox system, with increased intracellular levels of reduced GSH having an inhibitory effect, we measured the influence of TPA-induced differentiation on cellular GSH and ROS content in untreated and As$_2$O$_3$-treated cells. By using flow cytometry methods, we observed that the differentiation process was associated with a decrease in the basal level of GSH (Figure 7A) and an increase in the basal ROS content in the cells (Figure 7B). Arsenic exposure decreased GSH content and increased ROS level in parental cells. These effects were prevented by Bcl-2 overexpression. In differentiated cells, As$_2$O$_3$ treatment further decreased GSH content without increasing the cellular ROS content, as measured by using dihydro-ethidium (Figure 7A-B).

Lonidamine and arsenic trioxide also bypass the resistance of TPA-differentiated HL60 cells to classical chemotherapeutic drugs

The results obtained in U937 cells were reproduced in HL60 cells whose exposure to TPA also induces a macrophagic-like differentiation process with a resistant phenotype to classical anticancer drugs. TPA-differentiated HL60 cells, which were resistant to etoposide-induced apoptosis, demonstrated a sensitivity to lonidamine that was only slightly delayed compared to that of parental cells, whereas their sensitivity to As$_2$O$_3$-induced cell death was greatly increased (Figure 8; Table 1).
Discussion

Classical chemotherapeutic drugs induce apoptosis by compromising mitochondrial function in an indirect fashion—for example, by inducing DNA damage that triggers p53 activation and interferes with cell cycle progression by stimulating the Fas-dependent pathway, or by triggering ceramide production, and by inducing shifts in redox potentials. A series of new chemotherapeutic drugs, many of them currently tested in clinical trials, have been shown to act, at least in part, by directly targeting the mitochondria, as demonstrated by their ability to activate the permeability transition pore complex and to trigger the release of soluble intermembrane proteins from purified mitochondria in vitro. This observation led to the theoretical suggestion that these compounds may bypass the resistance to anticancer drugs induced by alterations of indirect death-inducing pathways. The current study demonstrates that TPA-mediated differentiation of leukemic cells reproduces a situation in which the pathway to cell death, triggered by a classical anticancer drug such as etoposide, is inhibited upstream of the mitochondrial events. Analysis of this model system confirms the hypothesis that mitochondria-targeting drugs can overcome such a resistant phenotype.

TPA-mediated inhibition of etoposide-induced apoptosis increased as leukemic cells progressed toward the macrophagic differentiation phenotype. Short-term effects of TPA are mainly the activation of several protein kinase C (PKC) isoenzymes. The differentiation process involves several events downstream of the transient activation of PKCs. Thus, the differentiation-related resistance of leukemic cells to etoposide-induced cell death might be distinguished from the direct consequences of PKC activation. A differentiation-related resistance to apoptosis induced by various stimuli has been described in leukemic cells exposed to either 1,25-dihydroxyvitamin D3 or DMSO. A resistance to growth factor deprivation-induced cell death was also identified in differentiated myocytes. Thus, several differentiation processes in different cell types were associated with a decreased sensitivity to apoptosis induced by a series of distinct stimuli.

Most pathways to apoptosis converge on the mitochondrial release of apoptogenic molecules to the cytosol and the activation of the caspase cascade. The current study confirms that etoposide activates a variety of caspases while inducing apoptosis of

Table 1. Effects of phorbol ester 12-O-tetradecanoylphorbol 13-acetate (TPA)-induced differentiation and Bcl-2 overexpression on etoposide, lonidamine, and arsenic trioxide–induced apoptosis in U937 and HL60 human leukemic cells

<table>
<thead>
<tr>
<th>Condition</th>
<th>U937 (mean SD)</th>
<th>U-TPA (mean SD)</th>
<th>U-BCL2 (mean SD)</th>
<th>HL60 (mean SD)</th>
<th>HL-TPA (mean SD)</th>
<th>HL-BCL2 (mean SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>2.8 ± 1.6</td>
<td>1.0 ± 0.6</td>
<td>0.8 ± 0.7</td>
<td>0.8 ± 0.7</td>
<td>1.3 ± 1.2</td>
<td>0.3 ± 0.7</td>
</tr>
<tr>
<td>VP16 6 h</td>
<td>71.7 ± 6.7</td>
<td>4.3 ± 2.1</td>
<td>—</td>
<td>46.3 ± 1.2</td>
<td>2.3 ± 2.6</td>
<td>—</td>
</tr>
<tr>
<td>LND 24 h</td>
<td>94.0 ± 16.0</td>
<td>22.3 ± 3.1</td>
<td>—</td>
<td>96.7 ± 1.2</td>
<td>9.5 ± 0.9</td>
<td>—</td>
</tr>
<tr>
<td>LND 48 h</td>
<td>39.3 ± 3.5</td>
<td>16.3 ± 3.3</td>
<td>2.8 ± 1.1</td>
<td>79.7 ± 4.1</td>
<td>25.2 ± 5.2</td>
<td>3.3 ± 1.2</td>
</tr>
<tr>
<td>As2O3 24 h</td>
<td>93.3 ± 2.7</td>
<td>81.3 ± 5.5</td>
<td>4.5 ± 1.4</td>
<td>98.7 ± 0.7</td>
<td>67.5 ± 5.4</td>
<td>13.2 ± 2.6</td>
</tr>
<tr>
<td>As2O3 48 h</td>
<td>12.7 ± 2.1</td>
<td>26.8 ± 5.8</td>
<td>2.3 ± 2.0</td>
<td>8.0 ± 1.6</td>
<td>20.3 ± 3.7</td>
<td>0.3 ± 0.5</td>
</tr>
</tbody>
</table>

Undifferentiated (U937, HL60). TPA-differentiated (U-TPA, HL-TPA), and Bcl-2 overexpressing (U-BCL2, HL-BCL2) U937 and HL60 cells were left untreated (control) or treated for indicated times with 50 μM etoposide (VP16), 200 μM lonidamine (LND), or 4 μM arsenic trioxide (As2O3). Results are expressed as percentages of apoptotic cells (mean ± SD of triplicate determinations).
undifferentiated U937 cells. Various mechanisms negatively interfere with this final pathway, including overexpression of antiapoptotic proteins of the Bcl-2 family that prevents the cytosolic release of mitochondrial soluble molecules and overexpression of heat-shock proteins such as HSP27 and HSP70 that prevent cytochrome c–mediated activation of the caspase cascade by interacting with cytochrome c and the Apaf-1 adaptor molecule, respectively. Despite the fact that the TPA-induced differentiation process induces some changes in the subcellular localization of procaspases (O.S. et al, manuscript in preparation) and modulates the expression of inhibitory proteins—for example, by inducing transient changes in the expression of the Bcl-2–related protein Mcl-1 and the Bcl-2 overexpressing U937 cells (U-BCL2) were treated with increasing concentrations of As$_2$O$_3$ for 72 hours (left panel) or with 4 μM As$_2$O$_3$ for indicated times (right panel). Apoptotic DNA fragmentation was quantified with the use of a filter elution assay. Values from 1 of 2 independent experiments are shown (mean ± SD of triplicate samples). (B) Western blot analysis of cytochrome c and Smac/Diablo expression in cytosolic fractions from undifferentiated (U937), and TPA-differentiated (U-TPA) U937 cells and U937 cells overexpressing Bcl-2 (U-BCL2) treated with 4 μM As$_2$O$_3$ for indicated times. (C) Mitochondrial membrane depolarization was visualized by flow cytometry in U937, U-TPA, and U-BCL2 cells treated with 4 μM As$_2$O$_3$ for indicated times. Mitochondrial depolarization was identified by an increased level of cytosolic green monomer (FL1-H; dashed lines indicate the controls). (D) Western blot analysis of procaspase-3 and caspase-3 active fragments and PARP expression in whole-cell extracts from U937, U-TPA, and U-BCL2 cells treated with 4 μM As$_2$O$_3$ for indicated times. Numbers are molecular weights in kilodaltons. *Cleavage products. Loading was checked with the use of an antihuman HSC70 mAb.

Lonidamine (1-[(2,4-dichlorophenylmethyl) methyl]-1H-indazole-3-carboxylic acid) is an antineoplastic drug derived from indazole-3-carboxylic acid. This compound, which was shown to exacerbate the response of human tumor cells to cisplatin, irradiation, doxorubicin, or cyclophosphamide, has been tested in phase II and III trials of metastatic breast cancer and ovarian cancer. Although several distinct mechanisms were proposed to account for lonidamine cytotoxic effects, recent data suggest that mitochondria could be the main subcellular target of this compound. Lonidamine directly acts on the permeability transition pore complex when tested on purified organelles, an effect inhibited by Bcl-2. Since we have shown that the mitochondrial pathway to cell death remains unchanged in TPA-differentiated cells, lonidamine may have demonstrated a similar efficacy on undifferentiated and TPA-differentiated leukemic cells. Actually, TPA-induced differentiation slightly, but reproducibly, delays lonidamine-induced apoptosis in U937 and HL60 cells, suggesting a negative effect of the differentiation process on some other potential consequences of lonidamine treatment, such as cytosolic calcium increase and intracellular lactate accumulation. This effect of the differentiation process on lonidamine-induced cell death remains limited compared with the strong inhibition of apoptosis triggered by classical anticancer drugs.

As$_2$O$_3$ is another therapeutic compound that was suggested to

![Figure 6. TPA-induced differentiation sensitizes U937 cells to arsenic trioxide (As$_2$O$_3$)-induced differentiation.](image)

![Figure 7. Intracellular GST and ROS levels.](image)
directly target the mitochondria. Although As$_2$O$_3$-induced apoptosis was initially described in acute promyelocytic cell lines, this compound induces the death of many other leukemia cell types. The hypothesis that As$_2$O$_3$ directly targets the mitochondria is based on the observation that exposure of purified organelles to arsenic triggers opening of the permeability transition pore and the release of soluble intermembrane proteins. In whole cells, As$_2$O$_3$ induces disruption of the mitochondrial transmembrane potential and caspase activation, events that are inhibited by Bcl-2 overexpression. As$_2$O$_3$-induced apoptosis was also related to Bcl-2 down-regulation, suppression of the apoptosis inhibitory Ras/MAP kinase cascade, and enhanced translocation of PML or PML/RARα chimeric proteins to nuclear bodies, but these events depend on As$_2$O$_3$ dose, cell type, and cellular environment. We show here that TPA-induced differentiation of leukemic cells increases the ability of As$_2$O$_3$ to target the mitochondria, to induce the release of cytochrome c and Smac/Diablo, to activate downstream caspases, and to induce apoptotic cell death. A similar synergy was described between As$_2$O$_3$ and retinoic acid, another differentiating agent, in an arsenic-resistant NB4 subline. The differentiation-related decrease in GSH cellular content may contribute to the sensitization of leukemic cells to As$_2$O$_3$-mediated cell death.

In conclusion, we have shown that the TPA-mediated differentiation of human myeloid leukemic cell lines induces a resistance phenotype to classical anticancer drugs such as etoposide without inhibiting the mitochondrial pathway to cell death. The differentiation process has limited influence on cell death induced by the mitochondria-targeting agent lonidamine and even sensitizes the cells to arsenic-mediated activation of the mitochondrial pathway.

Figure 8. Lonidamine and arsenic trioxide bypass the resistance of TPA-differentiated HL60 cells to etoposide. Undifferentiated (A, HL60), TPA-differentiated (B, HL-TPA), and Bcl-2 overexpressing (C, HL-BCL2) HL60 cells were treated with 50 μM etoposide (VP16), 200 μM lonidamine (LND), or 4 μM arsenic trioxide (As$_2$O$_3$) for indicated times. (A) Apoptotic DNA fragmentation was quantified with the use of a filter elution assay. Values from 1 of 2 independent experiments are shown (mean ± SD of triplicate samples). (B) Western blot analysis of PARP cleavage in whole-cell extracts from HL60, HL-TPA, and HL-BCL2 cells, either untreated (−) or treated (+) with 50 μM VP16 for 6 hours, 200 μM LND for 48 hours, or 4 μM As$_2$O$_3$ for 48 hours. Numbers are molecular weights in kilodaltons. *Cleavage product. Loading was checked with the use of an anti-human HSC70 mAb.

Figure 9. Influence of TPA-induced differentiation on drug-induced cell death. TPA-mediated differentiation of leukemic cells strongly decreases the ability of classical anticancer drugs, such as those inducing DNA damage, to activate the mitochondrial pathway to cell death. In contrast, the differentiation process has limited influence on cell death induced by the mitochondria-targeting agent lonidamine and even sensitizes the cells to arsenic-mediated activation of the mitochondrial pathway.

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References


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