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Osmocenyl-tamoxifen derivatives target the thioredoxin system leading to a redox imbalance in Jurkat cells

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Keywords

Osmocene, osmocenyl-tamoxifen, thioredoxin reductase, ROS, Jurkat cancer cells

ABSTRACT:

To complete our study of the iron, ruthenium and osmium metallocene triad derived from hydroxytamoxifen, we report here the synthesis and study of the biological effects of two ferrocifen analogs in the osmium series, namely the monophenolic complex 1, the tamoxifen-like complex 2 and their oxidized quinone methide derivatives, 1-QM and 2-QM. Studies carried out in vitro on purified thioredoxin reductase (TrxR) show that the characteristic inhibition of TrxR is observed with 1 and 2 only after their enzymatic oxidation by the H₂O₂/HRP system. The observed inhibition is very similar for the two complexes (IC₅₀ = 2.4 and 1.2 μ M for 1 and 2 after treatment by HRP/H₂O₂) but different from that obtained with the corresponding quinone methides (IC₅₀ = 5.4 and 3.6 μ M for **1-QM** and **2-**QM). The UV spectra of 1 or 2 incubated in the presence of HRP/H₂O₂ show that the species generated by enzymatic oxidation is not a quinone methide but more probably its corresponding cation. On Jurkat cells, complexes 1 and 2 behave differently. Indeed, while 2 shows high toxicity (IC₅₀ = 7.4 μ M), that of 1 is much more modest (IC₅₀ = 42 μ M). Interestingly, a significant inhibition of TrxR activity is observed in cells incubated with 2 (about 70% inhibition for incubation in the presence of 15 μ M) while that induced by 1 is much weaker (about 30% inhibition for incubation in the presence of 50 μ M). This strong inhibition of TrxR activity induced by 2 leads to accumulation of thioredoxin and peroxiredoxin 3 in oxidized form as well as a significant decrease of the mitochondrial membrane potential (MMP). These results show that the cytotoxicity of the osmocifens depends on their oxidation within the cell and that inhibition of thioredoxin reductase is a key factor in rationalizing the cytotoxicity of these complexes on Jurkat cells.

1. INTRODUCTION

Owing to the poor outcomes seen in the treatment of a number of cancers such as glioblastoma, melanomas, ovarian and pancreatic cancers etc., the search for new active principles outside the established avenues is a burning current concern. For this reason, the unusual mechanistic properties of organometallic medicinal chemistry are presently the subject of an exponentially growing number of studies [1]. These entities offer inventive strategic approaches [2-6] in terms of optimized space filling, intracellular redox behavior and antitumoral activity, with pharmacophores possessing varied and multiple targeting options that make it possible to counteract the phenomena of resistance to proapoptotic stimuli [7]. To that can be added the variety of available metals [2] and, for the transition elements, the potential availability of several degrees of oxidation.

With organometallic complexes of gold [8, 9], those from the group 8 triad, particularly of Fe and Ru, are amongst the most widely studied [10-13] with the notable exception to date of Os [14-17]. This heavy element offers a capacity for good stabilization of certain high degrees of oxidation (+VI) and for an excellent metal-to-ligand back-donation at low oxidation states (+II, +III) leading to more stable complexes than those of Fe or Ru in biological media. The potential of this so far neglected metal in biomedically-oriented organometallic complexes deserves to be better studied especially when comparisons with neighboring series of Fe and Ru are easy to obtain [18, 19].

Amongst the promising derivatives of ferrocene, the ferrocifen series, derived initially from tamoxifen, the drug used to combat hormone-dependent breast cancers, has been the subject of a number of studies [20-22]. These organometallic species can operate on cancerous cells via the signaling pathways of senescence [23, 24] or apoptosis, with markedly different effects depending on the concentration at which they are used and the type of cell involved, while healthy cells are affected only slightly or not at all [25].

Ferrocifen equivalents in the Ru and Os series (mono- and diphenolic, and tamoxifen-like) have recently been proved possible to synthesize [19, 26]. Their electrochemical behavior as well as their cytotoxicity on breast cancer cells (MDA-MB-231 and MCF-7) have also been studied. In the two series, the tamoxifen-like complexes have high cytotoxicity while the phenolic complexes have much more modest cytotoxicity. This result is very different from that of the ferrocene analogs which have similar high cytotoxicities. In addition, unlike ferrocifens, where the active metabolite has been identified as being a quinone methide [27, 28], with these heavy metals the electrochemical signature of a quinone methide has not been found. Studies previously carried out on the chemical and biochemical properties of these complexes (lipophilicity, solubility and rate of formation of quinone methides, differences in the acidity of the phenolic protons) have not been able to elucidate their mechanism of action [26].

In cells, the thiol redox balance and removal of hydrogen peroxide and other hydroperoxides is exerted by the thioredoxin and glutathione systems. To remove hydrogen peroxide these systems utilize glutathione peroxidase and peroxiredoxin, respectively [29]. Of note, both glutathione peroxidase and thioredoxin reductase contain a selenocysteine residue at their catalytic site. In particular, thioredoxin reductase which is present in two isoforms, a cytosolic (TrxR1) and a mitochondrial one (TrxR2), is endowed with a C-terminal motif Cys-SeCys-Gly. In this sequence, selenocysteine is considered the

major reactive site of the molecule towards metals, metal complexes and electrophilic organic compounds all able to inhibit the enzyme activity at micro/submicromolar concentrations [30]. In fact, the selenocysteine residue possesses a low pKa (5.24) that enhances its nucleophilic properties towards the above mentioned classes of compounds [31].

The recent study of thioredoxin reductases as possible targets of the ferrocifens has allowed new perspectives concerning their mechanism of action [32]. In the present paper, we examine the biological effects of two osmocifens, namely the monophenolic and the tamoxifen-like complexes **1** and **2**, their oxidized derivatives obtained after enzymatic oxidation, and their corresponding quinone methides obtained by chemical oxidation (Chart 1). In particular, their effects on thioredoxin reductases both isolated and obtained from treated cancer cells, the redox state of thiol enzymes and the effects on ROS production and mitochondrial bioenergetics were considered. Inhibition of thioredoxin reductase seems to play a critical role in inducing the observed cytotoxic effects exerted by **1** and **2** in cultured cancer cells.



Chart 1: Osmocifens (monophenol and tamoxifen-like osmium complexes)

2. MATERIALS AND METHODS

2.1. Synthesis of 1, 2 and of their corresponding QMs

General remarks: All reactions and manipulations were carried out under an argon atmosphere using standard Schlenk techniques. Prior to use, acetone was dried over CaCl₂, and pentane was dried over 4 Å molecular sieves. Ag₂O was freshly prepared according to the literature procedure [33]. ¹H and ¹³C-NMR spectra were acquired on a Bruker 300 MHz spectrometer. Mass spectrometry was carried out at the "Service de Spectrométrie de Masse" at ENSCP, Paris. Elemental analysis and HRMS determinations were performed by the Laboratory of Microanalysis and the Small Molecule

Mass Spectrometry Platform of IMAGIF (Centre de Recherche de Gif www.imagif.cnrs.fr). ICP-OES analysis were performed on the on a JY2000 apparatus (Jobin-Yvon Horiba) of the Metis Laboratory of the University Pierre et Marie Curie. Complexes **1** and **2** were prepared as previously described [26].

2.1.1. QM of 1-phenyl-1-(hydroxyphenyl)-2-osmocenyl-but-1-ene (1-QM)

1 (60 mg, 0.11 mmol) was dissolved in dry acetone (5 mL). Ag₂O (154 mg, 0.66 mmol) was added and the mixture was stirred for 2 h at room temperature. The red solution of quinone methide formed was separated from the solid by centrifugation (room temperature, 400 g, 5 min) and evaporated. A red oil (53 mg, 89% yield) of quinone methide was obtained. 1 mL of acetone was added to dissolve the quinone methide. Pentane was added to precipitate the quinone methide (red solid), Z + E isomers; E isomer is supposed to be the major isomer (E:Z 73:27). ¹H NMR (300 MHz, CD₃COCD₃) δ : 1.59 (d, 3H, J = 7.0 Hz, CH₃, major isomer), 1.99 (d, J = 7.0 Hz, 3H, CH₃, minor isomer), 4.50 (s, 5H, C₅H₅, major isomer), 4.50 – 4.64 (m, 4H,C₅H₄), 4.64 (s, 5H, C₅H₅, minor isomer), 5.75 (q, J = 7.0 Hz, 1H, C<u>H</u>-Me, minor isomer), 6.28 (q, J = 7.0 Hz, 1H, C<u>H</u>-Me, major isomer), 6.30-6.39 (m, 2H, C₆H₄), 7.33 (dd, J = 10.0 Hz + J = 2.5 Hz, 1H, C₆H₄, minor isomer), 7.40-7.53 (m, 7H, C₆H₅+C₆H₄). ¹³C NMR (75.47 MHz, CD₃COCD₃) δ : 15.8 (CH₃), 63.6 + 64.0 + 64.5 66.5 (C₅H₄), 65.8 (C₅H₅, major isomer), 66.0 (C₅H₅, minor isomer), 85.7 (C, C₅H₅), 125.2 (CH₃-<u>C</u>H=C), 129.2, 129.4, 130.7, 131.6, 132.1, 139.9 (<u>C</u>H of C₆H₅ and C₆H₄), 138.4 (C), 139.1 (C), 140.0 (C), 157.3 (C), 186.8 (C=O). EI-MS: 542.3 [M]·⁺ HRMS (ESI, [C₂₆H₂₂OOs+H]⁺) calcd: 543.1359, found: 543.1338. Anal. Calc. for C₂₆H₂₂OOs.0.5acetone (%): C, 57.97; H, 4.42. Found: C, 58.06; H, 4.

2.1.2. *QM* of 1-[(4-(3-dimethylaminopropoxy)phenyl)]-1-(hydroxyphenyl)-2-osmocenyl-but-1-ene (**2-***QM*)

2 (71 mg, 0.11 mmol) was dissolved in dry acetone (10 mL). Ag₂O (154 mg, 0.66 mmol) was added and the mixture was stirred for 2 h at room temperature. The red solution of quinone methide formed was separated from the solid by centrifugation (room temperature, 400 g, 5 min) and evaporated. A red oil (63 mg, 90% yield) of quinone methide was obtained. 1 mL of acetone was added to dissolve the quinone methide. Pentane was added to precipitate the quinone methide (red solid). ¹H NMR (300 MHz, CD₃COCD₃): δ 1.54 (d, *J* = 7.0 Hz, 3H, CH₃), 1.93 (m, 2H, CH₂), 2.17 (s, 6H, NMe₂), 2.41 (t, *J* = 6.9 Hz, 2H, CH₂), 4.14 (t, *J* = 6.4 Hz, 2H, OCH₂), 4.27 (broad s, 1H, C₅H₄), 4.53 (s, 5H, C₅H₅), 4.53-4.66 (m, 2H, C₅H₄), 4.97 (broad s, 1H, C₅H₄), 6.27 (q, *J* = 7.0 Hz, 1H, C<u>H</u>-Me), 6.27-6.36 (m, 2H, C₆H₄), 7.08 (d, J = 8.8 Hz, 2H, C₆H₄), 7.38 (d, J = 8.8 Hz, 2H, C₆H₄), 7.41 (dd, J = 10.0 and 2.8 Hz, 1H, C₆H₄), 7.51 (q, J = 10.0 and 2.8 Hz, 1H, C₆H₄). ¹³C NMR (75.46 MHz, CD₃COCD₃) δ : 15.7 (*CH*₃), 28.2 (OCH₂*CH*₂CH₂C), 30.1 (*CH*), 45.7 (N(CH₃)₂), 56.7 (OCH₂CH₂*CH*₂*CH*₂N), 63.8 (2CH, C₅H₄), 65.8 (7CH, C₅H₅ + C₅H₄), 67.1 (O*CH*₂CH₂CH₂N), 115.4 (2CH, C₆H₄), 125.4 (CH₃-<u>C</u>H=C), 128.7 (CH), 129.4 (CH), 133.6 (2CH, C₆H₄), + 138.3 (CH), 161.9 (C-OCH₂). MS (CI): m/z = 644.89 [M+H]⁺. MS (CI): m/z = 644.89 [M+H]⁺. HRMS (ESI, [C₃₁H₃₃O₂Os+H]⁺) calcd: 644.2200, found: 644.2187. Anal. Calc. for C₃₁H₃₃NO₂Os.0.5acetone (%): C, 58.19; H, 4.41; N, 2.09. Found: C, 58.31; H, 5.11; N, 2.05.

For biochemical experiments, 10 mM stock solutions of 1, 2, 1-QM and 2-QM in DMSO were prepared just prior to use.

2.2. TrxR1 preparation and enzymatic activity assay

Highly purified cytosolic thioredoxin reductase (TrxR1) was prepared from rat liver according to the method of Luthman and Holmgren [34] and the protein content was measured according to Lowry *et al.* [35]. Thioredoxin reductase activity was determined by estimating the DTNB-reducing property of the enzyme in the presence of NADPH. Aliquots of highly purified TrxR1 (60 nM) in 0.2 M Tris HCl buffer (pH 8.1), 1 mM EDTA, and 0.25 mM NADPH were preincubated for 5 min with the various compounds, at 25 °C. The reaction was started with 1 mM DTNB and followed spectrophotometrically at 412 nm for about 10 min. TrxR2 activity was assayed with the same procedure.

2.3. Enzymatic oxidation of 1 and 2 by the H₂O₂/HRP system

Complexes 1 or 2 were incubated at a concentration of 25 μ M at 25 °C in 1 mL of 0.2 M Tris HCl buffer (pH 8.1), 1 mM EDTA, 2.5% DMSO with a solution of 22 nM HRP and 0.1 mM H₂O₂. Kinetics of the reaction was monitored by UV-VIS spectrometry on a Cary 50 spectrometer (Varian).

2.4. Cell cultures

Jurkat cells were grown at 37 °C in a 5% carbon dioxide atmosphere using RPMI 1640 medium containing 10% fetal calf serum and supplemented with 2 mM L-glutamine.

2.5. MTT assay

Cell viability was determined with the MTT reduction assay. Jurkat cells (8×10^4) were treated with increasing concentrations of osmocifens for 24 h. At the end of incubation, cells were treated for 3 h at 37 °C with 0.5 mg/mL MTT dissolved in PBS buffer. Afterward, 20 µL of stop solution (50% dimethylformamide, 20% SDS, 2% acetic acid and 25 mM HCl pH 4.7) were added to each well. After 15 min, the absorbance at 595 nm was estimated using a plate reader (Tecan Infinite® M200 PRO).

2.6 ICP-OES analysis

Jurkat cells were incubated for 24 h in the presence of 50 μ M of **1** or 15 μ M of 2 and washed twice in PBS buffer. Pellets of cells (10⁷ cells) were suspended in 0.5 mL of HCl (37%, Fluka for trace analysis) and digested at 60°C in an ultrasound bath for 1 h. Then the samples were adjusted to HCl 2% by addition of 8.75 mL of water and filtered on a 0.2 μ m filter. Quantification of osmium was performed at 225.585 nm. The results are the mean of 3 measurements. HCl was used instead of HNO₃ classically used for ICP-OES experiments as HNO₃ leads to the formation of the very toxic and volatile OsO₄.

2.7. Thioredoxin reductase and glutathione reductase assays in Jurkat cell lysates

Jurkat cells (2×10^6) were incubated with **1** and **2** for 18 h. After incubation cells were harvested and washed with PBS buffer. Each sample was lysed with a modified RIPA buffer containing 150 mM NaCl, 50 mM Tris HCl (pH 7.4), 1 mM EDTA, 1% TRITON, 0.1% SDS, 0.5% DOC, 1 mM NaF, 0.1 mM PMSF and an antiprotease cocktail ("Complete" Roche, Mannheim, Germany). After 40 min of incubation at 0 °C, lysates were centrifuged at 14000 g for 5 min. The supernatants were tested for enzyme activities. Aliquots of lysates were subjected to thioredoxin reductase determination with the insulin reduction test [36]. Briefly, 12 µg proteins of cell lysates were incubated for 40 min in a final volume of 50 µL in 100 mM HEPES buffer (pH 7.6) and in the presence of 15 mM EDTA, 1.5 mM NADPH, 0.20 mM insulin, and 100 µM Trx from *E. coli*, at 37 °C. The reaction was stopped by addition of 0.2 mL of 1 mM DTNB in 0.2 M Tris HCl buffer (pH 8.1) with 1 mM EDTA and 7.2 M guanidine. Samples were estimated at 412 nm using a plate reader.

Glutathione reductase activity (80 µg of cell lysates) was measured in 0.2 M Tris HCl buffer (pH 8.1), 1 mM EDTA, and 0.25 mM NADPH. The assay was initiated by addition of 1 mM GSSG and followed spectrophotometrically at 340 nm.

2.8. Redox Western blot analysis of Trx1, Trx2 and Prx3

The redox state of Trx was detected using a modified Western blot analysis [37]. Briefly, Jurkat cells (2×10^6) were treated with **1** (50 µM) and **2** (15µM) for 18 h, washed with PBS and then lysed with 100 µL of urea buffer (8 M urea dissolved in 100 mM Tris HCl, pH 8.3) containing 1 mM EDTA and 10 mM IAM in order to alkylate free thiols. Incubation was carried out at 37 °C for 20 min. After centrifugation, cell lysates were precipitated by ice-cold acetone–1 N HCl (98:2). The obtained pellets were washed with ice-cold acetone –1 N HCl-H₂O (98:2:10), resuspended in 60 µL of urea lysis buffer with 3.5 mM DTT and incubated for 30 min at 37 °C. Afterward, 3 µL of 600 mM IAA (30 mM final concentration) were added to the samples, followed by incubation for 30 min at 37 °C. Protein concentration was determined by the Lowry assay [35]. Proteins were separated by urea-PAGE gel (7% acrylamide/bis(acrylamide) in 7 M urea) and blotted using Turbo System (Bio-Rad Laboratories, Hercules, CA, USA). Membranes were probed with the primary antibodies respectively for Trx1 (FL 105) and for Trx2 (H75) (Santa Cruz Biotechnology, Santa Cruz, CA, USA).

For the determination of Prx3 redox state, after incubation with **1** (50 μ M) and **2** (15 μ M) for 18 h, Jurkat cells were collected, centrifuged at 500 g for 5 min, washed with cold PBS and then treated with 1 mL of 10% trichloro-acetic acid. The samples were kept at 4 °C for 30 min, and then centrifuged for 15 min at 15800 g at 4 °C. Pellets washed with 1 mL of ice-cold acetone, were then centrifuged at 10000 g for 10 min at room temperature. The samples were dissolved in 670 mM Tris HCl (pH 7.5) 2% SDS medium containing 10 mM AIS (4-acetamido-40-((iodoacetyl)amino)stilbene-2, 20-disulfonic acid) (Invitrogen). Derivatization lasted for 20 min, at room temperature, followed by further 45 min at 37 °C. Samples were loaded, without reducing agents, onto Bis–Tris Gel NUPAGE (12%) and blotted. To assess the redox state of Prx3, a monoclonal antibody LF-MA0044 (Histoline) was used.

2.9. ROS formation

The generation of reactive oxygen species (ROS) was assessed by the fluorogenic probe CM-DCFH2-DA (Molecular Probes, Invitrogen). Jurkat cells (4×10^4 per well) were seeded in PBS/10 mM glucose. After 1 h cells were loaded with 1 μ M CM-DCFH2-DA and with increasing concentrations of the compounds. Fluorescence increase, 485 nm (Ex) and 527 nm (Em), was followed for 2 h using a plate reader (Tecan Infinite® M200 PRO).

2.10. Flow cytometry analysis

Drug-influenced cell mitochondrial membrane potential was analyzed by flow cytometry. Jurkat cells (1×10^6 cells) were treated for 18 h in presence of **1** (15 or 50 µM) and **2** (10 or 15 µM).

Cells were then collected, resuspended in PBS/10 mM glucose (2.5 x 10⁵/mL) and loaded with 25 nM TMRM at 37 °C in the dark, for 15 min. Induced changes of membrane potential were estimated with a FACSCantoTM II flow cytometer (Becton-Dickinson, CA, USA) using an argon laser at 585 nm.

2.11. Quantitative enzymatic oxidation of 1 by the HRP/H₂O₂ system

Enzymatic oxidation was performed on **1** (100 μ M, 976 μ g) in H₂O/DMSO 9:1 (18 mL). Solutions of HRP (110 nM final) and H₂O₂ (0.5 mM final) were pre-incubated for 5 min then added to the aqueous solution of **1**. The colorless solution progresseively turned to yellow. The kinetics of the reaction was followed by the appearance of a peak at 329 nm in the UV-VIS and the reaction was complete in 20 min. Then, 2 equivalents of NaBPh₄ were added, leading to the formation of a yellow precipitate. This precipitate was recovered by centrifugation at 11500 g for 10 min, and washed twice with a small volume of cold water (100 μ L). Finally, the precipitate was dissolved in methanol (100 μ L). This solution was analyzed by UV-VIS and mass spectrometry.

3. RESULTS

3.1. Synthesis of the osmocifens 1, 2 and of their corresponding quinone methides (QMs)

The two osmium-based tamoxifen derivatives, **1** and **2** were prepared following the protocol recently published by some of us [26]. Their corresponding quinone methides **1-QM** and **2-QM** were synthesized by chemical oxidation with freshly prepared Ag₂O at room temperature and obtained with high yield (89 and 90% respectively; see experimental section).



Scheme 1 : Synthesis of quinone methides (QMs)

The stability of **1** and **2** was studied by UV-VIS spectrometry. They are both stable for at least 3 days in aqueous medium (25 μ M in Tris HCl (pH 8.1), 1 mM EDTA, 2.5% DMSO) (decrease of the characteristic absorbance of **1** and **2** at 300 nm less than 20%). (Fig. S1)

3.2. Enzymatic oxidation of 1 and 2 by the HRP/H₂O₂ system

In the presence of the enzymatic oxidizing system HRP/H₂O₂, complexes **1** and **2**, undergo oxidation with formation of new derivatives called **1*** and **2***. The rate of formation of **1*** and **2***, can be followed by UV-VIS spectrometry (Fig. 1 and Fig. S2). Following addition of the HRP/H₂O₂ mixture, we observed the rapid decrease of the shoulder characteristic of **1** and **2**, around 300 nm together with the appearance of a peak at 323 nm that characterizes **1*** or **2***. The formation of these species is rapid and slightly faster for **2*** than for **1***. Mathematical treatment of the UV-VIS data according the first order law gave t_{1/2} = 6.7 and 4.0 min, respectively (Fig. S3, Table S1). As shown on Fig. 1, UV-VIS spectra of **1*** and **2*** are clearly different from those of their corresponding quinone methides **1-QM** and **2-QM** which show peaks at 365 nm and 417 nm, respectively. Interestingly, while slow evolution of the UV-visible spectrum of **1*** and **2*** was observed with time they did not convert in the quinone methides even after 10 h (Fig. S4). It should also be noticed that the rate of formation of **2*** is significantly higher than that of **2-QM** obtained by chemical oxidation (Table S1) (t_{1/2} = 4 min for **2*** versus 24 min for **2-QM**).



Fig. 1. Superimposition of the UV-VIS spectra of 25 μ M of 1 (A) or 2 (B), their quinone methides (1-QM and 2-QM) and 1*, 2* the compounds obtained by enzymatic oxidation of 1 and 2 of with the HRP/H₂O₂ system after 20 min.. Spectra were recorded with 25 μ M compounds in 0.2 M Tris HCl buffer (pH 8.1), 1 mM EDTA and 2.5% DMSO. 1 and 2 are characterized by a shoulder around 300 nm, 1* and 2* by a peak at 323 nm, 1-QM and 2-QM by peaks at 365 and 417 nm, respectively.

3.2. Inhibitory effect on isolated thioredoxin reductase of osmocifens 1 and 2, their oxidized derivatives 1* and 2* and the quinone methides 1-QM and 2-QM

The inhibitory effect of **1** and **2** on thioredoxin reductase 1 (TrxR1) activity before and after enzymatic oxidation with HRP/H₂O₂ was first evaluated. As shown on Fig. 1A, **1** and **2** are scarcely effective on TrxR1 up to a concentration of 20 μ M, while they both acquire remarkable inhibitory

properties after oxidation with the HRP/H₂O₂ system. Oxidized complex 2 (2*) is a little more active than oxidized complex 1 (1*) (IC₅₀ = 1.2 μ M and 2.4, respectively). Quinone methides, 1-QM and 2-QM, are also effective inhibitors of the enzyme although to a lower extent (Fig. 1B; IC₅₀ = 5.4 and 3.6 μ M respectively). In addition, compounds 1, 2 and 1* and 2* were tested on mitochondrial thioredoxin reductase activity *in vitro* (Fig. S7) and only the transformed compounds treated with HRP and H₂O₂ show a significant inhibition of TrxR2.





1, **2** and the compounds obtained after pre-treatment for 15 min of **1** and **2** with 0.1 mM H_2O_2 and 22 nM HRP (**1***, **2***), were incubated in 0.2 M Tris HCl buffer (pH 8.1), 1 mM EDTA with 0.25 mM NADPH and 60 nM TrxR1. After 5 min, TrxR1 activity was estimated as described in Materials and methods. Under the same experimental conditions the HRP/ H_2O_2 mixture is ineffective on TrxR1 activity in the absence of osmocifen compounds (A). Thioredoxin reductase activity was estimated in the same experimental conditions after a 5 min treatment with **1-QM** or **2-QM** (B).

3.4. Antiproliferative effect and ROS production of 1 and 2 in Jurkat cells

The antiproliferative effect of **1** and **2** was studied on Jurkat cells incubated for 24 h. As shown in Fig. 3 A, complex **2** exhibits a strong antiproliferative effect ($IC_{50} = 7.4 \mu M$) while **1** is significantly less cytotoxic ($IC_{50} = 42 \mu M$). ROS production was then estimated in Jurkat cells after 2 h incubation (Fig. 3 B) and, similarly to the antiproliferative effect compound, **2** is more effective than **1**.



Fig. 3. Estimation of Jurkat cell viability (A) and ROS production (B) after treatment with compound 1 or 2. Cell viability was measured with the MTT test in $8x10^4$ cells treated for 24 h with the osmocifens as indicated in Materials and methods. For ROS production $4x10^4$ cells were incubated in PBS/10 mM glucose medium in the presence of 1 μ M CM-DCFH2-DA and of the osmocifens at the indicated concentrations. Reaction was followed as fluorescence increase and the reported values were obtained after 120 min of incubation.

3.5. Cellular uptake of 1 and 2

Quantification of Os in whole cells incubated for 24 h in the presence of 1 (50 μ M) or 2 (15 μ M) was performed by ICP-OES (Table S2). The amount of Os found in one cell was 3.9 fmol for 1 and 4.2 fmol for 2. Interestingly the amount of osmium is higher in cells incubated with 15 μ M of 2, the most cytotoxic complex, than in cells incubated with 50 μ M of 1 the less cytotoxic molecule.

3.6. Effect of 1 and 2 on the activity of thioredoxin reductase (TrxR) and glutathione reductase (GR) in Jurkat cells

Similarly to what observed *in vitro* with the H_2O_2/HRP system, compounds 1 and 2 can be oxidatively metabolized in cultured cells into species able to interact with the various enzymatic targets. On this basis, we have evaluated the total thioredoxin reductase and glutathione reductase activities in Jurkat cell lysates incubated with compound 1 or 2. Both GR and TrxR are pyridine disulfide oxidoreductases, however GR lacks the easily accessible C-terminal -Cys-SeCys- motif which characterises TrxR and is the target of a great deal of inhibitors.



Fig. 4. TrxR (*A*) and *GR* (*B*) activities in Jurkat cell lysates. Jurkat cells $(2x10^6)$ were incubated for 18 h with **1** (50 µM) or **2** (15 µM) and then lysed. Activities of the two redox enzymes were estimated as reported in Materials and methods.

As shown in Fig. 4 A both 1 and 2 inhibit total TrxR activity. However, 2 is far more effective than 1 as, at 15 μ M, it inhibits the enzyme activity by almost 70%, while 1, even at a higher concentration (50 μ M), is effective by only 30%. The activity of GR is not inhibited by 1 or 2, but

instead is slightly stimulated (Fig. 4 B). Interestingly, **1** and **2** at the reported concentrations (50 and 15 μ M, respectively) do not inhibit TrxR activity in the non-tumor cell line HEK293 as shown in Fig. S8.

3.7. Effect of compounds 1 and 2 on the oxidized state of thioredoxin (Trx1 and Trx2) and peroxiredoxin 3 (Prx3) in Jurkat cells

Thioredoxin reductase belongs to the thioredoxin system, which is a relevant metabolic pathway for the maintenance of cellular thiols in a reduced state. The major substrate of thioredoxin reductase, thioredoxin, in turn delivers electrons to peroxiredoxin which removes hydrogen peroxide and participates in signaling processes. We therefore examined the thiol redox state of both cytosolic and mitochondrial thioredoxins (Trx1 and Trx2) and the redox conditions of mitochondrial peroxiredoxin (Prx3). For estimation of Trx1/2 a modified redox Western blot, specific for thiol oxidation states, was used (see Materials and methods). In this blot the upper band indicates the fully reduced enzyme while the lowest indicates the completely oxidized form. The intermediate bands indicate the partial formation of disulfides, mixed disulfides with protein thiols or glutathione, or nitrosylated residues. The number of bands is related to the amount of thiols originally present in each thioredoxin isoform. As shown in Fig. 5 and consistently with the results of Fig. 4 A, complex 2 appears to be completely oxidized (dimeric form) after treatment of cells with 2. Complex 1, even at higher concentrations is far less effective than 2 as, although inducing some oxidation of Trx1 and Trx2, it appears unable to stimulate the oxidation of Prx3.



Fig. 5. Jurkat cells (2x10⁶), treated for 18 h with **1** and **2**, were derivatized with 10 mM IAM to titrate free SH groups. Then, after treatment with DTT (3.5 mM), an excess of IAA (30 mM) was added, giving rise to derivatized groups corresponding to the oxidized residues. The determination of redox state of Trx1, Trx2 was performed with urea-PAGE in non reducing conditions and the bands, detected with WB analysis, correspond to the different oxidation states of Trx1 or Trx2. For the estimation of the redox state of Prx3, cell lysates were derivatized with 10 mM AIS to alkylate free thiols and subjected to SDS-PAGE in non reducing conditions as described in Materials and methods. Dimer corresponds to disulfide-linked oxidised form while monomer refers to the reduced form.

3.8. Effect of 1 and 2 on mitochondrial membrane potential (MMP)

In order to confirm that mitochondria are a key target of the antiproliferative activity observed with 2, the effects of 1 and 2 on the mitochondrial membrane potential of Jurkat cells were studied. As seen in Fig. 6, complex 2, even at the lower concentration (10 μ M), is markedly more effective than 1 in decreasing mitochondrial membrane potential (MMP) as shown by the decline of fluorescence intensity (Fig. 6 and Table 1).



Fig. 6. Mitochondrial membrane potential (MMP) of Jurkat cells in the presence of **1** and **2**. Cells were treated for 18 h with **1** (15 and 50 μ M) or **2** (10 and 15 μ M) and MMP was measured using TMRM and detected by flow cytometry analysis. P1 corresponds to cell population with low membrane potential, while P2 is related to cell population with high membrane potential. a, control; b, 15 μ M **1**; c, 50 μ M **1**; d, 10 μ M **2**; e, 15 μ M **2**; f, 1 μ M CCCP used a positive control.

Table 1: Percentage of cells with high and low mitochondrial membrane potential (MMP) after treatment with 1 or 2. Jurkat cells were treated with 50 μ M 1 or 15 μ M 2 in the conditions described above. Data are reported as mean \pm SD of five experiments.

	high MMP (%)	low MMP (%)
Control	82 ± 5	18 ± 5
1	61±11	39±11
2	3 ± 0	98± 1

4. DISCUSSION

The results of the different tests performed with osmocifens **1** and **2** *in vitro* on purified thioredoxin reductase and *in cellulo* on Jurkat cells are summarized in Table 2.

	1	2	
	Inhibition of cytosolic TrxR (IC ₅₀ in µM)		
Complex alone	≥ 20	> 20	
Complex pre-treated with HRP/H ₂ O ₂	2.4 ± 0.3	1.2 ± 0.1	
Quinone methide	5.4 ± 0.7	3.6 ± 0.1	
	Tests on Jurkat cells		
IC ₅₀ values of MTT Test (µM)	42 ± 4	7.4 ± 0.8	
ROS production (20 µM of 1-2)	+	+++	
	Effect of 50µM of 1	Effect of 15µM of 2	
% of TrxR inhibition in cell lysates	32.8 ± 1.2	68.4 ± 1.2	
Trx1 redox state	40% oxidised	95% oxidised	
Trx2 redox state	50% oxidised	82% oxidised	
Prx3 redox state	Mainly reduced (60% monomer form)	Completely oxidised (98% dimer form)	
GR activity in cell lysates	slight increase	slight increase	
% of cells with high MMP	61 ± 11	3 ± 0.1	

Table 2 : Summary of the biological studies performed with 1 and 2 in vitro or on Jurkat cells.

The results of the *in vitro* study on the inhibition of purified TrxR clearly show that complexes 1 and 2 have no inhibitory effect until after their enzymatic oxidation by the H₂O₂/HRP system and that the inhibition observed is very similar for the two complexes (IC₅₀ = 2.4 and 1.2 μ M). This inhibition is stronger than that found after incubation in the presence of the quinone methides 1-QM and 2-QM (IC₅₀ = 5.4 and 3.6 μ M). In addition, the UV-VIS spectra of 1* and 2*, the species obtained by treatment of 1 and 2 with HRP/H₂O₂, do not correspond to those of their quinone methides (Fig. 2) as it was the case in the ferrocifen series [32]. This result should also be viewed from the perspective of the recent observation that the electrochemical oxidation of 1 corresponds, in the presence of a base, to a 2-electron transfer but does not lead to the corresponding QM [26]. Looking again at the oxidation pathway previously established for the ferrocifens, the hypothesis can be made that in the case of the complex, the removal of the phenolic proton and a second oxidation, could be the reactive species responsible for the inhibition of TrxR (Scheme 2). This corresponds to a cessation of the reaction sequence at an earlier step than that of neutral quinone methide. This hypothesis is in agreement with

the stabilization sequence of carbocations and radicals in the metallocene triad, in the order Fe < Ru < Os, with osmium being the most stabilized [38, 39].



Scheme 2: Hypothesis for the oxidation sequence of 1 and 2 (adapted from ref [4]).

If this hypothesis is correct, **1*** and **2***, i.e. the species generated by enzymatic oxidation by the HRP/H₂O₂ system of **1** and **2**, should have a reasonable chance to correspond to these quinone methide cations. In order to check this assumption we tried to precipitate **1*** (which is more stable than **2*** cf. Fig. S3 and Fig. S4) by addition of BPh₄⁻, an anion previously used to get single crystals of osmocenyl carbenium ion [39]. This experiment was performed as follows: **1*** was generated by incubation of **1** in the presence of a mixture of HRP/H₂O₂ for 20 min. Then addition of 2 equivalents of NaBPh₄ afforded a yellow precipitate that was recovered by centrifugation, washed, then dissolved in methanol leading to a bright yellow solution (see Material and methods). The UV-VIS spectrum of this solution is similar to that obtained for **1*** (Fig. S5). Electrospray mass spectroscopic analysis of the solution gave a main molecular ion at m/z = 543.5 amu in the positive mode and an isotope pattern matching that of **1-QM⁺**. In the negative mode, the molecular peak was at 319 amu, corresponding to the BPh₄⁻ counterion. (Fig. S6). Thus this hypothesis of a reactive quinone methide carbocation in the osmocifen series appears plausible.

The series of experiments performed *in cellulo* on Jurkat cells shows that in the majority of trials performed, complexes **1** and **2** behave in different ways. This is particularly true for their cytotoxicity. While the phenolic complex **1** has low toxicity ($IC_{50} = 42 \mu M$), the tamoxifen-like complex **2** exhibits a high level of cytotoxicity ($IC_{50} = 7.4 \mu M$). Interestingly, this difference in cytotoxicity can be compared with the efficiency of inactivation of TrxR by the complexes in Jurkat cells. In fact, while **2** induces very strong inhibition of TrxR (70% inhibition for 15 μ M incubation), that produced by **1** is much more modest (30% inhibition at 50 μ M incubation). This result explains the

redox state of Trx1, Trx2 and of Prx3 which are markedly oxidized upon treatment with **2**. This is likely a consequence of the inhibition of TrxR by oxidation products of **2** hampering the electron flow delivery to Trx and hence to Prx3. Consequently, the shift of Trx1/2 and Prx3 toward a more oxidized condition, associated to an increase of ROS, may contribute to the decrease of mitochondrial membrane potential (MMP) observed for cells incubated in the presence of **2**.

A large difference in toxicity between these two complexes has also recently been observed on MDA-MB-231 cells ($IC_{50} = 34 \mu M$ for **1** and 2.7 μM for **2**) [26]. In the citation given, the parameters studied to explain these differences in reactivity (lipophilicity, solubility and speed of formation of quinone methides, difference in the acidity of the phenolic protons), did not permit a satisfactory explanation of the difference in their cytotoxicity. Here, TrxR appears as a privileged target to explain the high cytotoxicity of **2** as well as the difference in cytotoxicity between **1** and **2**.

The fact that complex **2** induces strong inhibition of TrxR indicates that the latter undergoes oxidation in cells due to the presence of oxidizing systems such as cytochromes P450 (CYP 450). We have also recently shown that the oxidative metabolism of the ferrocifens by CYP 450 led to the formation of three principal metabolites, namely quinone methide, cyclic indene and allylic alcohol, which in some cases possess high cytotoxicity [28]. With the osmocifens we do not observe formation of quinone methides, but formation of other active metabolites could be envisaged, such as the quinone methide carbocation mentioned above. As for the difference in behavior between **1** and **2** observed in Jurkat cells, it is clearly associated with the presence of the $O(CH_2)_3N(CH_3)_2$ side chain. This may be related to a stabilization associated with the oxygen-donating mesomeric effect. A difference in the inhibition of TrxR activity has also been found in the ferrocifen series after incubation with Jurkat cells [32]. As here, the inhibition is more marked for the tamoxifen-like complex than for the phenolic, but it is not correlated with the antiproliferative effect of the complexes [32].

As far as production of ROS by Jurkat cells is concerned, complex 2 induces greater production than 1, which was also observed in the ferrocifen series [40]. This ROS production would be the source of the increase in activity of glutathione reductase. This absence of inhibition of GR also shows that the osmocifens 1 and 2 can only react with nucleophilic selenols and not with thiols, similarly to the ferrocifens.

In conclusion, the cytotoxicity of osmocifen compounds was dependent on its intracellular conversion to species able to interact with specific targets of cancer cells. TrxR appears as a pivotal site of inhibition that stimulates a series of events leading to the death of the cancer cells.

Abbreviations

AIS, 4-acetamido-4'-((iodoacetyl)amino)stilbene-2,2'-disulfonic acid; CCCP, carbonyl cyanide-mchlorophenylhydrazone; DTNB, 5,5'-dithiobis(2-nitrobenzoic acid); GR, glutathione reductase; HRP, horseradish peroxidase; CM-DCFH₂-DA, 5-(and-6)-chloromethyl-2',7'-dichlorodihydrofluorescein diacetate acetyl ester; IAA, iodoacetic acid; IAM, iodoacetamide; Prx, peroxiredoxin; ROS, reactive oxygen species; TAM, tamoxifen; TMRM, tetramethylrhodamine methyl ester; Trx, thioredoxin; TrxR, thioredoxin reductase; PMSF, phenylmethylsulfonyl fluoride.

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Supplementary information

Osmocenyl-tamoxifen derivatives target the thioredoxin system leading to a redox imbalance in Jurkat cells

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Fig. S1. UV-Vis spectra of 1 and 2 $(25\mu M)$ in 0.2M Tris HCl buffer, pH 8.1.



Fig. S2. UV-VIS spectra of 1 (A) and 2 (B) after treatment with HRP and H_2O_2 . Fresh solutions of 1 or 2 (25 µM in Tris HCl buffer (pH 8.1), 1 mM EDTA and 2.5% DMSO) were incubated at 25 °C in the presence of 22 nM HRP + 0.1 mM H_2O_2 . Kinetics of the reactions after addition of HRP and H₂O₂ was followed by UV-VIS spectroscopy between 250 and 450 nm, at different times from 1.5 to 11.5 min. Black lines represent the spectra of 1 or 2 alone (before addition of HRP and $H_2O_2)_{.}$ 28



Fig. S3. Time-dependent formation and stability of 1^* and 2^* , the complexes obtained by enzymatic oxidation of 1 and 2 by HRP + H_2O_2 . Formation of 1^* and 2^* is monitored by the increase of absorbance at 323 nm. Formation of 2^* is faster than that of 1^* (see Table S1). It reaches its maximum at 40 and 20 min respectively for 1^* and 2^* . 1^* is stable during at least 60 min while 2^* is less stable (7% decrease after 1 h).

Table S1. Kinetics of formation of 1^* and 2^* , the complexes obtained by enzymatic oxidation of 1 and 2 by HRP + H₂O₂. Comparison with the values obtained for 1-QM and 2-QM obtained by chemical oxidation of 1 and 2 with Ag₂O.

complex	Kinetics of formation of 1* or 2* a)		Kinetics of formation of 2- $QM^{b)}$	
	$k_{obs} (min^{-1})$	t _{1/2} (min)	$k_{obs} (min^{-1})$	t _{1/2} (min)
1	$0.105 \pm 0.01^{\circ)}$	$6.7\pm0.7^{^{c)}}$	-	-
2	$0.18 \pm 0.02^{^{\rm c)}}$	$4.0 \pm 0.4^{^{\rm c)}}$	0.029 ^{b)}	24 ^{b)}

a) Obtained by enzymatic oxidation with 22 nM HRP and 0.1 mM H_2O_2 in Tris HCl (pH 8.1), 1 mM EDTA. b) Obtained after chemical oxidation by Ag₂O; data from ref. [26]. c) Mean of two experiments.



Fig. S4. UV-visible spectra of 1 (A) and 2 (B) (25 μ M) (t = 0 min), and of 1* and 2* obtained by incubation of 1 and 2 in the presence of 22 nM HRP and 0.1 mM H₂O₂ in 0.2 M Tris HCl (pH 8.1) 1mM EDTA containing 2.5% DMSO after 80 and 590 min. Inset: evolution of the absorbance at 330 nm as a function of time.



Fig. S5. Superimposition of the UV-VIS spectra of 1^* after precipitation with NaBPh₄ and dissolution in *MeOH* (solid line) and of 1^* obtained under the experimental conditions of Fig. S2 (dashed line).





Fig. S6. Analysis by mass spectrometry (ESI-MS analysis) of the precipitate obtained after treatment of 1 by $HRP+H_2O_2$ followed by addition, after 20 min of NaBPh₄.

Compound	Amount of Os in the sample (in ppb) ^{a)}	Amount of Os in 10^7 cells (in nmol)	Amount of Os per cell (fmol)	LogPo/w ^{d)}
1 ^{b)}	796 ± 6	38.7	3.9	6.3
2 ^{c)}	855 ± 5	41.6	4.2	4.3

Table S2 : Quantification by ICP-OES of Osmium in 10^7 Jurkat cells after 24h of incubation

a) mean of 3 measurements, control < 50 ppb, analysis performed on a volume of 9.25 mL; b) concentration of incubation of 1 : 50 μ M; c) concentration of incubation 2 : 15 μ M; d) value from ref [26]





1, **2** and the compounds obtained after pre-treatment for 15 min of **1** and **2** with 0.1 mM H_2O_2 and 22 nM HRP (**1***, **2***), were incubated in 0.2 M Tris HCl buffer (pH 8.1) and 1 mM EDTA with 0. 25 mM NADPH and 60 nM TrxR1. After 5 min, TrxR1 activity was estimated as described in Materials and methods. Under the same experimental conditions the HRP/ H_2O_2 mixture is ineffective on TrxR1 activity in the absence of osmocifen compounds.



Fig. S8. Effect of compounds 1 and 2 on TrxR activity in HEK293 cell lysates. $1x10^6$ cells were treated with the indicated concentrations of 1 and 2 for 18 h. Cell lysates were tested for thioredoxin reductase activity with the DTNB procedure as described in Materials and methods.