Cisplatin Preferentially Binds Mitochondrial DNA and Voltage-Dependent Anion Channel Protein in the Mitochondrial Membrane of Head and Neck Squamous Cell Carcinoma: Possible Role in Apoptosis

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Abstract

Purpose: Cisplatin adducts to nuclear DNA (nDNA) are felt to be the molecular lesions that trigger apoptosis, but the mechanism linking nDNA adduct formation and cell death is unclear. Some literature in the last decade has suggested a possible direct effect of cisplatin on mitochondria independent of nDNA interaction. In this study, we define separately the sequelae of cisplatin interactions with nDNA and with mitochondria in head and neck squamous cell carcinoma (HNSCC) cell lines.

Experimental Design: Cisplatin binding to mitochondrial DNA (mtDNA) and proteins was analyzed by atomic absorption spectroscopy and other methods.

Results: Following 1 hour of exposure to cisplatin, platinum adducts to mtDNA were 300- to 500-fold more abundant than adducts to nDNA; these differences were not due to differences in rates of adduct repair. Whereas HNSCC cell cytoplasts free of nDNA retained the same dose-dependent cisplatin sensitivity as parental cells, HNSCC pΔ cells free of mtDNA were 4- to 5-fold more resistant to cisplatin than parental cells. Isolated mitochondria released cytochrome c within minutes of exposure to cisplatin, and ultrastructural analysis of intact HNSCC cells by electron microscopy showed marked mitochondrial disruption after 4 hours of cisplatin treatment, whereas the nucleus and other cellular structures remain intact. The very prompt release of cytochrome c from isolated mitochondria implies that apoptosis does not require alteration in mitochondrial gene transcription. Further, cisplatin binds preferentially to mitochondrial membrane proteins, particularly the voltage-dependent anion channel.

Conclusions: Cisplatin binding to nDNA is not necessary for induction of apoptosis in HNSCC, which can result from direct action of cisplatin on mitochondria.

Cisplatin ([cis-diaminedichloroplatinum II (CDDP)] has been in widespread clinical use for more than a generation and is one of the most important chemotherapeutic agents ever introduced. Despite the importance of cisplatin in the treatment of head and neck cancer and a broad range of other malignancies, there are many uncertainties about its molecular pharmacology and ultimate mechanism of action. This has been an area of very active investigation for >2 decades.

Upon entering the low chloride intracellular environment, cisplatin is hydrated to form a positively charged species, which can react with nuclear DNA (nDNA) and other nucleophilic species within the cell (1).

Cisplatin has been most extensively characterized as a DNA-damaging agent, and the cytotoxicity of cisplatin has generally been attributed to the ability to form interstrand and intrastrand nDNA cross-links. Formation and repair of these cisplatin/nDNA adducts have been widely studied for the last 2 decades. For some time after the introduction of cisplatin, its cytotoxicity was felt to result from inhibition of DNA synthesis by cisplatin/DNA adducts, but several lines of evidence showed that this was not the case (2). More recently, it was shown that tumor cell exposure to cisplatin ultimately results in apoptosis (3, 4). However, the mechanism or mechanisms by which nuclear cisplatin/DNA adducts generate the cytoplasmic cascade of events leading to apoptosis have not been defined.

Although most investigations of the cellular and molecular pharmacology of cisplatin have focused on interactions between cisplatin and nDNA, only ~1% of intracellular platinum is bound to nDNA, with the great majority of the intracellular drug available to interact with nucleophilic sites on other molecules, including but not limited to phospholipids, cytosolic, cytoskeletal and membrane proteins, RNA, and mitochondrial DNA (mtDNA; refs. 5, 6).
Further evidence that nDNA adduct formation may not be the sole determinant of cisplatin-induced cytotoxicity comes from recent clinical studies showing that combined therapy with other agents, such as taxanes, significantly enhances the clinical efficacy of cisplatin while actually inhibiting formation of cisplatin adducts with nDNA (7).

Resistance to cisplatin can result from several mechanisms, including decreased uptake, inactivation by nucleophilic compounds, such as glutathione, or accelerated DNA repair (8). Inhibiting glutathione synthesis with buthionine sulfoximine (BSO) has been known for some time to enhance cisplatin cytotoxicity in tumor cells as well as increasing normal cell toxicity (9, 10).

We previously showed that BSO treatment of head and neck tumor cell lines was accompanied by complete loss of detectable glutathione and marked increase in cisplatin cytotoxicity. However, this enhanced apoptotic cell killing was not accompanied by significant changes in cisplatin DNA adduct formation (11).

Overexpression of Bcl-2 is associated with cisplatin resistance in several model systems (12). Recently, we showed that although Bcl-2 transfection was associated with significant acquired cisplatin resistance, it did not produce measurable alterations in nuclear cisplatin/DNA adducts (13). Because of the lack of clarity on the role of nuclear cisplatin/DNA adducts in mediating cytotoxicity and because of the Bcl-2 data suggesting the importance of mitochondrial pathways in cisplatin action, we elected to look more carefully at cisplatin interactions with mitochondria and mtDNA.

Limited studies have examined cisplatin activity in cells selectively depleted of mtDNA, with conflicting results. Loss of mtDNA has been associated with increased sensitivity to cisplatin-induced apoptosis (14), but more recent literature has shown that cells depleted of mtDNA show significant resistance to cell death mediated by a range of chemotherapeutic agents (15). Indeed, mtDNA is significantly more sensitive than nDNA to the damage induced by a range of agents (16).

Recent reports have also shown the ability of various compounds to act directly on mitochondria, inducing loss of membrane potential and release of apoptogenic proteins from isolated mitochondria (17–19). Most recently, such effects are mediated by changes in membrane potential and release of apoptogenic proteins from isolated mitochondria (20–22).

Mitochondrial damage by cisplatin has increasingly been studied as a mediator of toxicity in normal tissues in animals receiving cisplatin. Gastrointestinal toxicity (23), ototoxicity (24), and nephrotoxicity (25, 26) have all been attributed to mitochondrial effects of cisplatin.

In this study, we attempt to define separately the sequelae of the interaction of cisplatin with nDNA and with mitochondria in head and neck cancer cell lines.

Materials and Methods

Reagents. DMEM, phenol red–free DMEM, PBS, trypsin/EDTA, and glutamine were obtained from MediaTech (Herndon, VA). Fetal bovine serum was obtained from Quality Biological, Inc. (Gaithersburg, MD). BSO, CHAPS detergent, cytochalasin B, ethidium bromide, Epon 812 substitute, EDTA, Ficoll, glutathione, HEPES, leupeptin, pepstatin, phenazine methosulfate, phenylmethylsulfonyl fluoride, propidium iodide, sucrose, 5-sulfosalicylic acid, sodium pyruvate, Spurr resin, uridine, and 2,3-bis[2-methoxy-4-nitro-5-sulfophenyl]-2H-tetrazolium-5-carboxanilide inner salt (XTT) were obtained from Sigma-Aldrich (St. Louis, MO). Apopain/CPP32 fluorometric substrate was obtained from Upstate Biotechnology (Lake Placid, NY).

Cell lines. PCI-13 and PCI-51 cells were obtained from Dr. Theresa Whiteside (University of Pittsburgh, Pittsburgh, PA). UMSCC-17B cells were obtained from Dr. Thomas Carey (University of Michigan, Ann Arbor, MI). Cells were routinely maintained in DMEM supplemented with 10% fetal bovine serum and 2 mM/L glutamine.

Cisplatin production (cell emulsiﬁcation). Enucleated cells were produced by cytochalasin F treatment and Ficoll centrifugation (27–29). PCI-13 cytoplasts obtained by this method were allowed to recover for 2 hours in a 37°C CO2 incubator and then incubated for a further 24 hours after the addition of another 2 mL medium containing the indicated dose of cisplatin. For caspase-3 assay, cells and cytoplasts were harvested with trypsin/EDTA, rinsed with PBS, pelleted, and resuspended in 100 to 200 μL lysis buffer [100 mM/L HEPES (pH 7.5), 10% sucrose, 1 mM/L EDTA, 0.1% CHAPS, 1 mM/L phenylmethylsulfonyl fluoride, 1 μg/mL pepstatin, 10 μg/mL leupeptin]. Samples were frozen (−70°C) for 30 minutes and thawed on ice to lyse. Supernatant was recovered by a 5-minute spin in a 4°C microcentrifuge at maximum speed (17,500 × g).

Caspase-3 assay. Cell or cytoplasm lysate (25 μg protein) and apopain/CPP32 fluorometric substrate (50 μmol/L) were added to buffer [10% sucrose, 0.1% CHAPS, 100 mM/L HEPES (pH 7.5), 1 mM/L EDTA] to a total volume of 250 μL and incubated at 37°C for 1 hour. Samples were transferred to a microplate, and fluorescence measurements (excitation, 360 nm/emission, 460 nm) were obtained using a Bio-Tek Synergy HT multimode plate reader (Bio-Tek, Winooski, VT). Results are shown in relative fluorescence units.

Cytochrome c release from isolated rat liver mitochondria. Rat liver mitochondria were isolated as described (30). Assays were done within 4 hours of isolation. Samples were prepared to examine both caspase-3 activation (data not shown) and cytochrome c release; therefore, PCI-13 cell cytosol was included as a source of cellular enzymes for the caspase assay. PCI-13 cytosol was prepared as described (31). Protein concentration for mitochondria and cytosol was determined by the Lowry assay (32). PCI-13 and PCI-51 cells were plated to 50% confluence in DMEM supplemented with 10% fetal bovine serum and 2 mM/L glutamine; 100 μM/L BSO or control medium was added the following day. After 24 hours, cells were trypsinized, washed twice, and resuspended in cold PBS (200 μL per single well of a six-well culture plate). For mitochondrial GSH determination, mitochondria from six T175 flasks (treated with BSO or control as above) were isolated by the protocol described below for DNA purification. Pelleted mitochondria were resuspended in 100 μL PBS. Mitochondria or cells were lysed by the addition of 5% 5-sulfosalicylic acid and placed on ice for 10 minutes. The lysates were cleared by a 10-minute, 4°C spin at 17,500 × g. Lysates (25 μg protein) were then used for GSH determination according to a standard curve of purified GSH.

Generation and culture of PCI-13 and UMSCC-17B cells. \( \rho^0 \) cells. \( \rho^0 \) cell lines were derived from PCI-13 and UMSCC-17B cells by the established method of long-term exposure to ethidium bromide (33–37). Cells were cultured in DMEM supplemented with 10% fetal bovine serum, 2 mM/L glutamine, 1 mM/L sodium pyruvate, 50 μg/mL uridine, and 50 ng/mL (PCI-13) or 100 ng/mL (UMSCC-17B) ethidium bromide (EB medium). After 12 weeks, elimination of mtDNA was confirmed (see PCR and Western blot methods). mtDNA-depleted cells were cultured in medium with ethidium bromide.
were routinely maintained in EB medium; however, for XTT assays, ethidium bromide was omitted to prevent interference with absorbance measurement.

**Cytotoxicity assay.** Parental and pO cell lines were plated, in triplicate, in 96-well plates at 7,500 per well in 100 μL modified EB medium (without phenol red and ethidium bromide). Following overnight incubation, 100 μL medium containing cisplatin or irinotecan at the indicated concentrations was added. After 3 to 5 days, 50 μL medium containing 1 mg/mL XTT and 15 μg/mL phenazine methosulfate was added per well. After 1 to 2 hours of further incubation, absorbance was measured at 450 nm and expressed as a percentage of the absorbance reading of control cells (% survival). The IC50 and IC80 absorbance was measured at 450 nm and expressed as a percentage of the absorbance reading of control cells (% survival). The IC50 and IC80 values were determined using GraphPad Prism (GraphPad Software, San Diego, CA) from a sigmoidal dose-response curve (variable slope) fitted to the plot of % survival versus log [cisplatin].

**Clonogenic cell growth assay.** The cells were plated in 24-well plates at a density of 200 to 400 per well and incubated overnight to allow the cells to adhere. The following day, the medium was removed from each well and replaced with fresh medium containing different concentrations of CDDP. After incubating in a 37°C CO2 incubator for 1 hour, the CDDP-containing medium was removed and replaced with normal medium. The cells were further incubated for 7 to 12 days to allow surviving cells to form colonies. To stain the colonies, the cells were incubated for another 24 hours following the addition of p-toluidine blue solution (final dilution, 0.143 mg/mL). The Protocol COL colony detection system (Microbiology International, Frederick, MD) was used to count the number of colonies having a minimum diameter of 50 μm. Error bars represent results from triplicate wells for each experimental treatment.

**Purification of mtDNA and nDNA and measurement of platination.** PCI-13 and PCI-51 cells were treated with 50 μmol/L cisplatin for 1 or 2 hours. For each cell line, cells from 10 T175 flasks were harvested by scraping on ice. Cells were washed twice with cold PBS, and resuspended in 4 to 5 mL chilled homogenizing buffer [250 mM/L sucrose, 10 mM/L EDTA, 30 mM/L Tris-HCl (pH 7.5)]. Cells were homogenized by motorized pestle in a glass homogenizer. Nuclei were pelleted at 1,000 × g for 10 minutes at 4°C, and nDNA was purified using the PureGene DNA Isolation Cell and Tissue kit (Gentra Systems, Minneapolis, MN). The supernatant was centrifuged at 12,000 × g for 10 minutes at 4°C to pellet the mitochondria. mtDNA was purified by alkaline lysis as in Tamura (38). nDNA concentration was determined by absorbance at 260 nm. mtDNA concentration was determined by the following procedure (because the quantity of DNA was too low for absorbance measurement). The DNA preparation (2-3 μL) was digested with BamHI (New England Biolabs, Beverly, MA) to linearize the circular DNA and electrophoresed on a high sensitivity agarose gel (Reliant HS Gel System, Cambrex, East Rutherford, NJ). The linearized band was scanned and quantified using an imaging system (Alpha Innotech, San Leandro, CA) along with bands of known mass from a DNA ladder (MassRuler, Fermentas, Burlington, Ontario, Canada) on the same gel.

The absorbance values of the known bands were used to create a standard curve from which the DNA content of the mitochondrial band was determined. Platinum adduct content was measured by flameless atomic absorption spectroscopy as described (11). Platination measurements were made in duplicate.

**Bcl-2 transfection of head and neck squamous cell carcinoma.** PCI-13 cells (which are Bcl-2 negative) were transfected with a full-length cDNA for Bcl-2 as described by us previously (13).

**PCR analysis of mtDNA.** The mtDNA content of PCI-13 wild-type and pO cells was determined by PCR with primers specific to human mtDNA (upstream, 5'-CTTACGGATAACAGCGCAAT; downstream, 5'-TGAAGAAGCGATGTTGAGAG). mtDNA was detected using primers for the insulin-like growth factor-II gene (upstream, 5'-TGGCGTCTGAGTCCCTGAGT; downstream, 5'-CTCTGAGTCTGCTGCTGATA). Reactions (50 μL) contained 25 μL Herculase Hotstart PCR Master Mix (Stratagene, La Jolla, CA), 100 ng DNA, and 15 pmol of each primer. Reaction cycles were as follows: initial cycle, 5 minutes at 94°C; 30 cycles [30 seconds at 94°C, 30 seconds at 60°C, 40 seconds at 72°C]; final cycle, 5 minutes at 72°C.

**Western blotting.** For cytochrome c immunoblots, samples were prepared as described above. For Cox II immunoblots, lysates were prepared from parental and pO cells harvested by scraping on ice. Cells were washed twice with cold PBS, pelleted at 1,500 × g at 4°C, and resuspended in lysis buffer [1% NP40, 1% sodium deoxycholate, 0.1% SDS, 150 mM/L NaCl, 10 mM/L Na2HPO4 (pH 7.4)]. Proteins were separated by SDS-PAGE and transferred to nitrocellulose. Immunoblotting was done with primary antibodies at the indicated dilutions: cox II, 1:1,000 (anti-Oxphos complex IV subunit II, clone 12C4, Molecular Probes, Eugene, OR); cytochrome c, 1:200 (anti–holocytochrome c, clone 2CYTC-199, R&D Systems, Minneapolis, MN); and hsp70, 1:20,000 (anti–heat shock protein 70, clone BRM-22, Sigma-Aldrich).

Horsearid peroxidase–conjugated anti-mouse IgG (Amersham Biosciences, Piscataway, NJ) was applied 1:10,000 and detected using enhanced chemiluminescence plus (Amersham Biosciences).
Electron microscopy. PCI-13 cells in T75 flasks were treated with 50 μmol/L cisplatin in DMEM supplemented with 10% fetal bovine serum and 2 mmol/L glutamine for the indicated time. Before fixation, the cell monolayer was washed thrice with PBS. Fixative (2.5% glutaraldehyde and 2% paraformaldehyde in 5 mL PBS) was added to the flask for 30 minutes at room temperature and then replaced with PBS for three 5-minute washes. Cells were collected by scraping and pelleted (1,500 g, room temperature) with 2% agarose in PBS. Cell pellets were postfixed in 1% osmium tetroxide for 1 hour and washed with distilled water and en bloc stained with 2% uranyl acetate for 30 minutes in darkness. Samples were dehydrated with graded ethanols and propylene oxide, then embedded in a mixture of Epon 812 substitute and Spurr (Sigma-Aldrich), and polymerized overnight in an oven at 65°C. Ultrathin sections (90 nm) were cut on a Reichert-Jung microtome (Vienna, Austria) and transferred onto copper grids. Grids with sections were stained with lead citrate and examined under a Hitachi H7600 transmission electron microscope (Hitachi, Tokyo, Japan) at 60 kV. Photos were taken with a 6.7-megapixel AMT digital camera (Advanced Microscopy Techniques Corp., Danvers, MA).

Measurement of cisplatin binding to cell and mitochondrial fractions by atomic absorption spectrometry. Head and neck squamous cell carcinoma (HNSCC) cell line PCI-13 was exposed to 50 μmol/L cisplatin for 1 hour at 37°C. After cisplatin exposure, cells were

**Fig. 2.** Detection of active apoptotic pathway in cell cytoplasts. A and B, flow cytometry of propidium iodide fluorescence in PCI-13 cells (A) and cytoplasts (B). Cells (91%) in the cytoplast preparation lacked nuclei (region M1). C, cisplatin (CDDP) treatment induced dose-dependent caspase activity in enucleated cytoplasts as well as in whole cells. Columns, mean of duplicate measurements in a single assay; bars, SE. Results are representative of two independent experiments. Results are in relative fluorescence units. Caspase induction by cisplatin was statistically significant in both intact cells and cytoplasts (one-way ANOVA analysis: whole cells, P = 0.0001; cytoplasts, P = 0.0237).

**Fig. 3.** A, Western blot analysis of cisplatin-induced cytochrome c release from isolated rat liver mitochondria. Cytochrome c was measured in the supernatant of isolated mitochondria treated with cisplatin (CDDP) as indicated. B, time course of cytochrome c release from isolated rat liver mitochondria treated with 15 μmol/L cisplatin.

**Fig. 4.** Cytochrome c is rapidly released from HNSCC mitochondria treated with increasing doses of cisplatin; mitochondria from cisplatin-sensitive head and neck cancer cell line PCI-13 were treated with increasing doses of cisplatin for 40 minutes at 30°C. Supernatant from the treated mitochondria was collected and assayed for cytochrome c release by Western blot. Cisplatin doses as low as 12.5 μmol/L cause significant cytochrome c release. A, whereas cisplatin-induced rapid cytochrome c (Cyto C) release, an identical dose of the inactive stereoisomer transplatin (TDDP) did not. B, similarly, a cytotoxic dose of the topoisomerase I inhibitor irinotecan did not cause significant cytochrome c release from mitochondria.
harvested, washed to remove unbound cisplatin, and fractionated as follows. Whole-cell lysate was prepared from one tenth of the cells by standard detergent lysis. Whole-cell protein was prepared by DNase treatment of the whole-cell lysate. The rest of the cells were used to prepare isolated mitochondria by centrifugal fractionation as described for Fig. 3. One tenth of the mitochondrial preparation was used to prepare mitochondrial protein lysate in lysis buffer containing 1% Triton X-100. Voltage-dependent anion channel (VDAC) was isolated to homogeneity from the remaining mitochondrial preparation by affinity chromatography (Fig. 7; ref. 39) This method isolates all VDAC isoforms (1–3). VDAC concentration was estimated by SDS-PAGE from a standard curve of protein mass standards on the same gel. Cisplatin content in all samples was measured by atomic absorption spectrometry as described previously.

Results

Head and neck cell lines for nuclear and mitochondrial analysis. Table 1 presents relevant characteristics of two HNSCC cell lines, PCI-51 and PCI-13, selected for analysis of mtDNA platination based on our previous studies. PCI-51 overexpresses Bcl-2. Its cisplatin IC50 is 3 μmol/L. Cell line PCI-13 has a cisplatin IC50 of 0.3 μmol/L and is Bcl-2 negative. Whole-cell extracts of PCI-51 contained slightly elevated levels of glutathione compared with PCI-13 (6.8 and 5.3 pmol/μg, respectively). However, mitochondrial glutathione content was 6-fold higher in line PCI-51 compared with PCI-13.

Nuclear versus mitochondrial cisplatin/DNA adducts in head and neck tumor cell lines. Following treatment of PCI-13 and PCI-51 cells for 1 hour with 50 μmol/L cisplatin, cells were harvested, washed to remove unbound cisplatin, and fractionated as described. Agarose gel electrophoresis showed that mtDNA was highly purified with minimal nDNA contamination (Fig. 1A). Our results showed that any contaminating nDNA in the mtDNA samples would generate an insignificant contribution to the level of adducts measured in mtDNA (see below).

In the representative experiment shown in Fig. 1B, the cisplatin adduct level in the nDNA of PCI-13 and PCI-51 cells was 116 ± 1 fmol/μg and 121 ± 21 fmol/μg, respectively, whereas the level of cisplatin adducts in mtDNA was 40,700 ± 12,800 fmol/μg and 34,000 ± 3,500 fmol/μg, respectively.

Regardless of time of treatment (1 or 2 hours), pretreatment with BSO to deplete glutathione, or removal of drug for 24 hours to allow DNA repair, the level of adducts in mtDNA in both cell lines was consistently at least 2 orders of magnitude greater than the level in nDNA. MtDNA platinum adducts were a mean of 460 ± 160 times greater in PCI-13 (n = 14) and 350 ± 200 times greater in PCI-51 (n = 12). Consistent with our previous published results (11), pretreatment of both PCI-13 and PCI-51 cells with BSO resulted in a significant enhancement of cisplatin-induced cytotoxicity but no significant difference in cisplatin DNA adducts either in the nucleus or in the mitochondria (data not shown).

PCI-13 nucleus-free cytoplasts retain dose-dependent cisplatin sensitivity. PCI-13 cytoplasts were obtained through the centrifugation technique described in Materials and Methods. Following centrifugation, the nucleus was removed from 91% of the cells as confirmed by flow cytometry (Fig. 2B). Despite the absence of a nucleus, the cell cytoplasts retained dose-dependent cisplatin sensitivity as determined by caspase-3 activation. Although the cytoplasts show a somewhat elevated basal level of caspase-3 activation, they maintained a statistically significant response to escalating doses of cisplatin when compared with the parental cell line (Fig. 2C).

Cisplatin induces rapid dose-dependent release of cytochrome c from mitochondria isolated from rat liver or human HNSCC. Western blot analysis showed dose-dependent release of cytochrome c from isolated rat liver mitochondria into the supernatant (Fig. 3A). Substantial release occurred at the physiologically relevant dose of 12.5 μmol/L cisplatin [peak plasma platinum levels of 10 to 20 μmol/L are achieved clinically (40, 41)] and increased to maximal at doses of ≥ 25 μmol/L. At a dose of 15 μmol/L cisplatin, cytochrome c release was seen within as little as 5 minutes of cisplatin exposure (Fig. 3B). Similarly, mitochondria isolated from human HNSCC release cytochrome c into the supernatant after brief (40 minutes) exposure to CDDP in a dose-dependent fashion (Fig. 4A and B). Whereas cisplatin induced cytochrome c release, identical doses of the inactive transplatin or the topoisomerase inhibitor irinotecan did not cause release of cytochrome c. This dose of irinotecan is lethal to HNSCC.
4- to 5-fold resistant. PCI-13 and UMSCC-17B showed significant resistance compared with parental values of 1.8 ± 0.05 µmol/L and 2.8 ± 0.08 µmol/L, respectively; UMSSC-17B pDT had IC50 values of 5.5 ± 0.8 µmol/L and 12.7 ± 1.8 µmol/L compared with parental values of 1.8 ± 0.05 µmol/L and 2.8 ± 0.08 µmol/L. As a control, the cytotoxicity of irinotecan, which targets topoisomerase I in the nucleus, was also assayed in PCI-13 parental and pDT cells. As expected, there was little difference in response to this drug in cells depleted of mtDNA (Fig. 6).

Ultrastructural changes in mitochondria are evident by 4 hours after treatment with cisplatin. PCI-13 cells treated with 50 µmol/L cisplatin for 1 and 4 hours were examined by electron microscope (Fig. 7). Cells in each group consistently displayed the features evident in the images presented (at least 10 cells were examined at high resolution in each sample). At 1 hour, no change in mitochondrial or nuclear morphology is seen. Mitochondria in the cells treated for 4 hours (Fig. 7C and F) are pale, rounded, and lack defined cristae structure, all consistent with permeability transition and swelling (42). In contrast, nuclei and nucleoli in the same cells are intact.

Cisplatin binds preferentially to mitochondrial proteins, especially VDAC. Atomic absorption spectroscopic analysis of cellular fractions isolated from cisplatin-treated HNSCC (PCI-13) showed that the cisplatin concentration in the mitochondrial protein fraction was 10-fold higher than in the whole-cell protein fraction. Interestingly, the amount of cisplatin bound to VDAC was 24 times higher than in the whole-cell protein fraction and >200-fold higher than in the whole-cell protein fraction (Table 2). Figure 8 shows the affinity purification of VDAC for this experiment.

Discussion

Although there has been comparatively little study of direct cisplatin action on mitochondria, some studies have indicated that mtDNA-cisplatin adducts may be significantly more common than cisplatin adducts with nDNA in the same cell line treated with the same concentration of cisplatin (43, 44). This has been attributed to a lack of mtDNA repair following cisplatin exposure (45).

Mitochondria, as the sites of aerobic respiration, are the principal generators of reactive oxygen species in the cell. Mitochondria are dependent on glutathione to detoxify reactive oxygen species, preventing oxidative damage (46, 47). Despite this dependence, mitochondria are unable to synthesize glutathione. Glutathione stores within mitochondria are derived from active transport across the mitochondrial membrane against an electrochemical gradient (48). Mitochondrial glutathione concentrations are regulated and have been implicated in apoptotic regulation (49, 50). At baseline, the concentration of glutathione in mitochondria has been found to be similar to that of the cytoplasm. However, in cells exposed to oxidative stress, the concentration of mitochondrial glutathione is maintained at the expense of a decreasing cytoplasmic pool (51–53). Depletion of the mitochondrial (but not cytoplasmic) glutathione pool is associated with markedly increased sensitivity to antimycin A, which blocks oxidative phosphorylation in complex III of the electron transport chain, leading to generation of reactive oxygen species. These observations suggest that mitochondrial glutathione stores are highly regulated by the cell and may affect the cellular sensitivity to apoptotic stimuli.

![Image](63x668 to 230x708)

Fig. 6. \( p^0 \) (mtDNA depleted) cell lines are resistant to cisplatin. A, cox II Western blot showing absence of mitochondrially coded protein in \( p^0 \) cell lines. Lane 1, parental PCI-13 cells; lane 2, PCI-13 \( p^0 \) cells; lane 3, parental UMSCC-17B cells; lane 4, UMSCC-17B \( p^0 \) cells. B, PCR to detect mtDNA (lanes 2-4) and nDNA (lanes 6-8) in PCI-13 (top) and UMSCC-17B (bottom) parental and \( p^0 \) cell lines. C, XTT cytotoxicity assays (continuous CDDP exposure). Each is representative of at least three independent experiments. mtDNA absence was confirmed by PCR as in (B) before each assay. D, cisplatin dose response of UMSCC-17B parental and \( p^0 \) cells. \( p^0 \) cells are 2.1-fold resistant at the IC50 (1.2 ± 0.56 µmol/L) and 4.9-fold resistant at the IC50 (3.6 ± 0.74 µmol/L). E, irinotecan dose response of PCI-13 parental and \( p^0 \) cells. \( p^0 \) cells are 2.1-fold resistant at the IC50 (1.2 ± 0.56 µmol/L) and 4.9-fold resistant at the IC50 (3.6 ± 0.74 µmol/L). F, clonogenic assay of UMSCC-17B and UMSCC-17B \( p^0 \) cells following 1 hour of cisplatin exposure. In this experiment, mitochondrially depleted UMSCC-17B \( p^0 \) cells are approximately one log more resistant to cisplatin than the parental cells. This mirrors the results seen in the continuous exposure experiment.
Our findings support and expand recent literature reports, which have hinted that cisplatin-induced cytotoxicity may be independent of nDNA binding. In line with our results presented in Fig. 2, Mandic et al. recently showed that enucleated cells retain dose-dependent cisplatin induction of caspase-3 activation in colon and melanoma cell lines. Interestingly, in that study, although the enucleated cells retained sensitivity to cisplatin, they became resistant to the DNA-damaging topoisomerase II inhibitor etoposide (27). Similar results have been reported in colon cancer cell cytoplasts treated with oxaliplatin (54). Other agents may also exert proapoptotic effects through direct interactions with mitochondria (55).

Although we show that head and neck tumor cells lacking mtDNA become cisplatin resistant, they are not completely cisplatin insensitive. A previous study with \( \mu^0 \) osteosarcoma cells showed that the loss of mtDNA does not result in a complete absence of cytochrome \( c \) release from tumor mitochondria when damaged by the toxin staurosporine (56). This result combined with our own suggests that cisplatin-induced mitochondrial toxicity is not entirely DNA dependent. Although we show that mitochondrial platinum adducts are measured at a concentration at least 2 orders of magnitude greater than is measured in the nucleus, the observations that cytochrome \( c \) release from isolated mitochondria can be seen within 5 minutes of cisplatin exposure and that, in whole cells, ultrastructural damage of mitochondria is clearly visible within a few hours of cisplatin treatment both argue that a direct effect on mitochondrial gene transcription is not necessary for the

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<th>Fraction</th>
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<td>65</td>
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*DNase/RNase treated.

Cisplatin binding to whole-cell protein, mitochondrial protein, and VDAC protein isolated from cisplatin-treated HNSCC

Fig. 8. Cisplatin binds preferentially to mitochondrial proteins, especially VDAC. HNSCC (PCI-13) cells were exposed to 50 µmol/L cisplatin for 1 hour, and various cell fractions were then isolated. Total cisplatin binding to the specific fractions was measured by atomic absorption spectroscopy. VDAC was affinity purified from cisplatin-treated cells as described in the text. M, marker; MP, SDS-PAGE of starting mitochondrial protein fraction; AP, SDS-PAGE of affinity purified VDAC.
mitochondrial toxicity of cisplatin. Importantly, we show that cytochrome c release is seen at cisplatin concentrations seen following bolus injection of the drug clinically.

Corresponding to our observation that Bcl-2 transfection is associated with resistance to cisplatin-induced release of cytochrome c from the mitochondria, a recent study showed that Bcl-2 overexpression was associated with diminished disruption of mitochondrial ultrastructure following cisplatin treatment as determined by electron microscopy (57). Another recent study of normal middle ear hair cells clearly showed that otoxicity resulting from cisplatin treatment is associated with severe disruption of mitochondrial structure (58).

Although mitochondria lack the capability for nucleoside excision repair, the marked elevations in mtDNA versus nDNA platinum adducts are not explained by differences in repair alone. Adduct levels in both mtDNA and nDNA decreased at similar rates after cisplatin exposure (data not shown). The electrochemical gradient resulting in a net negative charge within mitochondria may play a role in the significant accumulation of the positively charged cisplatin that we measured. Indeed, alterations in mitochondrial membrane potential have been associated with significant shifts in cisplatin sensitivity (59, 60).

Although we have shown a direct effect of cisplatin on mitochondria, which is not dependent on nuclear or cytoplasmic signaling, our results do not exclude the importance of such effects in generating apoptotic signals following cisplatin exposure. Cisplatin exposure is associated with localization of p53 to the mitochondria, which enhances binding of mitochondrial transcription factor A to cisplatin-damaged mtDNA (61). The p38 mitogen-activated protein kinase pathway may also have a role in cisplatin-induced apoptosis. Activation of p38 has been shown following cisplatin exposure in several studies, and this can be inhibited by RACK1 and AKT2 (62–64).

Cells deficient in nDNA repair are sensitive to cisplatin, but this may be secondary to DNA damage caused by reactive oxygen species generated following release of mitochondrial cytochrome c, accelerating apoptosis.

In addition to interacting with nucleophilic sites in DNA, cisplatin binds to nucleophilic amino acid residues in proteins, including cysteine, methionine, and histidine. Cross-linking or modification of critical intracellular proteins by cisplatin is a potentially important, and little studied, cytotoxic mechanism.

The effect of cisplatin on mitochondria may be mediated through protein modification or cross-linking. Components of the permeability transition pore, such as the VDAC and the adenine nucleotide translocator, are interesting candidates. Both contain vulnerable residues, and modification of a cysteine residue of adenine nucleotide translocator by thiol cross-linking agents has been shown to cause mitochondrial membrane permeabilization and apoptosis (65). Our finding that the concentration of cisplatin bound to VDAC is more 200-fold higher than the amount bound by total cellular proteins raises the interesting possibility that cisplatin binding to this key mitochondrial membrane protein could significantly alter its structure or function, facilitating the release of cytochrome c.

In conclusion, our data indicate that, in HNSCC cell lines, the cytotoxic effect of cisplatin is, in significant part, mediated through direct action on mitochondria. Although cisplatin interactions with nDNA may have important cellular effects contributing to apoptosis, cisplatin binding to nDNA is not necessary for induction of apoptosis. Indeed, the data presented here, in combination with recent findings discussed above, suggest that mitochondria may be the principal and sufficient target of cisplatin in this group of diseases. Further studies will be necessary to determine the precise mitochondrial target(s) of cisplatin and whether these observations are applicable to other tumor types. In the future, mitochondria-based drug screening assays may be important in the evaluation of new chemotherapeutic agents.

References


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