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N-Hydroxyphthalimide catalysts as bioactive prooxidants[†]

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The catalytic role of *N*-hydroxyphthalimide (NHPI) in promoting free-radical hydrogen atom transfer (HAT) reactions, well-documented for processes of industrial and synthetic interest, is here investigated for the first time in a biological environment. While NHPI by itself did not show any bioactivity, selected NHPI-derivatives (NHPIDs) revealed the ability to activate the intracellular formation of reactive oxygen species (ROS), causing the depletion of glutathione (GSH) levels and an increase in oxidative stress (OS). The evident bioactivity of some of these derivatives resulted in a significant reduction of the viability in osteosarcoma MG-63, suggesting a new, potential role of NHPIDs as pro-oxidant drugs. The key role of the N–OH group in promoting oxidative stress is demonstrated.

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1. Introduction

NHPI is a cheap molecule, which is emerging as a valuable organocatalyst capable of promoting free-radical aerobic oxidation by activating hydrogen atom transfer (HAT) processes.¹

The efficiency and selectivity of this catalyst have been related to the favourable combination of enthalpic, polar, and entropic effects.²

As depicted in Scheme 1, the catalytic cycle implies the *in situ* generation of phthalimide-*N*-oxyl (PINO) radical by several metal¹ and metal-free initiators.³

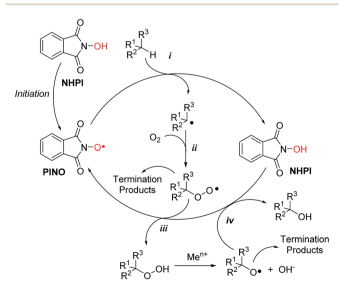
PINO, whose formation was proved by Electron Spin Resonance (ESR) spectroscopy,⁴ is the real responsible of the catalytic activity of NHPI in oxidative processes, being directly involved in the free-radical propagation step by hydrogen atom abstraction from different activated organic substrates (Scheme 1, path i). The resulting carbon centred radicals undergo molecular oxygen addition, leading to the formation of the corresponding peroxyl radicals (Scheme 1, path ii).

The catalytic cycle is completed by the HAT reaction from a new NHPI molecule to the peroxyl radical (Scheme 1, path iii), favouring the concomitant formation of hydroperoxides and a new PINO unit, so that the radical chain is maintained.

Moreover, the possible presence of transition metal ions would favour the reduction of the hydroperoxide, leading to the formation of alkoxyl radicals which in turn are even faster in abstracting hydrogen from C–H bonds and from NHPI itself (Scheme 1, path iv). Due to its unique catalytic role, in the last decade this derivative has attracted increasing attention, and its use has been also suggested for processes of industrial relevance.^{5,6}

We also showed how NHPI and related derivatives (NHPIDs) are effective in catalysing the peroxidation of polyunsaturated fatty acids (PUFAs).^{7,8} In that case the final purpose was the selective synthesis of PUFAs' corresponding hydroperoxides. However, by considering that PUFAs are important components of the cellular membrane and environment, the manifested efficiency of this process prompted us to further investigate the possible catalytic action of NHPIDs in promoting oxidative stress (OS) in a biological environment *in vitro*.

The reasons of this interest rely on the still controversial role played by OS in regulating cancer development and apoptosis.



Scheme 1 General free-radical cycle for the NHPI-catalysed oxidation of organic substrates.

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OS occurs when redox homeostasis within the cell is altered.⁹ While experimental evidences show that an increase in the cellular concentrations of reactive oxygen species (ROS), like alkoxyl and peroxyl radicals, can promote DNA damage, angiogenesis, and metastasis,¹⁰ more recently researchers recognized the possible role of ROS in cancer therapy, leading to the conclusion that, to consider OS as outright toxic would be a mistake.¹¹ In fact, it was reported that an increase of intracellular ROS content to cytotoxic levels would selectively induce apoptotic cell death in cancer.⁹ Thereby, besides the investigation of natural and synthetic agents with antioxidant activity,^{12,13} the development of pro-oxidant agents capable of inducing OS is nowadays emerging as an intriguing anticancer strategy.^{14–16}

Herein, for the first time we demonstrate that the prooxidant catalytic activity of NHPIDs, originally developed for industrial applications, can be exerted also *in vitro* to address relevant biological issues. Striking outcomes of the study include (i) the key role that lipophilic tails on the aromatic ring play in favouring the interaction of NHPI with cells; (ii) the association between the cytotoxic action of NHPIDs and their catalytic activity as pro-oxidants, evidenced by the increase of OS, followed by the depletion of intracellular glutathione (L- γ -glutamyl-L-cysteinyl-glycine, GSH) levels and consequently apoptotic cell death; (iii) the importance of having free N–OH moieties in order to exhibit a pro-oxidant activity, suggesting that the catalytic action *in vitro* could still follow the free-radical HAT mechanism described in Scheme 1.

2. Results and discussion

Human osteosarcoma MG-63 cells were chosen as target recipients because being responsive to OS insult.^{17,18}

Disappointingly, first experiments carried out in the presence of increasing amounts of NHPI for 24 h showed that it did induce significant cytotoxicity only at extremely high concentrations ($EC_{50} = 7 \text{ mM}$), as determined by AlamarBlue® fluorimetric assay (Fig. 1A). We thus hypothesized that this behavior would be associated to the strong polar character of NHPI, a key factor already investigated for industrial applications.^{19,20} In biological systems, high polarity could drastically reduce the reactivity of NHPI towards the plasma membrane, preventing its effective interaction with the cell.

In order to ascertain this, we synthesized NHPIDs bearing different lipophilic tails (Fig. 2), reproducing the same synthetic

solutions suggested to overcome the industrial limit of NHPI related to its solubility. 20

Compounds **1-OH** and **2-OH** were obtained starting from trimellitic anhydride chloride, after optimization of the experimental procedure reported by Ishii for the synthesis of **2-OH**.²¹

Similarly, compound **3-OH** was synthesized in the presence of the purposely prepared (*R*)-2-(1-hydroxybutan-2-yl)-1*H*-benzo [de]isoquinoline-1,3(2*H*)-dione, following the approach recently reported by our group.²² Interestingly, specifically functionalized naphthalimides have already shown good anticancer activity upon a variety of human cancer cells due to their high intercalation affinity towards DNA.²³ The cytotoxic properties of these derivatives were usually ascribed to the side chains present on the naphthalene moiety. However, data reported in the literature suggested that this target group would favor cellular internalization, rendering derivative **3-OH** attractive for our purpose. Moreover, we recently reported how catalyst **3-OH** could be photo-activated by UV radiation, further promoting aerobic oxidation.²²

In addition, we also synthesized compound **4-OH**, a bis-NHPI lipophilic catalyst, by reacting hydroxylamine hydrochloride with 4,4'-(4,4'-isopropylidenediphenoxy)bis (phthalic anhydride), a cheap and commercially available di-anhydride obtained from bisphenol A. The catalytic efficiency and the high lipophilic character of compound **4-OH** have been already documented by our group, for the solvent-free selective aerobic oxidation of alkyl aromatics.²⁰

NHPIDs were tested in MG-63 cancer cell line to check their potential, concentration-dependent, cytotoxicity. The results are reported in Fig. 1. Compounds **2-OH**, **3-OH**, and **4-OH** displayed EC_{50} at 100, 200 and 110 μ M, respectively (Fig. 1C–E, full black cycles), well below the values for the poorly cytotoxic compound **1-OH** (Fig. 1B).

These observations prompted us to further investigate the reasons for such a bioactivity. If our original assumption was correct, the decrease in cell viability following challenging with compounds **2-OH**, **3-OH**, and **4-OH** had to be due to a significant elevation of the OS. In fact, being NHPIDS **2-OH**, **3-OH**, and **4-OH** more lipophilic than, and as pro-oxidant as **1-OH**, they would be expected to interact with cells and exert potential pro-oxidant effects to a greater extent than the latter molecule. Moreover, onto consideration of the HAT mechanism according to which the NHPIDs promote oxidation (Scheme 1), we wanted to be sure that the eventual pro-oxidant action was specific to these compounds and relied on the presence of the N–OH

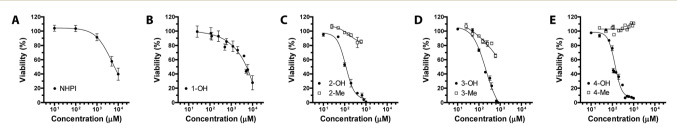
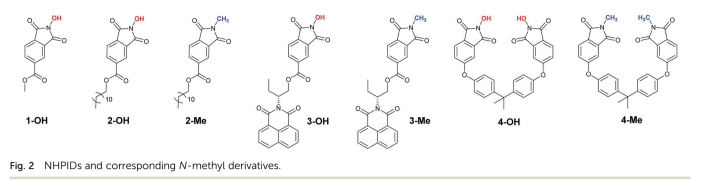


Fig. 1 Cytotoxicity of (A) NHPI, (B) 1-OH, (C) 2-OH vs. 2-Me (full black cycles vs. empty squares, respectively), (D) 3-OH vs. 3-Me (full black cycles vs. empty squares, respectively) and (E) 4-OH vs. 4-Me (full black cycles vs. empty squares, respectively) after 24 h of exposure as a function of concentration in MG-63 human osteosarcoma cells.



group, rather than to the introduction of the lipophilic tails. To do so, we challenged osteosarcoma cells with compounds **2-OH**, **3-OH**, and **4-OH** and the purposely synthesized *N*-methyl molecules **2-Me**, **3-Me**, and **4-Me** (Fig. 1C–E, empty squares *vs.* full black cycles, respectively). In this way we could evaluate and compare the eventual cytotoxicity exclusively associated to the grafting of the phthalimide aromatic ring (Fig. SI8†). In line with our expectations, **2-Me** to **4-Me** derivatives were far less cytotoxic than parent **N-OHs**.

Following these observations, we moved to consider the possible occurrence of OS both directly, by evaluating the increase of ROS, and indirectly, by measuring GSH depletion. In fact, chemotherapy agents (*e.g.* cisplatin) are detoxified by antioxidant defense mechanisms, such as enzymatic conjugation to the tripeptide GSH.²⁴ Abrogation of such drug-resistant mechanisms by redox modulation would thus have significant therapeutic implications.

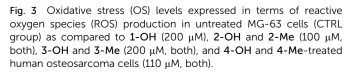
Interestingly, we herein show the generation of large quantities of ROS induced by the treatment with compounds **2-OH**, **3-OH**, and **4-OH** used each at its respective EC_{50} (130 ± 5%, 294 ± 20%, 234 ± 23%, all p < 0.05 vs. CTRL cells). Furthermore, our results about OS induced by derivatives **2-Me**, **3-Me**, and **4-Me** (all not statistically significant vs. CTRL cells) (Fig. 3) pointed out that the mechanism whereby NHPIDs **2-OH**, **3-OH**, and **4-OH** induce cytotoxicity is associated to the N–OH moiety. Indeed, injury to cells occurs only when ROS overwhelm the biochemical defense of the cell.⁹ Moreover, GSH consumption that mirrors the depletion of cellular antioxidant defenses,²⁵ is

related to such OS insult, as confirmed by the significantly greater GSH depletion with the parent molecule **4-OH**, if compared to **4-Me** compound (71 \pm 4% *vs.* 92 \pm 7%, respectively, *p* < 0.05) (Fig. SI9†).

We would like to point out that these results cannot be taken for granted. From one point of view, the aqueous environment could inhibit NHPIDs' catalytic properties by means of the hydrogen bonding involving N–OH moiety. On the other hand, we operated for the first time at really lower concentrations of the catalyst with respect to classical synthetic applications. It is in fact well known that peroxidation kinetics promoted by NHPI are significantly dependent on catalyst concentration.¹

The difference noticed among the EC_{50} of **2-OH** (100 μ M), **3-OH** (200 μ M) and **4-OH** (110 μ M) highlights one more the key role played by the lipophilic tails, suggesting that a proper design of the nitroxide pro-oxidant would allow to improve its biological performance. Such a difference is emphasized by the inherent behavior of the resulting **2-OH** to **4-OH** but not **1-OH**, to aggregate. In our experimental conditions, **2-OH**, **3-OH**, and **4-OH** compounds did form monodisperse aggregates with hydrodynamic diameters comprised between 136 and 628 nm (Fig. SI10–SI12†).

It is worth noting that, although also **2-Me**, **3-Me**, and **4-Me** do form aggregates in water in our experimental conditions (Fig. SI13–SI15†), sometimes (**2-Me**) with lower, sometimes (**3-Me**, **4-Me**) with higher hydrodynamic diameters than the corresponding *N*-hydroxy derivatives, they are in any case much less cytotoxic (Fig. 1).



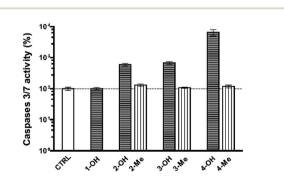


Fig. 4 Caspase-3 and -7 activities in untreated MG-63 cells (CTRL group) as compared to 1-OH (200 μ M), 2-OH and 2-Me (100 μ M, both), 3-OH and 3-Me (200 μ M, both), and 4-OH and 4-Me-treated human osteosarcoma cells (110 μ M, both).

This confirms once again the key function of any N–OH moiety. Overall, the inherent tendency of pro-oxidants **2-OH** to **4-OH** to aggregate do have a twofold advantage over less lipophilic molecules as **1-OH**. First, submicrometric aggregates can be easily endocytosed by cells. Moreover, apart from being pro-oxidant drugs *per se*, –**OHs** do form aggregates in water that might offer place to accommodate hydrophilic, lipophilic as well as amphiphilic drugs. This colloidal behavior would extend their use as chemotherapeutic carriers or drug delivery systems.²⁶

Excess OS kills or damages cells mostly by apoptosis, and eventually by other mechanisms.²⁷⁻²⁹ In the former case, alterations in the redox status of the cell to a more oxidizing environment occurs prior to activation of the final phase of cysteine aspartic acid-specific protease known as caspases that orchestrate the execution phase of apoptosis by cleaving multiple structural and repair proteins.⁹ Between the fourteen caspases identified in mammals, a subset of downstream, or executioner caspases (Caspase-3, and -7) is actually responsible for the demolition phase of apoptosis.³⁰ We thereby focused our attention on the primary effector caspases necessary for triggering the cleavage of the vast majority of subcellular proteins.

As shown in Fig. 4, Caspase-3 and -7 activity was significantly higher in cells treated with any of the NHPIDs as compared to each respective –**Me** derivative (597 \pm 47% *vs.* 126 \pm 7% for **2-OH** and **2-Me**, 679 \pm 25% *vs.* 104 \pm 6% for **3-OH** and **3-Me**, 6.664 \pm 223% *vs.* 126 \pm 7% for **4-OH** and **4-Me**, all *p* < 0.05), strengthening the idea that overt cytotoxicity specifically induced by **2-OH**, **3-OH**, and **4-OH** was linked to OS insult. This, in turn, led to some GSH depletion and engages cells in caspase-mediated apoptotic cell death.

3. Conclusions

In summary, lipophilic NHPIDs **2-OH**, **3-OH**, and **4-OH** were synthesized and their activity as pro-oxidant catalysts was substantiated *in vitro* in osteosarcoma MG-63 cells.

The overt cytotoxicity of these derivatives was associated to an increase of OS levels, and to a consequent depletion of intracellular GSH content, leading to caspase activation and apoptotic cell death. Interestingly, the markedly lower bioactivity of the *N*-methyl derivatives **2-Me**, **3-Me**, and **4-Me** shed light on the key role played by N–OH group in promoting oxidation even *in vitro*, suggesting the occurrence of the classical HAT process.

Taken together, these observations provide unambiguous evidence that NHPI, an oxidation catalyst widely used in a plethora of abiotic industrial applications, once tethered to suitable lipophilic moieties is a promising pro-oxidant *in vitro*.

It is worth noting that, in this preliminary investigation, our aim was to verify an hypothesis, that is the potential bioactivity of NHPIDs, rather than to design the best drug solutions for *in vitro* and *in vivo* applications.

Even though further investigation is necessary to make these catalysts more effective and selective towards target cells, we believe that these preliminary results may open up the possibility of the development of novel anticancer chemicals and strategies based on NHPI catalysis.

4. Experimental

4.1. General

All reagents used in this work are commercially available (Sigma-Aldrich, Italy) and were used as received without further purification unless otherwise stated. ¹H and ¹³C-NMR of the products were recorded at room temperature (r.t.) with a Bruker Avance-400 NMR spectrometer. Compounds **1**–4 were stored dried at 4 $^{\circ}$ C and solubilized in dimethyl sulfoxide (DMSO) before use.

4.2. Synthetic procedures

4.2.1. Synthesis of NHPIDs 1-OH, 2-OH, and 3-OH. Compound 3-OH was obtained according to the procedure previously described,22 by adding dropwise and under stirring at 0 °C for \sim 30 min, 20 mL of an anhydrous THF solution of (*R*)-2-(1-hydroxybutan-2-yl)-1H-benzo[de]isoquinoline-1,3 (2H)-dione (10 mmol) into 20 mL of an anhydrous solution of THF containing trimellitic anhydride chloride (10 mmol, 2.106 g) and 4 mL of anhydrous pyridine. The mixture was kept at 0 °C for 6 h and then overnight at r.t. After filtrating the white precipitate, the clear liquid was vacuum evaporated obtaining a white solid that was dissolved in pyridine (~40 mL) and reacted with NH₂OH · HCl (50 mmol) in microwave for 1 h under reflux. The evaporation of the solvent and the washing of the mixture with 0.1 M HCl at 0 °C resulted in a white solid that was purified by chromatography on silica-gel using $9:1 (v:v) CHCl_3: MeOH$ as eluent ($R_{\rm f}$: 0.60 – orange color spot if TLC is exposed to NH₃). Yield: 90%. Mp: 115.0 °C. ¹H-NMR (400 MHz, DMSO-d6) δ (ppm): 10.91 (s, 1H), 8.49 (d, 2H), 8.42 (d, 2H), 8.18 (d, 1H), 8.01 (s, 1H), 7.84 (m, 1H + 2H), 5.43 (s, 1H), 4.95 (dd, 1H), 4.76 (dd, 1H), 2.21 (m, 1H), 2.018 (m, 1H), 0.97 (t, 3H). ¹H-NMR and ¹³C-NMR spectra of this derivative is reported in Fig. SI4.[†]

Compounds **1-OH** and **2-OH** were synthetised similarly to **3-OH**, following the procedures previously described,²⁰ first preparing the ester from MeOH and dodecyl alcohol respectively and trimellitic anhydride chloride. The corresponding NHPIDs **1-OH** and **2-OH** were obtained using NH₂OH·HCl and purified by chromatography on silica-gel using 9 : 1 (v : v) CHCl₃ : MeOH as eluent (R_{f} : 0.60 (3), 0.76 (4) – orange color spot if TLC is exposed to NH₃). Yield: 94%. Mp: 172.0 °C (**1-OH**), 85.0 °C (**2-OH**). ¹H NMR of **1-OH** (400 MHz, acetone-d6) δ (ppm): 9.95 (broad s, 1H), 8.43 (d, 1H), 8.28 (s, 1H), 7.96 (d, 1H), 3.98 (s, 3H). ¹H-NMR of **2-OH** (400 MHz, DMSO-d6) δ (ppm): 8.44 (d, 1H), 8.31 (s, 1H), 7.96 (s, 1H), 4.40 (t, 2H), 1.83 (q, 2H), 1.6–1.2 (m, 18H), 0.88 (t, 3H). ¹H-NMR and ¹³C-NMR spectra of these derivatives are reported in Fig. SI1 and 2,† respectively.

4.2.2. Synthesis of NHPID 4-OH. Compound 4-OH was prepared according to the procedure previously described.²⁰ 4,4'-(4,4'-Isopropylidenediphenoxy)bis(phthalic anhydride) was purified by recrystallization from acetic anhydride. 5 g of dianhydride (commercially provided as yellowish flakes) were dispersed in 40 mL of acetic anhydride and refluxed at 139 °C

for 1 h in a microwave reactor with the automatic control of power (Micro-SYNTH Labstation – Milestone Inc., USA). After cooling at r.t. pure crystals were recovered by filtration on filter paper and extensively washed with cold diethyl ether in order to remove the acetic anhydride. The white crystals were dried under vacuum and stored in desiccator until next use.

In a 50 mL two-necked flask, 2.60 g (5 mmol) of purified 4,4'-(4,4'-isopropylidenediphenoxy)bis(phthalic anhydride) were added to 30 mL of anhydrous pyridine. After adding an excess of hydroxylamine hydrochloride (30 mmol, 2.85 g), the mixture was heated under reflux (119 °C) for 1 h in a microwave reactor. After cooling the pyridine was evaporated under vacuum, obtaining a viscous orange product which was dissolved in ethylacetate (150 mL). The organic phase was extensively washed with 0.5 M HCl (100 mL, 3 times) and deionized water (dH₂O, 100 mL, 3 times) in order to remove the residual pyridine and finally dried with anhydrous sodium sulphate. The evaporation of the solvent gave a pale yellow foam, which was purified by chromatography on silica-gel using 9:1 (v:v) $CHCl_3$: MeOH as eluent (R_f : 0.54 – orange color spot if TLC is exposed to NH₃). Yield: 93%. ¹H-NMR of 4-OH (400 MHz, DMSO-d6): $\delta = 10.77$ (s, 2H), 7.80 (d, 2H), 7.34 (d, 4H), 7.29 (d, 2H), 7.24 (s, 2H), 7.08 (d, 4H), 1.69 (s, 6H). ¹H-NMR and ¹³C-NMR spectra of this derivative is reported in Fig. SI6.[†]

4.2.3. Synthesis of methylated derivatives. Compounds 2-Me, 3Me and 4-Me were prepared similarly to 2-OH, 3-OH and 4-OH using MeNH₂·HCl in place of hydroxylamine hydrochloride. The NMR spectra of all these derivatives are reported in ESI.[†]

2-Me. Yield: 95%; ¹H-NMR (400 MHz, CDCl₃): δ (ppm): 8.44 (d, 1H), 8.37 (s, 1H), 7.88 (s, 1H), 4.44 (t, 2H), 3.19 (s, 1H), 1.77 (q, 2H), 1.45–1.10 (m, 18H), 0.85 (t, 3H).

3-Me. Yield: 94%; ¹H-NMR (400 MHz, CDCl₃) δ (ppm): 8.59 (d, 2H), 8.33 (s, 1H), 8.26 (d, 1H), 8.21 (d, 2H), 7.79 (d, 1H), 7.75 (t, 2H), 5.55 (m, 1H), 5.05 (dd, 1H), 4.85 (dd, 1H), 2.34 (m, 1H), 2.09 (m, 1H), 1.01 (t, 3H).

4-Me. Yield: 94%; ¹H-NMR (400 MHz, CDCl₃) δ (ppm): 7.77 (d, 2H), 7.36 (d, 4H), 7.29 (d, 2H), 7.24 (s, 2H), 7.00 (d, 4H), 3.15 (s, 6H), 1.73 (s, 6H).

¹H-NMR and ¹³C-NMR spectra of these derivatives are reported in Fig. SI3, SI5 and SI7,[†] respectively.

4.3. Biological procedures

4.3.1. Cell cultures. MG-63 cells (human osteosarcoma cell line, CRL-1427) were purchased from the American Type Culture Collection (ATCC, USA) and cultured at 37 °C, 99% humidity with regular supply of 5% CO₂, in complete medium consisting in Dulbecco's Modified Eagle Medium (D-MEM) added with 10% (v : v) of Fetal Bovine Serum (FBS), 1 mM sodium pyruvate, 10 mM Hepes buffer, 100 U mL⁻¹ penicillin, 0.1 mg mL⁻¹ streptomycin, and 2 mM glutamine.

4.3.2. Cell treatment. Cultures were continuously kept in exponential-phase growth before experiments. The day before the beginning of the treatments, cells were trypsinized, plated at the density of 1.5×10^4 cells per cm² and allowed to adhere overnight. 24 h post-seeding, the medium was replaced with

a fresh aliquot (300 μ L cm⁻²), supplemented with 1.5% (v : v) DMSO-containing compounds at the desired concentration and kept in contact with cells for 24 h before analysis. Cells grown in complete medium containing 1.5% (v : v) DMSO were considered as untreated controls (CTRL).

4.3.3. Cell viability test. Cytotoxicity was evaluated using AlamarBlue® cell viability assay (Thermofisher Scientific, Italy) according to manufacturer's guidelines. Briefly, at the end of treatment, the cell culture medium was replaced with fresh medium containing 10% of AlamarBlue® reagent and after 2 h of incubation at 37 °C, fluorescence ($\lambda_{ex} = 540$ nm; $\lambda_{em} = 595$ nm) was measured by a multimethod plate reader (GENios Plus, TECAN, Italy). Experiments were performed at least 3 times in triplicate and cell viabilities were expressed as percentage of CTRL cells (assigned as 100%). Cytotoxicity (%) was calculated as follows: 100 (%) – viability (%).

4.3.4. Evaluation of oxidative stress (OS) levels. OS was monitored by evaluating the fluorescence of 2',7'-dichlorofluorescein (DCF) produced by intracellular deesterification and oxidation of 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA). Briefly, at the end of treatment, cells were washed with PBS, incubated for 15 min at 37 °C with 10 µM DCFH-DA in Phosphate Buffered Saline (PBS), washed twice with the same buffer and lysed with 0.5% (v : v) Tween in 50 mM Tris-HCl buffer, pH 7.5 for at least 10 min at r.t. Cells were then scraped, collected and subjected to 3 consecutive freezing/thawing cycles (-80 °C/r.t.) and the fluorescence in cell lysates measured by means of a microplate reader ($\lambda_{ex} = 485 \text{ nm}$; $\lambda_{em} = 530 \text{ nm}$), then normalized over the total protein content of each sample as determined by bicinchoninic acid (BCA) protein assay kit (Pierce, USA). Experiments were performed at least 3 times in quadruplicate and values expressed as percentage of CTRL cells.

4.3.5. Quantification of intracellular GSH content. For evaluating the total intracellular glutathione (GSH) content after treatment, cells were trypsinized, harvested, centrifuged at 5×10^3 g for 5 min and washed once with PBS. Pellets were lysed in 5% (w : v) sulfosalicylic acid (SSA) in dH₂O and subjected to 2 freezing/thawing cycles (-80 °C/r.t.). After an incubation of 10 min at 4 °C, lysates were finally deproteinized by centrifugation for 10 min at 6×10^3 g and GSH levels determined in supernatants with Glutathione assay kit according to the manufacturer's instructions. Experiments were performed at least twice in triplicate and values, normalized for protein content, were expressed as percentage of CTRL cells.

4.3.6. Biophysical characterization of aggregates. The existence and the size (hydrodynamic diameter) of aggregates were determined by Dynamic Light Scattering (DLS) utilizing Zetasizer Nano ZS (ZEN3500, Malvern Instruments Ltd., UK). Briefly, all compounds were solubilized in DMSO, diluted 66 times in 10 mM Hepes buffer to the final concentration of 200, 100, 200 μ M, and 110 μ M for compounds **1**, **2**, **3**, and **4**, respectively, and incubated 30 min at 37 °C before analysis. All measurements were performed at 37 °C in a final volume of 500 μ L. Single measurements were repeated at least twice in independent experiments. Representative intensity distributions are shown in Fig. SI10–SI15.†

4.3.7. Evaluation of cell death mechanism. Caspase-3 and -7 activity was measured using the luminogenic Caspase-Glo® 3/7 assays (Promega Corp., Italy) according to manufacturer's guidelines. Briefly, at the end of treatment, the medium was replaced with a fresh aliquot and the cells were allowed to equilibrate at r.t. for 15 min before adding an equal volume of reagent. After gentle mixing for 1 h at 100 rpm on an orbital shaker, the luminescence of each sample was measured by means of a microplate reader (GENios Plus, TECAN, Italy). Values were subtracted of no-cell background, standardized over cell viability, and expressed as percentage of CTRL cells. Each experiment was performed at least in quadruplicate.

4.3.8. Statistical analysis. Results were expressed as mean \pm SD and statistical analyses carried out by GraphPad version 5.04 (GraphPad software, USA). Comparisons among groups were performed by one-way ANOVA with post-hoc Bonferroni test. Significance was retained when p < 0.05.

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