Morphofunctional effects of mitotane on mitochondria in human adrenocortical cancer cells

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Abstract
At present, mitotane (MTT) represents the first-line pharmacological approach for the treatment of advanced adrenocortical carcinoma (ACC). Despite clear evidence that the drug can reduce the clinical signs of steroid excess in secreting ACC, the mechanism mediating the possible toxic effect of MTT on tumor cells still remains obscure. This study investigated the intracellular events underlying the toxic effect of MTT by studying qualitative and quantitative alterations in mitochondrial morphology and functions in human adrenocortical cancer cell lines, H295R and SW13. Increasing concentrations of MTT resulted in rapid intracellular accumulation and conversion of the drug. Cytostatic and cytotoxic effects were evident at doses corresponding to the therapeutic window (30–50 μM) through an apoptotic mechanism involving caspase 3/7. Electron microscopic analysis of cell mitochondria displayed MTT-induced dose- and time-dependent alterations in the morphology of the organelle. These alterations were characterized by a marked swelling and a decrease in the number of respiratory cristae, accompanied by a significant depolarization of the mitochondrial membrane potential, finally leading to the disruption of the organelle. A drastic reduction of oxygen consumption was observed due to mitochondrial membrane damage, which was accompanied by a decrease in the levels of VDAC1 integral membrane channel. These findings contribute to better understand the intracellular mechanism of action of MTT in ACC cells, showing that its cytotoxic effect seems to be mainly mediated by an apoptotic process activated by the disruption of mitochondria.

Key Words
- adrenal cortex
- carcinoma
- intracellular signalling

Introduction
Adrenocortical carcinoma (ACC) is a rare and aggressive endocrine tumor with an incidence of 0.5–2 cases per million people every year. ACC prognosis is mainly dependent on the stage at diagnosis (Fassnacht et al. 2009), with the overall 5-year survival rate being drastically reduced from 35% (Allolio & Fassnacht 2006) to

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5–10% if the mass is diagnosed at the advanced metastatic stage (Fassnacht et al. 2009). At present, the complete surgical removal of the tumor remains the most valuable option for ACC cure as most of the cytotoxic chemotherapeutic regimens proposed so far have displayed very limited efficacy. However, the use of the adrenotoxic agent mitotane (MTT or dichlorodiphenyldichloroethane (DDD)) as the first-line therapy, alone or in combined chemotherapy (Fassnacht et al. 2012), for advanced carcinoma as well as in adjuvant regimen (Terzolo et al. 2007) has been demonstrated to significantly enhance progression-free survival.

MTT (DDD), a derivative of the insecticide dichlorodiphenyltrichloroethane (DDT), has been successfully introduced for the treatment of Cushing’s syndrome (Temple et al. 1969, Baudry et al. 2012) and lately for that of ACC (Metz 1973) due to its adrenotoxic effect: this therapeutic strategy results in an overall response in 25–30% of ACC patients (Hahner & Fassnacht 2005). MTT metabolism and protein binding are species specific and higher in cattle and dogs than in humans as demonstrated in adrenal cortex homogenates (Cai et al. 1995a,b, Lindhe et al. 2002). Inhibition of steroidogenesis, in particular, of cortisol production, has been associated with the inhibition of specific enzymes responsible for cholesterol metabolism in the adrenal (StAR, P450scc, and CYP11B1; Asp et al. 2010, Lin et al. 2012). However, the mechanism underlying the adrenolytic activity of MTT has not yet been elucidated. Some early sporadic studies have reported the qualitative effects of MTT on animal adrenals (Kaminsky et al. 1962, Moore et al. 1980, Krüger et al. 1984) or on the human atypical estrogen-secreting adrenal cancer cell line Fang-8 (Fang 1979). Recently, by evaluating MTT-induced variation of the proteomic profile of the ACC H295R cell line (Stigliano et al. 2008), MTT has been suggested to interact with some proteins belonging to the classes of energetic metabolism, stress response, cytoskeletal structure, and tumorigenesis. Thus, the aim of the present study was to evaluate the potential toxic activity of MTT and its cellular targets in human cell models of ACC.

Materials and methods

Materials

Purified stereoisomers of MTT, o,p’-DDD (DDD), and of its metabolite dichlorodiphenyldichloroethylene (DDE), o,p’-DDE (DDE), and of DDT, o,p’-DDT (DDT), were purchased from Sigma–Aldrich. 2-Propanol, n-hexane, acetonitrile, and water of HPLC grade were obtained from Panreac Quimica (Barcelona, Spain). All other reagents were procured from Sigma–Aldrich unless differently indicated.

Cell cultures

Human SW13 and H295R ACC cell lines (American Type Culture Collection, Manassas, VA, USA) were cultured in 10% FBS (Euroclone, Pero, Milan, Italy)–DMEM/F-12 medium supplemented with 2 mM L-glutamine, 100 U/ml penicillin, and 100 µg/ml streptomycin (Sigma–Aldrich) and enriched with a mixture of insulin/transferrin/selenium for H295R cells.

Human primary fibroblasts were obtained from patients undergoing elective abdominal surgery, after getting written informed consent. The cells were cultured in 10% FBS (Euroclone)–DMEM supplemented with 2 mM L-glutamine, 100 U/ml penicillin, and 100 µg/ml streptomycin (Sigma–Aldrich) and used at passage 3.

For all the experiments, the cells were cultured for the indicated time under the indicated treatment with 1% FBS at 37 °C in a humidified 5% CO2 atmosphere.

HPLC measurement of intracellular levels of DDD and DDE

Stock solutions (1 mg/ml) of DDD and DDE were made up in methanol and stored at 4 °C. Working solutions (100 µg/ml) were obtained by 1:10 dilution of the stock solutions in methanol. Standard solutions (5, 10, and 20 µg/ml) were prepared by adding 5, 10, and 20 µl of the working solutions to 100 µl of a serum-free solution respectively. To both the standard solutions and methanol-extracted cell samples, 100 µl of the internal standard solution (o,p’-DDT, 20 µg/ml) were added. All the samples and standards were subsequently extracted by adding 2 ml n-hexane and 2 ml 2-propanol. After centrifugation at 2500 g for 5 min, the resultant supernatants (4 ml) were evaporated to dryness under a nitrogen flow. The residues were resuspended with 60 µl mobile phase, and 30 µl were injected into the HPLC system (Chromsystems, München, Germany) equipped with a CLC300 isocratic pump, a CLC200 autosampler, and a SPD-20A/20AV u.v.–Vis detector (Shimadzu, Kyoto, Japan). Data acquisition and integration were performed using the Geminyx Software (Chromsystems). All the separations were achieved with a C18 reversed-phase column (ref. 24100, Chromsystems). The mobile phase consisted of an acetonitrile–water–phosphoric acid mixture (85:15:0.05%/oo) delivered at a flow rate of 0.6 ml/min, and the eluate was monitored at 218 nm.
MTS assay
The H295R cells were seeded in 96-well plates at a density of 1×10^4 cells/well. After 24 h of starvation, the cells were treated with vehicles (control) or increasing doses of MTT for the indicated times. Cell viability was assessed by 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium salt (MTS) assay (Promega), according to the manufacturer's instructions. The samples were analyzed using an ELISA plate reader (Victor 1420, PerkinElmer, Monza, Italy) at 490 nm wavelength to measure optical density. Each experimental point was performed in quintuplicate in at least three independent experiments.

DNA synthesis assay: [3H]thymidine uptake
DNA synthesis was evaluated according to the amount of [3H]TdR incorporated into trichloroacetic acid (TCA)-precipitated materials. Cells grown in 10% FBS complete medium till 70% confluence were starved for 24 h and then treated with increasing concentrations of MTT for 48 h, pulsing them with 1.0 μCi/ml [3H]TdR (6.7 Ci/mmol) 4 h before stopping proliferation in ice-cold 10% TCA. After washing in TCA and then in methanol, the cells were solubilized in 0.2 M NaOH, and radioactivity was measured using a scintillation counter. Experiments were performed in triplicate and repeated at least three times.

Viable cell count
The cells were seeded in 12-well plates (1×10^5 cells/well) and were treated with vehicles (control) or with increasing doses of MTT for the indicated times after 24 h of starvation. At each time interval, the cells were trypsinized and counted using a hemocytometer. Mean cell number was obtained by counting triplicates in three different experiments. Dead cells were excluded by trypan blue exclusion test.

Transmission electron microscopy and morphometry of mitochondria
H295R and SW13 cells cultured under different experimental conditions were pelleted by centrifugation, fixed in 4% glutaraldehyde, post-fixed in 1% osmium tetroxide, and embedded in Epon 812. Ultrathin sections were stained with uranyl acetate and alkaline bismuth subnitrate and examined under a JEM 1010 electron microscope (Jeol, Tokyo, Japan) at 80 kV. Micrographs were taken at ×10 000 and ×50 000 final magnifications using a MegaView III digital camera (SIS-Soft Imaging System, Munster, Germany), and the ones obtained at the latter magnification were used for the morphometric analysis of mitochondria. Five cytoplasmic fields were randomly chosen in each experimental condition, for a total of about 20 mitochondrial profiles. Mitochondrial surface area and perimeter of cristae were measured using the iTEM image analysis software (SIS). Mitochondrial swelling/disruption was estimated by calculating the ratio of surface area to overall perimeter of cristae. Normal and swollen mitochondria were counted on at least five ×10 000 magnification micrographs for every sample.

Evaluation of apoptosis by Muse cytofluorimetric analysis
After 48 h of treatment with 30 and 50 μM MTT, H295R and SW13 cells were analyzed for the detection of early and late apoptosis and cell death using Muse Annexin V/Dead Cell (cat #MCH100105, Merck Millipore, Billerica, MA, USA) and Muse Caspase 3/7 (cat #MCH100108, Merck Millipore) assays respectively, according to the manufacturer's instructions. Based on positivity for caspase 3/7 activity or phosphatidylserine exposure (positivity for annexin V) in apoptotic cells and simultaneous detection of dead cells, positive for the nuclear dye 7-aminoactinomycin D (7-AAD), both assays enable to differentiate four populations in each sample by cytofluorimetric separation on a Muse automated cell analyzer (Merck Millipore): non-apoptotic live (lower left panel: 7-AAD negative, apoptosis negative), non-apoptotic dead (upper left panel: 7-AAD positive, apoptosis negative), 90 85 80 75 70 30 20 10 20 30 40 50 60 70 80 90 100 110 MTT (μM)

Figure 1
H295R cells capture and metabolize MTT. H295R cells were treated with increasing concentrations of MTT for 24 h. After several washes, the cells were lysed in HPLC buffer and the concentrations of intracellular DDD and DDE were evaluated by HPLC. Data are expressed as mean ± S.E.M. values. Experiments were performed in duplicates three times.
apoptotic live (lower right panel: 7-AAD negative, apoptosis positive), and apoptotic dead (upper right panel: 7-AAD positive, apoptosis positive) cells.

Preparation of mitochondrial fraction

The cells (1 × 10⁶) were washed once with cold PBS, harvested by scraping with cold PBS, and centrifuged for 5 min at 800 g at 4 °C. The pellet was suspended in 0.5 ml ice-cold medium (250 mM sucrose, 2 mM HEPES, and 0.1 mM EGTA, pH = 7.4), and the cells were disrupted by 40 passes in a homogenizer (tight-fitting). The homogenate was then centrifuged for 10 min at 800 g at 4 °C, and the mitochondria-rich supernatant was collected. The cell debris pellet was resuspended in a further 0.5 ml of the same medium, homogenized, and centrifuged as described earlier. The two supernatants were pooled and centrifuged for 20 min at 13 000 g at 4 °C. The resulting mitochondrial pellet was suspended in 100 μl medium containing 120 mM KCl, 20 mM HEPES, 2 mM MgCl₂, 1 mM EGTA, and 5 mg/ml BSA at pH = 7.4 (Kirby et al. 2007) and analyzed for oxygen consumption. Supernatants were quantified for protein content (Coomassie Blue reagent, Bio-Rad). All procedures were carried out on ice or at 4 °C.

Oxygen consumption analysis

Quantification of oxygen consumption by mitochondria isolated from cells exposed or not exposed to different concentrations of MTT for 24 and 48 h was carried out using the Oxygraph system (Hansatech Instruments, Norfolk, UK). Mitochondria (300 μg) were loaded into the chamber in 300 μl respiration buffer (125 mM KCl, 10 mM Tris/MOPS, 10 mM EGTA/Tris, and 1 mM potassium phosphate monobasic, pH = 7.4) (Frezza et al. 2007).

Estimation of mitochondrial membrane potential (Δψ m)

Mitochondrial membrane potential (Δψ m) was estimated in adherent live cells using the cationic, lipophilic dye

Figure 2

MTT inhibits cell viability and proliferation. H295R cells were grown in the presence or absence of increasing concentrations of MTT (5–30 μM). After different time intervals, the cells were harvested and live cells were directly counted (A) or cell viability was estimated by the MTS assay (B). Cell proliferation was evaluated by thymidine incorporation (2-h pulse) in H295R cells treated or not treated with MTT (10–30 μM) for 48 h. Data are expressed as mean % increase ± S.E.M. of MTS absorbance (A), cell count (B), and thymidine incorporation (C) vs the respective controls in five experiments performed in quintuplicates or in triplicates respectively. Statistical analysis was performed with ANOVA followed by Dunnett’s test: *P<0.01 and **P<0.0001 vs the respective controls. Dose-response curves of cell viability and count obtained for the indicated time points in treated H295R cells as described above were used to calculate the respective MTT IC₅₀ indicated in the inset (D). MTT and cell count curves did not differ statistically significantly.
JC-1 (5,5′,6,6′-tetrachloro-1,1′,3,3′-tetraethylbenzimidazole carboxyanine iodide). For spectrofluorimetric detection, the cells were plated into 96 wells, whereas for fluorescence microscopy, they were plated on glass cover-slips in six-well plates. After 48 h of treatment with different doses of MTT or 2 h of treatment with 10 μM valinomycin, the washed cells were incubated with a fresh culture medium containing 2.5 μg/ml JC-1 fluorescent dye for 30 min at 37 °C in the dark. After washing in PBS, fluorescence intensities were measured at 527 nm (green) and 590 nm (red) wavelength emission on an ELISA plate reader (Victor3 1420, PerkinElmer). For fluorescence microscopy, coverslips were washed and mounted in an antifade mounting medium (Prolong Gold, Life Technologies Italia) on glass slides. The two-channel fluorescence emission was acquired with a Leica DM4000 epifluorescence microscope (Leica Microsystems GmbH, Wetzlar, Germany).

Western blot
The cells were lysed in RIPA buffer, and after protein measurement, equal amounts of proteins (30 μg) for each sample were separated by SDS–PAGE and transferred onto PVDF membranes (Immobilon, Merck Millipore). Bound antibodies were revealed using ECL reagents (Immobilon, Merck Millipore) and analyzed on a Bio-Rad ChemiDoc Imaging System for dedicated chemiluminescent image acquisition (Imaging and Analysis Software by Bio-Rad, Quantity-One). The polyclonal anti-B subunit of succinate dehydrogenase (SDHB) antibody was obtained from Atlas Antibodies (Stockholm, Sweden). The polyclonal anti-VDAC1 (sc-8828), anti-actin (sc-1615), and anti-caspase 3 (sc7272) primary antibodies and all the peroxidase-conjugated secondary antibodies were obtained from Santa Cruz Biotechnology.

Statistical analysis
Statistical analysis was performed using SPSS Software 18.0 version (SPSS, Inc.). One-way ANOVA followed by Dunnett’s post hoc test was applied for multiple comparisons. A P value <0.05 was considered statistically significant. Results are expressed as means ± S.E.M. unless otherwise stated. On the basis of cell count and MTS curves using Origin Software 6.1 version (OriginLab Corporation, Northampton, MA, USA), 50% inhibitory concentration (IC50) of MTT was calculated.

Results
In order to demonstrate that MTT is metabolized not only in hepatocytes, but also locally in the adrenal tumor cells and is accumulated with its metabolites in the cells, we treated the ACC cell line H295R for 24 h with increasing doses of MTT and quantified DDD and its lipophilic metabolite DDE in the washed cell extracts by HPLC. Cell treatment with MTT (10–100 μM) resulted in the detection of intracellular levels of DDD and DDE, which increased in parallel with the added doses of MTT (Fig. 1). Correspondently, MTT treatment resulted in a dose-related and time-dependent reduction in cell number, as demonstrated by the direct cell count (Fig. 2A). This reduction was due to a combined inhibitory effect on viability and proliferation, as evaluated by the MTS assay (Fig. 2B) and thymidine uptake at 48 h (Fig. 2C) respectively. Respective
Figure 4

MTT induces apoptosis in adrenocortical cancer cells. After 48 h of treatment without (A and F) or with MTT of 30 (B and G) and 50 µM (C and H) as well as with H₂O₂ of 200 µM (D and I), used as a positive control for the induction of apoptosis, H295R cells were trypsinized and analyzed with a Muse automated cell analyzer (Merck Millipore), according to the manufacturer's instructions (A, B, C, D and E). Analogous independent experiments were analyzed with Muse Caspase-3/7 Assay (cat #MCH100108, Merck Millipore), according to the manufacturer's instructions (F, G, H, I and J). Based on positivity for caspase 3/7 activity or (cat #MCH100108, Merck Millipore), according to the manufacturer's instructions (F, G, H, I and J). Based on positivity for caspase 3/7 activity or phosphatidylserine exposure (positivity for annexin V) in apoptotic cells and simultaneous detection of dead cells (7-AAD dye positivity), both assays enabled to differentiate four populations in each sample by cytofluorimetric separation on a Muse automated cell analyzer (Merck Millipore): non-apoptotic live (lower left (LL): 7-AAD negative, apoptosis negative), non-apoptotic dead (upper left (UL): 7-AAD positive, apoptosis negative), apoptotic live (lower right (LR): 7-AAD negative, apoptosis positive), and apoptotic dead (upper right (UR): 7-AAD positive, apoptosis positive) cells. Positivity for annexin V (A, B, C, D and E) and caspase 3/7 (F, G, H, I and J) activity helped in identifying early (apoptotic live fraction, LR) and late phases of apoptosis (apoptotic dead fraction, UR). Data obtained using the two kits were not completely superimposable due to the different markers used for the detection of apoptosis. Representative cytofluorimetric dot plots of four independent experiments performed for annexin V (A, B, C and D) or caspase 3/7 (F, G, H, I and J) detection. Data are given as mean ± S.E.M. percentage of cells in the four populations identified with annexin V and caspase 3/7 activity respectively. Inset, mean ± S.E.M. total apoptosis % is indicated for each population. Statistical analysis was performed with Student’s t-test for independent data, n=4, *P<0.05, **P<0.01, and ***P<0.001 vs the respective controls.

IC₅₀ as calculated by the MTT inhibition curves for cell count and viability at the different time points, ranged between 10 and 20 µM (Fig. 2D). The toxic effect exerted by MTT was also evident in the ACC cell line SW13 (Fig. 3A). Conversely, increasing doses of MTT up to 50 µM did not significantly affect the cell viability of primary fibroblasts treated for comparable time intervals (Fig. 3B), suggesting the drug specificity for adrenal cancer cells.

Cytofluorimetric analysis of apoptotic events through the exposure of annexin V marker in the H295R cells (Fig. 4A, B, C, D and E) showed that 48 h of treatment with 30 and 50 µM MTT stimulated a net increase in the percentage of total apoptotic cells compared with the controls. H₂O₂ was used as a positive control. Interestingly, at 48 h, the effect of 30 µM MTT was more evident in the live fraction of apoptotic cells (Fig. 4B (LR) and E), while higher doses mainly affected the dead fraction of apoptotic cells (Fig. 4C (UR) and E). The mean ± S.E.M. increase in total apoptosis percentage was evident in response to MTT and H₂O₂ (Fig. 4E). These results were substantially confirmed by cytofluorimetric analysis performed using an apoptotic marker of later events in the apoptotic execution program, which evaluates caspase 3/7 activity (Fig. 4F, G, H, I and J). Figure 4J also reports the increase in the percentage of total apoptotic cells induced by 30 and 50 µM MTT and H₂O₂ after 48 h of treatment. Similar results were also obtained in the SW13 cells (not shown).

To evaluate which intracellular structures are mainly affected by MTT treatment, we performed electron microscopy on H295R cells treated with increasing doses of MTT for 24 h (Fig. 5A, B, C, D, E, F, G and H) and with 10 µM MTT up to 10 days (Fig. 5I, J, K, L, M, N, O and P). Electron microscopy clearly showed a progressive alteration in mitochondrial structure dependent both on the dose of MTT and on the length of treatment duration. Dose-related and time-dependent mitochondrial
abnormalities were visible at different magnifications, consisting of prominent swelling and rarefaction of cristae and electron-dense myelin-like remnants of the inner mitochondrial membranes. Vacuolation of cytoplasm was more pronounced than those observed at 24 h (B and F). At 48 h (J and N), mitochondrial swelling and disruption of cristae were the most evident changes; at 5 days (K and O), most mitochondria were swollen and often showed myelin-like membrane debris, while some cells appeared necrotic, with swollen nuclei and severely vacuolated cytoplasms (asterisk); at 10 days (L and P), almost all cells were lost, and the only ones present showed intermediate features between apoptosis and necrosis, such as hyperchromatic chromatin clumps typical of apoptotic nuclei and plasma membrane rupture typical of cell necrosis; mitochondria were barely recognizable as such. Control cells at 10 days of culture (I and M). Arrows indicate the cytoplasmic areas corresponding to the high-power panels (E, F, G, H, M, N, O and P). Bars = 1 μm.

Figure 5
MTT treatment dose- and time-dependently affects mitochondrial morphology in H295R cells. (A, B, C, D, E, F, G and H) Representative electron micrographs of cells grown for 24 h in the absence or presence of increasing concentrations of MTT (10–50 μM). The untreated 24-h control cells (Ctrl, A and E) showed rod-shaped mitochondria with numerous cristae and dense matrix. Conversely, in the MTT-treated cells, dose-related mitochondrial abnormalities were visible, consisting of prominent swelling, matrix clearing, and rarefaction and disruption of cristae. The most severe mitochondrial changes consisted of marked vacuolation containing electron-dense myelin-like remnants of the membranes (D and H). (I, J, K, L, M, N, O and P) Representative electron micrographs of cells grown for different times in a culture medium to which MTT was added (10 μM). Time-related mitochondrial abnormalities (I, K, L, N, O and P) were visible and evidently and progressively worsened at high doses and longer treatment durations. At 5 days, most of the cells were lost due to the highly toxic effect of MTT and only dead cells were present, typically showing intermediate...
features between apoptosis and necrosis (aponecrosis), e.g. nuclear chromatin condensation and plasma membrane rupture; mitochondria were no more recognizable as such (Fig. 5I, J, K, L, M, N, O and P). Similar ultrastructural alterations following increasing doses and time of MTT treatment were detected in the SW13 cells (Fig. 6). The total number of mitochondria decreased with increasing doses of MTT, showing a significant decrease in the number of normal mitochondria paralleled by a significant increase in that of swollen/disrupted ones (Figs 5 and 6), as also shown in Fig. 7A, at 48 h of MTT treatment. In order to better elucidate the nature of the mitochondrial lesions induced by MTT, we calculated the ratio of the whole area to the total length of the inner membrane cristae in individual mitochondria viewed by electron microscopy. Such a parameter takes into account both aspects of mitochondrial damage, i.e. mitochondrial swelling (increase in mitochondrial area) and decrease in the number and extension of the inner membrane cristae (decrease in the total length of cristae). This ratio significantly increased with increasing doses of MTT (Fig. 7B) and with the length of the treatment duration (Fig. 7C) in the H295R cells. Similar results were observed in the SW13 cells (not shown).

Since the reduction in the extension of inner cristae may affect mitochondrial functions, we next investigated whether the structural lesions also resulted in alterations in mitochondrial functions. Mitochondrial membrane potential ($\Delta \psi_m$) was estimated in adherent live cells using the cationic, lipophilic dye JC-1. Fluorescence microscopic analysis of H295R cells treated for 48 h with MTT resulted in a clear shift of the JC-1 molecule from red to green fluorescence (Fig. 8C and D). This shift was higher than that observed in the untreated control (Fig. 8A) and also more pronounced than that observed in the positive control, represented by 48-h valinomycin-treated cells (Fig. 8B). Quantitative analysis of fluorescence emission revealed a significant reduction in red fluorescence accompanied by an increase in green emission following MTT treatment (Fig. 8E), suggesting that MTT induced a reduction of mitochondrial membrane potential in the H295R cells.

We next evaluated whether the observed derangement of the mitochondrial membrane potential may affect mitochondrial respiration. Respiration was evaluated as inducible oxygen consumption in live mitochondria isolated from H295R cells treated or not treated for 48 h with increasing concentrations of MTT.
(10, 30, and 50 µM). Figure 9B shows mitochondrial oxygen consumption after addition of the respiratory substrate succinate. MTT strongly reduced oxygen consumption in a dose-dependent manner, with up to an 82 ± 10% inhibition being obtained with 50 µM MTT (Fig. 9A and B).

Since the reduced oxygen consumption observed in response to MTT treatment may be due to a general damage in the organization of the mitochondrial membranes and/or to a specific interference with the enzymes of the respiratory chain, we measured the effect of MTT on the expression of a structural channel in the outer membrane, the voltage-dependent anion channel, VDAC, and of one of the respiratory chain enzymes present in the inner mitochondrial membrane, SDH. Western blot analysis of total cell lysates showed that the levels of the SDHB remained rather stable after 48 h of MTT treatment (Fig. 10A and B). Conversely, the levels of VDAC were significantly reduced in a dose-dependent manner (Fig. 10A and B). In parallel, MTT increased the levels of cleaved caspase 3 (Fig. 10A and B), confirming the cytofluorimetric results obtained with the same 48 h of treatment with 30 and 50 µM MTT (Fig. 4).

Discussion

MTT has long been used as the first-line chemotherapeutic agent for the treatment of patients affected by ACC, which represents a rare and aggressive neoplasm with poor prognosis (Metz 1973). Despite clear evidence that the drug can reduce the steroid excess in the secreting forms of ACC as well as in Cushing’s syndrome, the mechanism underlying the possible toxic effect on human tumor cells still remains obscure. In this study, using H295R cell line as a model of human ACC, we demonstrated that MTT is actively taken up and metabolized by cancer cells, where it accumulates with its metabolites, as demonstrated by the detection of intracellular levels of DDD and DDE. In contrast to the hepatic biotransformation of MTT, which is mainly supported by microsomal P450 enzymes (Martz & Straw 1980), in adrenals, mitochondria are the primary source of enzymes producing MTT metabolites.
In vitro experiments in adrenal cell lines and ex vivo experiments on adrenal slides (Cai et al. 1995b, Lund & Lund 1995, Lindhe et al. 2002) have demonstrated that CYP11B1, in addition to its role in cortisol production, activates MTT and its metabolites through an initial hydroxylation step. Selective expression of this enzyme in the zona fasciculata and reticularis provides tissue specificity for MTT activation. Selective inhibition of CYP11B1 reduces the irreversible binding of MTT and its metabolites to the proteic component of adrenal cells and adrenocortical tissue (Cai et al. 1995a, Lund & Lund 1995, Lindhe et al. 2002). However, the relationship between this binding and the effect of MTT is yet to be established. Interestingly, there is a strict correlation between the variability in MTT activity in different species and the corresponding rank order of metabolic transformation/covalent binding. Bovine and canine adrenocortex seems to be the most sensitive to

**Figure 8**

MTT treatment results in disrupted mitochondrial membrane potential (ΔΨm) in H295R cells. Cells were treated with different doses of MTT or with valinomycin (VAL), as a positive control, for 48 h. Mitochondrial membrane potential (ΔΨm) was estimated using the cationic, lipophilic dye JC-1 (5,5′,6,6′-tetrachloro-1,1′,3,3′-tetraethylbenzimidazole carbocyanine iodide), which accumulated as red fluorescence-emission J-aggregates at high negative transmembrane potential in functional mitochondria or as diffused green fluorescence-emission monomers at low transmembrane potential in altered mitochondria. Epifluorescence microscopy of H295R cells not treated (A) or treated with 10 μM MTT (C), 30 μM MTT (D), or 1 μM VAL (B) in a representative experiment. Merged images were obtained with excitation/emission of 490/530 nm for green fluorescence and 525/590 nm for red fluorescence. (E) Quantitative evaluation of red and green fluorescence emission. Data represent optical density (OD) mean ± S.E.M. at the two emission wavelengths (530 and 590 nm for green and red fluorescence respectively) evaluated spectrofluorimetrically in at least six replicates for each sample in three independent experiments. Statistical analysis was performed with ANOVA followed by Dunnett’s test: *P < 0.005 and **P < 0.001 vs the respective red and green controls.
the metabolism and effects of MTT in comparison with human normal and cancer tissue (Cai et al. 1999a,b). This relationship may explain the variability in the response to MTT observed in patients with adrenal cancer, depending on the ability of the tumor cell to metabolize and activate the drug (Schtueingart 2007). The majority of data addressing the effects of MTT in adrenocortex report the inhibitory effect of this compound on steroidogenesis. In particular, mitochondrial enzymes, but not microsomal ones, in the steroidogenic pathways (STAR, CYP11A1, and CYP11B1) have been demonstrated to be inhibited in the H295 and H295R cells (Stigliano et al. 2008, Lin et al. 2012). Different strains of H295 cells have shown different degrees of inhibition in steroid secretion, depending on their steroidogenic profiles (Samanfari et al. 2007). In contrast to the inhibitory effect on steroid production, the cytotoxic effects of MTT responsible for adrenal necrosis are less clear. Our data demonstrate that increasing concentrations of MTT, in the range of the therapeutic window (14–20 mg/l corresponding to 43.7–62.5 μM MTT), result in cellular toxicity in H295R cells through the disruption of mitochondrial integrity and functions. Early studies have reported qualitative mitochondrial alterations in adrenals of animals treated with MTT (Kaminsky et al. 1962, Moore et al. 1980, Krüger et al. 1984) and in an estrogen-secreting human ACC cell line. Nevertheless, these effects were observed only at doses as high as 168 μM, whereas 80 μM MTT was not effective on either mitochondrial structure or steroid secretion (Fang 1979). It may be hypothesized that low intracellular levels of MTT or short treatment durations affect primarily mitochondrial steroidogenic enzymatic activity with little effects on cellular viability (Stigliano et al. 2008, Lin et al. 2012). Conversely, higher doses of MTT or longer treatment durations cause a mitochondrial toxic effect, in addition to reduced steroid hormone secretion (Fang 1979, present paper), resulting in cell death (Cai et al. 1995b, present paper). Our data clearly show that MTT levels in the therapeutic window (30–50 μM) exert a cytotoxic effect associated with the inhibition of cell proliferation. Increasing doses of MTT affected the normal morphology of H295R cell mitochondria starting from 24 h of incubation. Prolonged exposure to the drug up to 10 days enhanced the effect even at low doses (10 μM). The MTT IC₅₀ calculated for cell viability showed a substantial agreement among the MTS and cell count experiments (10–20 μM range), indicating that at the circulating therapeutic window of 14–20 mg/l, MTT is largely exerting its ACC cytotoxic effect.

Figure 9
MTT induces a decrease in oxygen consumption in H295R cell-isolated mitochondria. (A) Representative experiment of the effects of 48 h of MTT treatment (10, 30, and 50 μM) on oxygen consumption. Traces represent oxygen consumption during 5 min of respiration live mitochondria (1 mg of protein/ml) isolated from the different samples. The arrow indicates the time point when mitochondria were challenged with 5 mM succinate. Respiration was monitored for 5 min at 37 °C. (B) Oxygen consumption expressed as mean fold increase ± S.E.M. vs the controls (untreated cells) in four independent experiments. Statistical analysis was performed with ANOVA followed by Dunnett’s test: *P < 0.05 and **P < 0.001 vs the controls. In controls, endogenous oxygen consumption was 22 ± 8 nmol oxygen/ml, whereas it was 29 ± 8 nmol oxygen/ml following succinate challenge.
cristae and calculated the ratio of the mitochondrial area to the total length of cristae in individual mitochondria of the analyzed cells. This parameter was closely related to the well-known morphological aspects of mitochondrial functions, i.e. number of cristae and density of the inner matrix, yielding more detailed information than other criteria adopted in previous studies, such as the percentage of the visible mitochondrial section displaying intact cisternae (Cadwell et al. 2008).

The effects of MTT are evident in both ACC cell lines, whereas in non-tumoral cells a slight but non-significant inhibition of viability can be detected only after long exposure with high doses. These findings suggest a specificity of MTT for adrenal cancer, which may justify that the side effects, although variable, seem to be restricted to gastric absorption and neurotoxicity.

The apoptotic images evidenced during electron microscopy have been confirmed by the cytofluorimetric analysis, demonstrating how MTT induces early (annexin V exposure at plasma membrane) as well as late apoptosis (caspase 3/7 activation) events.

The MTT-induced alterations in mitochondrial morphology are accompanied by a correspondent functional damage, as demonstrated by a progressive depolarization of mitochondrial membrane potential, finally resulting in the disruption of the respiratory machinery. In order to evaluate the effect of MTT on respiration in live mitochondria avoiding the detection of the toxic effect on the organelle, we measured oxygen consumption in the mitochondrial fraction after normalization for the number of mitochondria. In the treated cells, we observed a net reduction of oxygen consumption in the mitochondria-rich fraction (nearly 80% inhibition with 50 µM MTT), which seems to be mainly due to a damage of the mitochondrial outer membrane (a significant decrease in the expression of VDAC is evident) rather than to a specific derangement of the respiratory chain complexes (stable levels of SDH). MTT has been previously demonstrated to downregulate the expression of proteins involved in cell redox potential, cholesterol trafficking, and stress response as well as in the maintenance of mitochondrial integrity (Stigliano et al. 2008).

Reduction in the expression of VDAC1 following MTT treatment may not be only a passive consequence of the disruption of the mitochondrial ultrastructure and increased permeability of the outer membrane. Indeed, VDAC1, a voltage-dependent anion channel of outer membrane involved in regulation of metabolic and energy functions of mitochondria, has been more recently described as a key player in mitochondria-mediated programmed cell death and apoptosis (Zaid et al. 2005). Tumors highly overexpress VDAC, promoting hexokinase interaction and stimulation of cancer cell glycolysis (Mathupala & Pedersen 2010). Downregulation of the expression of VDAC1 inhibits tumor cell growth both in vitro and in animal models (Koren et al. 2010), thus representing a novel potential anti-cancer target in tumor (Mathupala & Pedersen 2010). VDAC1 also stabilizes StAR at the mitochondrial outer membrane, enabling the
transport of cholesterol to the inner membrane to be converted to pregnenolone (Bose et al. 2008). Thus, reduction in the expression of VDAC1 may contribute to the steroidogenesis block observed in adrenal cancer cells and in ACC patients treated with MTT.

In conclusion, our findings contribute to better understand the intracellular mechanism of the action of MTT. We demonstrated that the drug significantly affects mitochondria in ACC cells and quantified the mitochondrial damage caused by MTT used at therapeutic doses. We also demonstrated that the cytotoxic effect of MTT implies the activation of the apoptotic process. Whether MTT has similar actions in the correspondent normal adrenocortex of patients under MTT treatment needs further investigation.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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