PRO-APOPTOTIC ACTIVITY OF NOVEL ISATIN-SCHIFF BASE COPPER(II) COMPLEXES DEPENDS ON OXIDATIVE STRESS INDUCTION AND ORGANELLE-SELECTIVE DAMAGE.

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We characterized the pro-apoptotic activity of two new-synthesized isatin-Schiff base copper(II) complexes, obtained from isatin and 1,3-diaminopropane or 2-(2-aminoethyl)pyridine: the [Bis(2-oxindol-3-ylimino)-1,3-diaminopropane-N,N',O,O']copper(II), Cu(isapn), and the Bis[(2-oxindol-3-ylimino)-2-(2-aminoethyl)pyridine-N,N']copper(II), Cu(isaepy)2. We demonstrated that these compounds trigger apoptosis via the mitochondrial pathway. The early induction of the p53/p21 system indicates a role for p53 in cell death, however experiments carried out with siRNA against p53, or with cells lacking p53, support that a p53-independent mechanism can also occur. The extent of apoptosis mirrors the kinetics of intracellular copper uptake. Particularly, Cu(isaepy)2 enters the cells more efficiently and specifically damages nuclei and mitochondria, as evidenced by atomic absorption analysis of copper content and by the extent of nuclear and mitochondrial integrity. Conversely, Cu(isapn), although less permeable, induces a widespread oxidative stress, as demonstrated by analyses of ROS concentration, and oxidation of proteins and lipids. The increase the antioxidant defense, through the overexpression of Cu,Zn SOD partially counteracts cell death; whereas retinoic acid-mediated differentiation completely rescues cells from apoptosis induced by both compounds. The activation of JNK- and Akt-mediated phosphorylative pathways has been found to be not functional for apoptosis induction. On the contrary, apoptosis significantly decreased when the analogous zinc complex was used or when Cu(isapn) was incubated in the presence of a copper chelator. Altogether, our data provide evidence for a dual role of these copper(II) complexes: they are able to vehicle copper into the cell, thus producing ROS and could behave as delocalized lipophilic cations (DLCs)-like molecules, thus specifically targeting organelles.

INTRODUCTION

In the last years synthesis and characterization of novel anti-tumor compounds have represented a field of research which has aroused expectations for more specific and less toxic therapies. Besides DNA and cellular replication, which so far represented the principal targets of cancer treatment, other intracellular compartments and other cell functions, as well as the microenvironment of cancer cells, have become the targets of new and more specific therapies (1-6). For instance, tumor cells are known to show different redox sensitivity or to have low levels of antioxidants, which may lead to an increase of radical species. This phenomenon represents a double-edged sword: on one hand it allows the activation of several redox-sensitive transcription factors inducing tumor proliferation and cell cycle progression (7-10); on the other it can represent an important tool in selectively inducing apoptosis (11-14). Many chemical agents still used in chemotherapy are exploiting this feature; in fact they easily undergo one-electron redox cycling with oxygen, giving rise to superoxide production and oxidative insult. Doxorubicin, daunorubicin, and bleomycin, are among the most used and well-known examples of such chemotherapeutics (15-17), but recently particular attention has been addressed also to transition metals, such as copper (18, 19).
Copper is a micro-nutrient essential for cell survival because it functions as cofactor of several metallo-enzymes (e.g. Cu,Zn SOD and cytochrome c oxidase), but it is also toxic when present at high concentrations (20). In fact, existing in two redox states, Copper(I) and Copper(II), it represents an excellent catalyst of redox cycles in the presence of oxygen (21), generating partially reduced and highly reactive O₂ derivatives, the so-called “reactive oxygen species” (ROS). Besides their well known detrimental effects, ROS can also act as second messengers, which, depending on their concentration and the protein target involved, can trigger different signal transduction pathways ultimately leading to cell survival or death response (22).

Since the clinical success of cisplatin for the treatment of several tumor cell types has been amply demonstrated, other metal-based compounds have been tested for their anti-tumor activity, in order to discover more effective and less toxic drugs than cisplatin (23, 24). In this context copper has long history of medical application, however its potential anti-tumor properties have been explored only in the last few decades (25). Anti-tumor activity of copper thiosemicarbazone complexes was reported as early as the 1960s (26), however, although many copper-based anti-tumor agents induced cell death in vitro, their further use was limited, due the low water solubility and relatively high toxicity in vivo (26-28).

We previously synthesized novel isatin-Schiff base copper(II) complexes and characterized their chemical, physical and biological properties, suggesting a potential role of some of them in the activation of the apoptotic program in different tumor cell lines (29, 30). In this study, we have investigated the molecular mechanisms underlying the activation of apoptosis upon treatment with two specific isatin-diimine copper(II) complexes: Cu(isapn), and Cu(isaepy), in SH-SY5Y neuroblastoma cells, demonstrating that their pro-apoptotic activity involves the mitochondrial pathway. Owing to the different copper coordination characterizing the two complexes, the molecular events upstream of the execution of apoptosis are peculiar of each compound and seem to be tightly associated with the relative kinetics of copper uptake inside the cells.

**EXPERIMENTAL PROCEDURES**

**Materials** – Isatin-diimine copper(II) complexes [Bis(2-oxindol-3-ylimino)-1,3-diaminopropane-N,N',O,O' ]copper(II) perchlorate, [Cu(isapn)][ClO₄]₂, and Bis[(2-oxindol-3-ylimino)-2-(2-aminoethyl)pyridine-N,N']copper(II) perchlorate, [Cu(isaepy)₂][ClO₄]₂, named here Cu(isapn) and Cu(isaepy)₂, respectively (Fig. 1 A and 1 B), were synthesized as previously described (30). The analogous isatin-imine zinc(II) complex [Zn(isaepy)Cl₂], designated as Zn(isaepy), was prepared similarly using zinc chloride to metallate in situ the isaepy ligand. (Fig. 1 C). Dimethyl sulfoxide (DMSO), dithiothreitol (DTT), EDTA, EGTA, paraformaldehyde, propidium iodide, tert-butyl hydroperoxide, sodium ortovanadate, triethylenetetramine (TRIEN), and Triton X-100 were from Sigma (St. Louis, MO). Goat anti-mouse and anti-rabbit IgG (H+L)-horseradish peroxidase conjugate was from Bio-Rad Lab. (Hercules, CA). N-Tris(hydroxymethyl)methyl-2-aminoethanesulfonic acid (TES) was from US Biological (Cleveland, OH). All other chemicals were obtained from Merck (Darmstadt, Germany).

**Cell culture** – Human neuroblastoma cells SH-SY5Y were purchased from the European Collection of Cell Culture and grown in Dulbecco’s MEM-F12 medium. Promonocytoma U937 were from the American Type Culture Collection; melanoma M14 were kindly provided by Dr. Gabriella Zupi from the Experimental Chemotherapy Laboratory, Regina Elena Cancer Institute of Rome, and grown in RPMI 1640 medium; cervical carcinoma HeLa cells stably transfected with pSUPER vector containing p53 siRNAs (pSUPER-p53), or with empty vector (pSUPER) were a gift of Dr. Anna Maria Biroccio from the Experimental Chemotherapy Laboratory, Regina Elena Cancer Institute of Rome and grown in DMEM. All cell media were supplemented with 10% fetal calf serum (FCS), and the cells were grown at 37 °C in an atmosphere of 5% CO₂ in air. Monoclonal SH-SY5Y cell lines transfected with human wild type Cu,Zn SOD (named hSOD) were obtained as previously described (31). During the experiments cells were plated at a density of 4 × 10⁵/cm² (for SH-SY5Y, HeLa and M14) or 2 × 10⁵/ml (for U937), unless otherwise indicated.

**Treatments** – A 5 mM solution of Cu(isapn), Cu(isaepy)₂ or Zn(isaepy) was prepared just before the experiments by dissolving the lyophilized compounds in DMSO. Treatments were performed with a concentration of 50 μM...
at 37 °C in medium supplemented with serum. This concentration was chosen for all the experiments because it gave a substantial degree of apoptosis at the times selected (29). As control, equal volumes of DMSO (1%) were added to untreated cells. The pancaspase inhibitor zVAD-fmk (Alexis Biochemicals, CA) was used at a final concentration of 100 mM, pre-incubated for 1 h before the addition of Cu(isapn) and Cu(isaepy)_2, and maintained throughout the experimental time. TRIEN, at a final concentrations of 150 μM, was added 3 h before the addition of Cu(isaepy)_2 and maintained throughout the experiment. A 5 mM solution of copper sulfate in water was prepared just before the experiments and added to culture media at a concentration of 50 μM. Treatments with the specific c-Jun-N-terminal kinase (JNK) inhibitor, SP600125 (Calbiochem-Novabiochem, La Jolla, CA), and the phosphoinositide-3-kinase (PI3K)/Akt pathway inhibitor wortmannin (Calbiochem-Novabiochem) were performed at concentrations of 10 μM and 5 μM, respectively, since under our experimental conditions they did not result to be toxic. They were added 30 min before the addition of Cu(isapn) or Cu(isaepy)_2 and maintained throughout the experiment. Retinoic acid (RA) was added to culture media at a concentration of 20 μM and maintained for seven days of culture in order to allow differentiation of cells, which was monitored by analyzing the increased expression of the differentiation marker, growth-associated protein-43 (GAP-43).

Analysis of cell viability and apoptosis – Adherent (after trypsinization) and detached cells were combined, washed with PBS and stained with 50 μg/ml propidium iodide prior to analysis by a FACScalibur instrument (Becton Dickinson, San José, CA). The percentages of apoptotic cells were evaluated according to Nicoletti et al. (32) by calculating peak area of hypodiploid nuclei (SubG1). Alternatively, cells were collected and counted after Trypan Blue staining by optical microscopy using a Thoma’s chamber.

Cell fractionation and protein extraction – Total protein extracts were obtained by rupturing cells with 30 min of incubation on ice in lysis buffer (50 mM Tris-HCl, pH 7.4, 1 mM EDTA, 1 mM EGTA, 1% Triton X-100, 10 mM NaF, 1 mM sodium orthovanadate) and protease inhibitor cocktail (Roche Applied Science, Monza, Italy) and centrifuged at 22,300 × g for 20 min at 4°C. Cell fractions were obtained as previously reported (33). Briefly, cells were incubated in hypotonic medium (10 mM Tris-HCl, pH 7.5, 15 mM MgCl_2, 10 mM KCl, and protease inhibitor cocktail). After 10 min of incubation on ice, equal volumes of “mitochondrial” buffer containing 400 mM sucrose, 10 mM TES, 0.1 mM EGTA and 2 mM DTT was added, and cells ruptured by 40 strokes in a glass Dounce. Pellets obtained after centrifugation of lysates at 900 × g were considered nuclear-enriched fractions, whereas supernatants further centrifuged at 10,000 × g produced a mitochondria- (pellet) and cytosol-enriched fraction (supernatant) For purity determination, total cell extracts and each fraction were analyzed by Western blot against: the α subunit of cytochrome c oxidase (CytOx) and the 39 kDa subunit of Complex I (specific for mitochondria); Cu,Zn SOD (mainly present into the cytosol); Lamin A/C (specific for nuclei) (see supplemental data, Fig. S1). For nuclear phospho-H2A.X determination, nuclear extracts were obtained by lysing the cells in nucleus buffer (1 mM K_2HPO_4, pH 6.4, 150 mM NaCl, 14 mM MgCl_2, 1 mM EGTA, 0.1 mM DTT and 0.3% Triton-X100), and the nuclear fractions obtained were further lysed with lysis buffer.

Western Blot analyses – Protein extracts were electrophoresed by SDS-PAGE and blotted onto nitrocellulose membrane (Bio-Rad). Polyclonal anti-caspase-9, anti-phospho-p42/44 (thr202/tyr204) (Cell Signaling Technology, Beverly, MA), anti-phospho-Akt1/2/3, anti-poly-ADP ribose polymerase (PARP), anti-p21, anti Cu, Zn SOD (Santa Cruz Biotechnology, Santa Cruz, CA), and monoclonal anti-caspase-3 (clone 3G2), anti-phospho-p38 (thr180/tyr182) (Cell Signaling Technology), anti-p53 (clone BP5312), anti-α-tubulin (clone DM1A) (Sigma), anti-Lamin A/C (UCS Diagnostics, Rome, Italy), anti-phospho-JNK (G7) (Santa Cruz), anti-phospho-H2A.X (ser-139) (clone JBW301) Upstate Biotechnology, Lake Placid, NY), anti-CytOx (α-subunit) and anti-39 kDa subunit of Complex I (Invitrogen-Molecular Probes, San Giuliano Milanese, Italy) were used as primary antibodies. The specific protein complex, formed upon specific secondary antibody treatment, was identified using a Fluorchem Imaging system (Alpha Innotech, Analitica De Mori, Milano, Italy) after incubation with ChemiGlowl chemiluminescence substrate (Alpha Innotech).

Measurement of glutathione, ROS and oxidative damage – Intracellular reduced (GSH) and oxidized (GSSG) forms of the tripeptide glutathione were assayed upon formation of S-carboxymethyl derivatives of free thiols with
iodoacetic acid, followed by the conversion of free amino groups to 2,4-dinitrophenyl derivatives by the reaction with 1-fluoro-2,4-dinitrobenzene as previously described (34). Detection of intracellular ROS by 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA, Invitrogen-Molecular Probes), analyses of protein carbonyls content as well as malondialdehyde (MDA) and 4-hydroxynonenal (4-HNE) levels were performed as previously described (35).

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**Fluorescence microscopy analyses –** Cells were plated on chamber slides at 6 × 10⁴/cm², fixed with 4% paraformaldehyde and permeabilized. Afterward, they were washed exhaustively with PBS, blocked with PBS containing 10% FCS and incubated with: a) monoclonal anticytchrome c antibody (clone G742A) (PROMEGA, Madison, WI) and polyclonal anti-AIF antibody (Chemicon International, Temecula, CA). Cells were then washed with PBS and probed with an Alexa fluor®-488 goat anti-mouse and an Alexa fluor®-568 conjugated goat anti-rabbit secondary antibodies (1:1000) (Invitrogen-Molecular Probes), then analyzed by fluorescence microscopy. b) Monoclonal anti-ser-139-phosphorylated histone H2A.X antibody, and further probed with an Alexa fluor®-488 goat anti-mouse secondary antibody (1:1000) (Invitrogen-Molecular Probes). To visualize nuclei, cells were also incubated with propidium iodide solution, washed with PBS and analyzed by fluorescence microscopy. To evaluate mitochondrial integrity, cells were stained with 50 nM of the mitochondrial transmembrane potential-sensitive probe MitoTracker Red® (Invitrogen-Molecular Probes), washed and then fixed with 4% paraformaldehyde. To determine the shape of nuclei, cells were also incubated with Hoechst 33342 (1:1000, Calbiochem-Novabiochem), and visualized by fluorescence microscopy. Images of cells were digitized with a Cool Snap video camera connected to Nikon Eclipse TE200 fluorescence microscopy. All images were captured under constant exposure time, gain and offset.

**Copper determination –** Cell pellets were diluted 1:2 with 65% HNO₃. After 1 week at room temperature, copper concentration was measured by atomic absorption spectrometry using an AAAnalyst 300 Perkin-Elmer instrument, equipped with a graphite furnace with platform HGA-800 and an AS-72 auto sampler. Concomitantly, cell media were separated from pellets and analysed by EPR spectroscopy. EPR spectra were recorded using 80 μl samples in flat glass capillaries (inner cross section 5 × 0.3 mm) to optimize instrument sensitivity. All measurements were made at 298 K with an ESP300 X-band instrument (Bruker, Karlsruhe, Germany) equipped with a high sensitivity TM₁₀₀-mode cavity. Spectra were measured over a 1000 G range using 50 mW power, 10 G modulation and a scan time of 42 s; normally 24 single scans were accumulated to improve the signal to noise ratio.

**siRNA transfections –** Twenty-four hours after plating, 50% confluent SH-SY5Y cells were transfected, with a 21-nucleotide siRNA duplex directed against the p53 mRNA target sequence, 5'-GACUCAGUGGAUUCUACTT-3' (sip53) (MWG Biotech, Ebersberg, Germany). Control cells were transfected with a scramble siRNA duplex, which does not present homology with any other human mRNAs (siScr). Cells were transfected by electroporation using a Gene Pulser Xcell system (Bio-Rad) according to the manufacturer’s instructions and immediately seeded into fresh medium. Transfection efficiency of siRNA into SH-SY5Y cells was estimated by co-transfecting p53 siRNA with non-specific rhodamine-conjugated oligonucleotides and found to be > 80%.

**RESULTS AND DISCUSSION**

Cu(isapn) and Cu(isaepy)₂ induce cell cycle arrest and caspase-dependent apoptosis – In order to dissect the mechanisms by which isatin-Shiff base copper(II) complexes induce cell death in tumor cell lines, on the basis of the results previously obtained (29), we selected the most effective molecules, Cu(isapn) and Cu(isaepy)₂ (Fig. 1 A and B), and used them at a concentration of 50 μM. Fig. 2A shows representative histograms from cytofluorimetric analyses of SH-SY5Y cells treated with Cu(isapn) and Cu(isaepy)₂ for 24 and 48 h, where a more effective increase in the apoptotic cells with Cu(isaepy)₂ treatment was evidenced (Fig. 2A). Western blot analyses of p53 and p21, showed a strong activation of these two proteins although with different kinetics: rapid for...
Cu(isaepy)$_2$ and more gradual for Cu(isapn) with a peak of induction at 12 and 24 h, respectively (Fig. 2B), indicating that the p53/p21 pathway is activated as response to cell damage. The typical mitochondrial localization of the apoptosis inducing factor (AIF) and endonuclease G showed that no “caspase-independent” mechanism was operative under our experimental conditions (data not shown); searching for the mechanism underlying apoptotic induction, we therefore used the anti-AIF antibody to probe mitochondria. As evidenced by immuno-fluorescence analyses of SH-SY5Y cells treated for 24 h with copper complexes, AIF fluorescence did not superimpose cytochrome c staining (Fig. 2C), indicating that cytochrome c was efficiently released from mitochondria into the cytosol, and that the intrinsic mitochondrial pathway was operative under our experimental conditions. Concomitantly, Western blot analyses of pro- and active caspase-9, as well as pro-caspase-3 and poly-ADP ribose polymerase (PARP) indicated that each step of the apoptotic program was executed upon treatment with either Cu(isapn) or Cu(isaepy)$_2$ (Fig. 2D). In fact, immuno-reactive bands of proteolyzed caspase-9 and PARP, together with a significant decrease of pro-caspase-3 were already detected after 24 h of treatment. As final evidence that a caspase-dependent apoptotic response occurred after treatment with both copper complexes, we pre-incubated the cells with 100 μM of the pancaspase inhibitor zVAD-fmk for 1 h and then added Cu(isapn) or Cu(isaepy)$_2$. Fig. 2E shows that a recovery of cell viability resulted from the inhibition of caspases activation upon treatment with both compounds, confirming that caspase-mediated apoptosis is the principal mechanism for cell death induction in our experimental system.

To evaluate the role of p53 in apoptosis induction, we transfected SH-SY5Y cells with siRNA against p53 (sip53) or with a scramble sequence which does not present homology with any other human mRNAs (siScr). Western blot analyses of p53 levels indicated that the concentration of the protein rapidly decreased after transfection (see Fig. 3B, upper panel). Therefore, we decided to treat the cells with Cu(isapn) or Cu(isaepy)$_2$ after 9 h from transfection, time which allowed the cells adhering to the flask and p53 interference being still in progress. Cytofluorimetric analyses show that sip53 cells were significantly resistant to apoptosis if compared to siScr (Fig. 3A). This was also confirmed by Western blot of PARP which indicated a more efficient cleavage of the protein occurring in siScr than in sip53 cells (Fig. 3B, bottom panel). We also performed experiments on HeLa cells stably transfected with a pSUPER vector containing siRNAs for p53 (pSUPER-p53) or with an empty vector (pSUPER). Fig. 3C shows cytofluorimetric analyses of apoptosis upon 24 h treatment with Cu(isapn) or Cu(isaepy)$_2$. As previously observed for SH-SY5Y, also for HeLa, the percentage of subG1 cells was significantly reduced upon p53 interference. This result was further confirmed by Western blot analyses of PARP (Fig. 3D), suggesting that p53 activation contributes to copper complexes-mediated apoptosis. Nevertheless, a considerable amount of p53-independent apoptosis was also observed.

Cu(isapn) and Cu(isaepy)$_2$ cross the cell membrane and selectively induce oxidative stress – To characterize the capability of Cu(isapn) and Cu(isaepy)$_2$ to enter the cells and the kinetics of their accumulation, we followed copper uptake by atomic absorption analyses. Fig. 4A shows that treatments with both compounds resulted in a rapid increase of intracellular copper content which reached a plateau after 12 h. This result was particularly significant, especially when compared with that obtained with copper sulfate, used as control of cellular incorporation of the metal ion. Cu(isaepy)$_2$ seems to be more efficiently incorporated within the cells with respect to Cu(isapn). These results demonstrated a direct relationship between copper uptake and the extent of apoptosis, with Cu(isaepy)$_2$ being more permeating and more efficient in inducing cell death than Cu(isapn). Since the molecules we have synthesized have characteristic EPR spectra (23), we measured the extracellular concentration of Cu(isaepy)$_2$ by EPR spectroscopy. Fig 4B reports the spectra of Cu(isaepy)$_2$ in cell culture media and its relative concentration up to 12 h of treatment. The data suggest that a decrease in the content of copper complex, most probably due to its uptake by cells, was operative under our experimental conditions. Moreover, these results also suggested that the complex was chemically stable during the experimental times selected, as no changes of the EPR parameters were evidenced. To further confirm both hypotheses, we performed the same experiments at 4°C, temperature at which metabolic processes are strongly reduced. Fig. 4C show that, under these
Cu(isaepy)$_2$ were pro-oxidants, as ROS generation in a cell system, we measured copper uptake, suggesting that the levels of glutathione, due to its ability to bind copper and detoxify xenobiotics, might be coupled with the intracellular content of copper compounds.

**Cu(isapn) and Cu(isaepy)$_2$-mediated cytotoxicity does not depend on phosphorylative pathways**

To determine if apoptosis was mediated by phosphorylative pathways downstream of oxidative stress produced by Cu(isapn) and Cu(isaepy)$_2$, we performed Western blot analysis of phospho-active levels of the different members of mitogen activated protein (MAP) kinases and protein kinase B, also known as Akt. No significant changes in the immunoreactive bands of active p38$^{\text{MAPK}}$ and ERK1/2 were detected upon treatment with the copper complexes (data not shown), but the phospho-isoforms of JNK and Akt increased significantly, particularly after Cu(isaepy)$_2$ treatment (Fig. 6A), suggesting a role for these proteins in the induction of the cell death program. However, since this increase occurred very early after treatment (3-6 h), no correlation between Akt or JNK activation and ROS production can be stated, as far as Cu(isaepy)$_2$ treatment is concerned. To determine the role of Akt in cell response to Cu(isapn) and Cu(isaepy)$_2$, we pre-incubated SH-SY5Y cells with wortmannin, a specific inhibitor of the Akt upstream kinase PI3K, and then treated the cells with the copper complexes. Fig 6B shows that inhibition of Akt before treatment with Cu(isapn) and Cu(isaepy)$_2$ resulted in a significant increase of subG1 cell population. Western blot analyses of cell lysates confirmed that wortmannin incubation produced blockage of the Akt-mediated pathway (Fig. 6C) and concomitantly showed no effect on the phospho-active form of JNK after Cu(isapn) and Cu(isaepy)$_2$ treatments, suggesting that Akt and JNK activation are independent responses. Pre-incubation with the specific JNK pathway inhibitor SP600125 did not result in modified cell survival to treatments with either Cu(isapn) or Cu(isaepy)$_2$ (Fig. 6D), indicating that JNK activation could represent just an epiphenomenon of the copper complexes-induced cytotoxicity, which does not directly relate to the apoptotic process.

**Increase of antioxidant defense or differentiation by retinoic acid rescues SH-SY5Y cells from Cu(isapn) and Cu(isaepy)$_2$-induced cytotoxicity**

– The results so far presented indicate a different time dependency and occurrence of the oxidative markers upon treatments with Cu(isapn) or Cu(isaepy)$_2$, which could reflect the different
structures of the copper complexes used. Particularly, the former could represent a more efficient ROS producer and the latter a more toxic and permeating compound. To evaluate the contribution of oxidative stress in the execution of apoptosis, we used SH-SY5Y cells transfected with an additional copy of the human SOD1 gene (hSOD cells) or differentiated by means of seven days of incubation with 20 μM retinoic acid (RA). Fig. 7A shows the results obtained by direct counts of dead cells after Trypan Blue staining, and demonstrates that hSOD cells were more resistant to both compounds than parental SH-SY5Y, even though to different extents. The increased anti-oxidant defense let hSOD cells to be highly resistant to Cu(isapn)-induced cytotoxicity, whereas a moderate decrease in dead cells upon Cu(isaepy)₂ treatment was observed at 48 h. These results give support to a different mode of action of the two copper complexes, in which ROS production represents a crucial upstream event in apoptosis induced by Cu(isapn), but only plays a downstream additional role in Cu(isaepy)₂ cytoxicity. The p53/p21 pathway was also activated upon both treatments, although with a different kinetics. In fact, p53 and p21 seemed highly expressed after 6 and 12 h of treatments with Cu(isapn) and Cu(isaepy), respectively; but no activation was observed after 24 h (Fig. 7B). These results suggest that Cu(isapn) and Cu(isaepy)₂ exert similar inhibitory effects on cell growth of parental SH-SY5Y and hSOD cells, a result confirmed by cell direct counts of attached viable cells (data not shown). To test whether such compounds functioned as non-specific toxic agents, or were less efficient towards untransformed cells, we induced differentiation with prolonged RA incubations. The results obtained would have provided significant information about a potential approach for a selective use of copper complexes in cancer treatment. Seven days incubation with 20 μM RA conferred efficient protection against both copper complexes. A significant decrease of subG1 cells to values close to untreated cells (about 4.25% after 48 h as shown in Fig. 7C) and no activation of the p53/p21 system was observed under these experimental conditions (Fig. 7D). The average one fold-increase of superoxide dismutase and catalase activity (data not shown), together with the inhibition of cell proliferation, features found in normal (untransformed) cells, are some of the modifications induced by RA incubation and may contribute to cell resistance against Cu(isapn) and Cu(isaepy)₂-mediated toxicity.

Cu(isapn) and Cu(isaepy)₂-mediated toxicity is a copper mediated event which induces nuclear and mitochondrial dysfunction – Cu(isapn) and Cu(isaepy)₂ are lipophilic compounds which may vehicle copper and allow the catalysis of redox reactions and ROS production to take place also within specific intracellular organelles. We evaluated the copper content in cellular fractions of SH-SY5Y cells (the purity of which is shown in supplemental data, Fig. S2) treated for different times with the copper complexes. Fig 8A shows atomic absorption analyses, which reveal a significant increase of copper in nuclear and mitochondrial fractions upon treatment with both compounds, with a trend mirroring total copper uptake and suggesting a likely copper-mediated site-directed injury. To evaluate the degree of the insult induced, we investigated if mitochondria and nuclei were damaged upon treatment with the two copper complexes. Fig. 8B shows SH-SY5Y nuclei stained with an antibody against the phospho-active histone H2A.X, which is phosphorylated on Ser139 after DNA double strand break. After 6 h and 12 h of treatment, the appearance of discrete foci, indicating the recruiting sites of the DNA repair machinery, revealed DNA-specific damage mediated by Cu(isaepy)₂ and, to a lesser extent by Cu(isapn) treatment, a phenomenon that was further confirmed by Western blot analyses of nuclear fractions (Fig. 8C). Concomitantly, we incubated SH-SY5Y cells with MitoTracker Red®, a red-fluorescent dye that stains mitochondria in viable cells and which accumulates depending upon transmembrane potential (ΔΨₘ). Fig. 8D shows that especially for Cu(isaepy)₂ treatment, a huge number of mitochondria resulted completely depolarized 12 h after treatment; ΔΨₘ-dependent fluorescence was localized only in single spots and did not look like a continuous network as in untreated cells. These results give strength to the hypothesis that although Cu(isaepy)₂ does produce less ROS, it is more efficient than Cu(isapn) in inducing apoptosis, owing to its ability to specifically accumulate and damage fundamental organelles, such as the nucleus and mitochondria.

To evaluate further the contribution of copper in Cu(isaepy)₂-induced apoptosis, we synthesized an analogous complex molecule in which zinc, a non-redox active metal ion, replaced copper in the complex. Alternatively,
before Cu(isaepy)₂ administration, we incubated the cells with 150 µM of the non-permeating copper chelator TRIEN, which is able to remove copper from the complex and to prevent its uptake by the cells. Cytofluorimetric analyses shown in Fig. 8E demonstrate that either upon treatment with Zn(isaepy) (Fig. 1C), or by pre-incubating the cells with TRIEN, the extent of subG1 cell population strongly decreased with respect to Cu(isaepy)₂-treated cells, although a significant percentage of apoptosis was still detectable (n = 6, p < 0.05 with respect to untreated cells). These results suggest that copper plays a fundamental role in Cu(isaepy)₂-mediated apoptosis, presumably due to its action as catalyst of redox reactions; however, the isatin-Schiff base could also contribute to the toxicity observed.

Cu(isapn) and Cu(isaepy)₂ induce apoptosis via the mitochondrial pathway in other tumor cell types – Finally, to verify a general pro-apoptotic activity of the more efficient copper complex Cu(isaepy)₂, we selected two other human tumor cells, the promonocytoma U937 and the melanoma M14. As previously done with SH-SY5Y cells, we treated U937 and M14 cells with 50 µM Cu(isaepy)₂. Fig 9A shows representative cytofluorimetric panels which demonstrate that treatment with this copper complex induced apoptosis after 24 h in both the tumor cell lines selected, with U937 cells being more susceptible. Western blot analyses of caspase-9, caspase-3 and PARP, performed at 24 and 48 h, further confirmed that the intrinsic mitochondrial pathway represents the preferential route for the induction of apoptosis (Fig. 9B). These results demonstrate the ability of Cu(isaepy)₂ in inducing cell death in different tumor cells, and suggest a general application of the molecule.

CONCLUSIONS

In this paper we report that two recent synthesized isatin-Schiff base copper(II) complexes Cu(isapn) and Cu(isaepy)₂ are able to induce apoptosis via the mitochondrial pathway in neuroblastoma SH-SY5Y cells and in other tumor histotypes, mainly by copper-dependent oxidative stress and nuclear/mitochondrial site-directed damage. Although both compounds are capable of producing such effects, the difference in the time of appearance of pro-apoptotic and oxidative markers and the extent of cell death depends on their efficiency to permeate the cell. Cu(isapn), although less permeating, seems more prone to produce oxyradicals and induce oxidative stress, whereas Cu(isaepy)₂ easily crosses cell membranes, accumulates and damages nuclear and mitochondrial compartments at very early times. In line with these features, we speculate that the absence of a detectable “cytosolic” oxidative stress upon treatment with Cu(isaepy)₂ explain the absence of a critical redox-activation of the JNK-mediated signaling cascade in the events leading to cell death.

Moreover, our data are equally supportive of parallel p53-dependent and independent pathways, as demonstrated by experiments carried out in p53 knocked-down cells by siRNA or in cell lines lacking p53. These results are of great interest since functional p53 is frequently lost in human tumorigenesis. The results obtained indicate that the role of copper in Cu(isapn) and Cu(isaepy)₂ cytotoxicity is fundamental, because of its capability to catalyze one-electron redox cycle reactions with oxygen thus producing ROS; the combination of this property with the chemical structure of the organic ligand binding the metal ion seems to give the specificity of the cellular damage induced. We suggest that the presence of copper is mainly required for nuclear damage, similar to what iron do in the presence of bleomycin (37, 38), whereas the isatin-imine ligand is important to carry the redox-active metal ion across cellular membranes, and could be more active at the mitochondrial level. This hypothesis is strengthened by the similarity of the chemical structure of Cu(isapn) and Cu(isaepy), with the delocalized lipophilic cations (DLCs), a class of molecules able to permeate the cell in response to negative transmembrane potentials and increase their concentration, particularly into mitochondria (39). The higher plasma and mitochondrial membrane potentials of tumor cells compared to normal cells account for the preferential accumulation of DLCs in carcinoma mitochondria (40, 41). Since most DLCs are toxic to mitochondria at high concentrations, their selective accumulation in the mitochondria of tumor cells, and consequent mitochondrial toxicity, provide the basis for selective tumor cell killing. The capability of this compounds to preferentially induce detrimental effects in transformed cells can been also suggested on the basis of the preliminary results obtained in the presence of RA, which is able to protect the cells by making them similar to differentiated lines.

Overall, the results obtained allow us to suggest a dual role for Cu(isapn) and
Cu(isaepy)₂ in the induction of apoptosis: on one hand they are able to vehicle copper into the cell, thus producing ROS; on the other they could behave as DLCs, thus specifically targeting mitochondria. Therefore the chemical structure of the isatin-Shiff base represents the “switch” between these properties. This suggests that by specifically changing the chemical characteristics of this ligand type, we may modulate the cytotoxic effects induced, thus exploiting the plasticity of this new class of compounds to improve the therapeutic selectivity to different tumor histotypes.

REFERENCES


**FOOTNOTES**

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#Recipient of the fellowship “Santina Grillini” from Italian Association for Cancer Research (AIRC-FIRC).

1The abbreviation used are: AIF, apoptosis inducing factor; Cu(isaepy)_2, Bis[(2-oxindol-3-ylimino)-2-(2-aminoethyl)pyridine-N,N’]copper(II); Cu(isapn), [Bis(2-oxindol-3-ylimino)-1,3-diaminopropane-N,N’,O,O’]copper(II); Cu,Zn SOD, copper, zinc superoxide dismutase; DCFH-DA, 2’,7’-dichlorodihydrofluorescein diacetate; DLCs, delocalized lipophilic cations; DMPO, 5,5-dimethyl-1-pyrroline N-oxide; EPR, electron paramagnetic resonance; ERK1/2, extracellular regulated kinase 1 and 2; GSH, reduced glutathione; GSSG, oxidized glutathione; JNK, c-Jun-N-terminal kinase; MAPK, mitogen activated protein kinase; PARP, poly-ADP ribose polymerase; PI3K, phosphoinositide-3-kinase; RA, retinoic acid; ROS, reactive oxygen species; TRIEN, triethylenetetramine; Zn(isaepy), [(2-oxindol-3-ylimino)-2-(2-aminoethyl)pyridine-N,N’]zinc(II); zVAD-fmk, benzoyloxycarbonyl-Val-Ala-Asp-fluoromethyl ketone.

**FIGURE LEGENDS**

Fig. 1. Structures of isatin-Schiff base copper(II) and zinc(II) complexes. A, [Bis(2-oxindol-3-ylimino)-1,3-diaminopropane-N,N’,O,O’]copper(II), [Cu(isapn)]^{2+}, and B, Bis[(2-oxindol-3-ylimino)-2-(2-aminoethyl)pyridine-N,N’]copper(II), [Cu(isaepy)]^{2+}. C, the analogous isatin-imine zinc(II) complex [Zn(isaepy)Cl_2], designated as Zn(isaepy), was prepared similarly using zinc chloride to metallate in situ the isaepy ligand.

Fig. 2. Cu(isapn) and Cu(isaepy)_2 induce a p53/p21-associated apoptosis in SH-SY5Y cells via the mitochondrial pathway. A, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)_2 for 24 and 48 h, washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACS Calibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 12, *p < 0.001. B, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)_2 for 6, 12 and 24 h. 25 μg of total protein extract was loaded onto each lane for detection of p53 and p21. α-tubulin was used as loading control. Western blots are from one experiment representative of three that gave similar results. C, SH-SY5Y cells were grown on chamber-slides, treated for 24 h with 50 μM Cu(isapn) or Cu(isaepy)_2 and concomitantly incubated with antibodies anti-AIF (red), to visualize mitochondria, and anti-cytochrome c (green). Images were digitized with a Cool Snap video camera connected to Nikon Eclipse TE200 fluorescence microscopy. White arrows indicate cells were cytochrome c was not
localized into mitochondria; therefore no superimposition of the two fluorescences was evidenced. D, Alternatively, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 and 48 h. 40 μg of total protein extract was loaded onto each lane for detection of pro- and active-caspase-9, procaspase 3 and PARP. Western blots are from one experiment representative of three that gave similar results. E, SH-SY5Y cells were incubated for 1 h with or without 100 μM pancaspase inhibitor zVAD-fmk, treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24, washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACScalibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 5, *p < 0.001.

Fig. 3. p53 RNA interference partially counteracts apoptosis induced by Cu(isapn) and Cu(isaepy)₂. A, SH-SY5Y cells were transiently transfected with siRNA duplex directed against the p53 mRNA target sequence (sip53) or with a scramble siRNA duplex, which does not present homology with any other human mRNAs (siScr). Cell adhesion has been allowed for 9 h, then the cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for the next 24, washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACScalibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 6, *p < 0.05. B, to evaluate the degree of p53 decrease, at the time points indicated, sip53 or siScr cells were harvested and lysed. 25 μg of total protein extract was loaded onto each lane for detection of p53. α-tubulin was used as loading control. After 9 h from transfection with siScr or sip53, SH-SY5Y cells were then treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h. 40 μg of total protein extract was loaded onto each lane for detection of p53 and PARP. α-tubulin was used as loading control. Western blots are from one experiment representative of three that gave similar results. C, HeLa cells, stably transfected with a pSUPER vector containing p53 siRNA (pSUPER-p53) or with empty vector (pSUPER) were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h, washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACScalibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 6, *p < 0.05. D, pSUPER and pSUPER-p53 cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h. 40 μg of total protein extract was loaded onto each lane for detection of p53 and PARP. α-tubulin was used as loading control. Western blots are from one experiment representative of three that gave similar results.

Fig. 4. Cu(isapn) and Cu(isaepy)₂ induce intracellular copper uptake. A, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 6, 12 and 24 h, exhaustively washed with PBS containing 1 mM EDTA to avoid contamination of extracellularly membrane-bound copper, diluted 1:2 with 65% HNO₃ and analysed for copper content by atomic absorption. 50 μM copper sulfate (CuSO₄) was used as control of copper uptake. Data are expressed as nmol copper/mg total protein and represent the mean ± S.D. n = 5, *p < 0.001. B, SH-SY5Y cells were treated with 50 μM Cu(isaepy)₂ for 12 h at 37 °C. At the same time points protein carbonyls were identified upon derivatization with dinitrophenylhydrazine (DNP) followed by immunoblot using anti-DNP antibody. 20 μg of derivatized proteins were loaded onto each lane. A representative Western blot of three that gave similar results is shown. C, alternatively lipid peroxidation was evaluated by measuring the levels of MDA and 4-HNE using a colorimetric method. Data are expressed as % of control and represent the means ± S.D. n = 5; *p < 0.05, **p < 0.001.

Fig. 5. Cu(isapn) and Cu(isaepy)₂ induce oxidative stress. A, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 3, 6, 12 and 24 h, and incubated with 50 μM DCF-DA at 37 °C. At indicated time points, cells were washed with PBS and ROS production was analyzed by FACScalibur instrument. Histograms shown are representative of three experiments that gave similar results. B, At the same time points protein carbonyls were identified upon derivatization with dinitrophenylhydrazine (DNP) followed by immunoblot using anti-DNP antibody. 20 μg of derivatized proteins were loaded onto each lane. A representative Western blot of three that gave similar results is shown. C, alternatively lipid peroxidation was evaluated by measuring the levels of MDA and 4-HNE using a colorimetric method. Data are expressed as % of control and represent the means ± S.D. n = 5; *p < 0.001.
Fig. 6. JNK and Akt are not involved in Cu(isapn) and Cu(isaepy)₂-induced apoptosis. A, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ up to 24 h. 20 μg of total protein extract was loaded onto each lane for detection of phosphorylated forms of JNK (P-JNK) and Akt (P-Akt). α-tubulin was used as loading control. Western blots are from one experiment representative of three that gave similar results. B, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h previous 1 h incubation with the PI3K/Akt pathway inhibitor, wortmannin. Cells were washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACS caliber instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 5; *p < 0.05, **p < 0.001. C, alternatively 20 μg of total protein extract was loaded onto each lane and used for the detection of phosphorylated forms of JNK (P-JNK) and Akt (P-Akt). Western blots are from one experiment representative of three that gave similar results. D, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h previous 1 h incubation with the specific JNK inhibitor, SP600125. Cells were washed and stained with propidium iodide. Analyses of subG1 (apoptotic) cells was performed by a FACS caliber instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 5.

Fig. 7. The increase of antioxidant defense or differentiation by retinoic acid protect SH-SY5Y cells against Cu(isapn) and Cu(isaepy)₂-induced cytotoxic effects. A, SH-SY5Y and hSOD cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 and 48 h, adherent and floated cells were collected, washed with PBS and counted upon Trypan blue staining. Data are expressed as % of control and represent the means ± S.D. n = 6; **p < 0.001, with respect to parental cell line. B, hSOD cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 6, 12 and 24 h. 25 μg of total protein extract was loaded onto each lane for detection of p53 and p21. Western blots are from one experiment representative of three that gave similar results. C, SH-SY5Y cells were incubated with 20 μM retinoic acid, to induce differentiation, then treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 24 h, washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACS caliber instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 6. D, SH-SY5Y cells were incubated with 20 μM retinoic acid, then treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 6, 12 and 24 h. 25 μg of total protein extract was loaded onto each lane for detection of p53 and p21. Western blots are from one experiment representative of three that gave similar results.

Fig. 8. Copper is fundamental in Cu(isapn) and Cu(isaepy)₂-induced apoptosis and is associated with the specific damage to nuclear and mitochondrial compartments. A, SH-SY5Y cells were treated with 50 μM Cu(isapn) or Cu(isaepy)₂ for 3, 6 and 12 h, and exhaustively washed with PBS containing 1 mM EDTA to avoid contamination of extracellularly membrane-bound copper. Cells were then subjected to separation into nuclear, mitochondrial and cytosolic enriched fractions, subsequently diluted 1:2 with 65% HNO₃, maintained at room temperature for 1 week and analyzed by atomic absorption for copper content. Data are expressed as nmol copper/mg total protein and represent the mean ± S.D. n = 5; *p < 0.05, **p < 0.001, with respect to cytosolic copper concentrations. B, SH-SY5Y cells were grown on chamber-slides, treated for 6 and 12 h with 50 μM Cu(isapn) or Cu(isaepy)₂ and incubated with an anti-phospho histone H2A.X antibody (green), to detect nuclear damage, and propidium iodide (red), to visualize nuclei. Images were digitized with a Cool Snap video camera connected to Nikon Eclipse TE200 fluorescence microscopy. C, alternatively, after 3, 6 and 12 h of treatment, nuclei of SH-SY5Y cells were isolated and lysed. 50 μg of total nuclear extract was loaded onto each lane and used for the detection of phosphorylated form of histone H2A.X. Lamin A/C was used as nuclear loading control. D, SH-SY5Y cells were grown on chamber-slides and treated for 12 h with 50 μM Cu(isapn) or Cu(isaepy)₂. Before fixation, cells were incubated for 30 min with 50 nM MitoTracker Red® and subsequently stained with the specific nuclear vital dye Hoechst 33342. White arrows indicate the formation of nuclear fragment characteristic of cells undergoing apoptosis. Images were digitized with a Cool Snap video camera connected to Nikon Eclipse TE200 fluorescence microscopy. All of images were captured under constant exposure time, gain and offset. E, SH-SY5Y cells were treated with 50 μM Cu(isapn), Zn(isaepy), or 50 μM Cu(isaepy)₂ with or without the non-permeating copper chelator TRIEN (150 μM) for 24 h. Cells
were then washed and stained with propidium iodide. Analysis of subG1 (apoptotic) cells was performed by a FACScalibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. Data are expressed as mean ± S.D. n = 6; **p < 0.001, with respect to Cu(isaepy)₂ treated cells.

Fig. 9. Cu(isapn) and Cu(isaepy)₂ induce mitochondria-dependent apoptosis in U937 and M14 cells. A, Human melanoma M14 and promonocytoma U937 were treated with 50 μM Cu(isaepy)₂ for 24 h, washed and stained with propidium iodide. Analysis of cell cycle and apoptosis was performed by a FACScalibur instrument and percentages of staining-positive cells were calculated using WinMDI version 2.8 software. The cell cycle plots reported are from a typical experiment done in triplicate out of five that gave similar results. The percentages of apoptotic nuclei are shown above each histogram. B, alternatively, M14 and U937 cells were treated with 50 μM Cu(isaepy)₂ for 24 and 48 h. 40 μg of total protein extract was loaded onto each lane for detection of pro- and active-caspase-9, pro-caspase 3 and PARP. Western blots are from one experiment representative of three that gave similar results.
FIGURE 1

A

[Cu(isapn)]^{2+}

B

[Cu(isaepy)_2]^{2+}

C

Zn(isaepy)
FIGURE 2

A

% subG1 cells

24 h 48 h

Control Cu(isapn) Cu(isapn)₂

* *

B

p53 p21 α-tubulin

Cu(isapn)₂ Cu(isapn)

6 12 24 hours

C

Cytochrome c AIF Merge

Control

Cu(isapn)

Cu(isapn)₂

D

Cu(isapn) Cu(isapn)₂

0 24 48 hours

pro-Casp-9 active Casp-9 pro-Casp-3

PARP (117 kDa) PARP (83 kDa)

E

% subG1 cells

zVAD-fmk

Control Cu(isapn) Cu(isapn)₂
FIGURE 3

A

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FIGURE 5

A

B

C

D

FIGURE 5

A

B

C

D
FIGURE 7

A

Dead cells
(% of control)

Cu(isapn)  
Cu(isaepy)₂

- +  
- +

24 h  48 h

B

C

% of subG₁ cells

Cu(isapn)  
Cu(isaepy)₂

- +  
- +
Pro-apoptotic activity of novel Isatin-Schiff base copper(II) complexes depends on oxidative stress induction and organelle-selective damage

Giuseppe Filomeni, Giselle Cerchiaro, Ana Maria Da Costa Ferreira, Angelo De Martino, Jens Z. Pedersen, Giuseppe Rotilio and Maria R. Ciriolo

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