Cyclohexa-2,5-diene-1,4-dione-based antiproliferative agents: design, synthesis, and cytotoxic evaluation

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Abstract

Background: Tumors are diseases characterized by uncontrolled cell growth and, in spite of the progress of medicine over the years, continue to represent a major threat to the health, requiring new therapies. Several synthetic compounds, such as those derived from natural sources, have been identified as anticancer drugs; among these compounds quinones represent the second largest class of anticancer agents in use. Several studies have shown that these act on tumor cells through several mechanisms. An important objective of this work is to develop quinoid compounds showing antitumor activity, but with fewer side effects. The parachinone cannabinol HU-331, is a small molecule that with its core 4-hydroxy-1,4-benzoquinone, exhibits a potent and selective cytotoxic activity on different tumor cell lines. A series of derivatives 3-hydroxy-1,4-benzochinones were thus developed through HU-331 chemical modifications. The purpose of the work is to test the ability of the compounds to induce proliferative inhibition and study the mechanisms of cell death.

Methods: The antitumor activities were evaluated in vitro by examining their cytotoxic effects against different human cancer cell lines. All cell lines tested were plated in 96-multiwell and treated with HU-100-V at different concentrations and cell viability was evaluated by MTT assay. Subsequently via flow cytometry (FACS) it was possible to assess apoptosis by the system of double labeling with PI and Annexin-V, and the effect of the compounds on ROS formation by measuring the dichlorofluorescein fluorescence.

Results: The substitution by n-hexyl chain considerably enhanced the bioactivity of the compounds. In details, 2-hexyl-5-hydroxycyclohexa-2,5-diene-1,4-dione (V), 2,5-Dimethoxy-3-hexyl-2,5-cyclohexadiene-1,4-dione (XII) and 2-hydroxy-5-methoxy-3-hexyl-cyclohexa-2,5-diene-1,4-dione (XIII) showed most prominent cytotoxicity against almost human tumour cell lines. Compound V was further subjected to downstream apoptotic analysis, demonstrating a time-dependent pro-apoptotic activity on human melanoma M14 cell line mediated by caspases activation and poly-(ADP-ribose)-polymerase (PARP) protein cleavage.

Conclusions: These findings indicate that 2-hexyl-5-idrossicicloesa-2,5-diene-1,4-dione can be a promising compound for the design of a new class of antineoplastic derivatives.

Keywords: Apoptosis, Cytotoxicity, Benzoquinones, Melanoma cell line M14, PARP, Reactive oxygen species

Introduction

Several benzoquinones have been found to be effective in the treatment of some forms of cancer; previous studies demonstrated that these drugs act on cells by numerous mechanisms, such as apoptosis, abrogation of the cell cycle, activation of caspases, stimulation of the production of reactive oxygen species (ROS), inhibition of topoisomerases I and II, activation of intracellular second messengers, and production of free radicals to attack DNA. However, their cumulative heart toxicity limits their use; therefore, an important goal of present and future work is to develop quinoid compounds that display anticancer activity but with less side effects.

Among the 1,4 benzoquinones, there are several naturally occurring quinones having potent anticancer activity.
A recent work [1] demonstrated that Ardisianone, a natural benzoquinone derivative, displayed anti-proliferative and apoptotic activities against human hormone-refractory prostate cancer cells (HRPC), PC-3, and DU-145. Previous investigations also showed that Primin isolated from the leaves of Miconia Lepidota present in Suriname forests, exhibited activity towards mutant yeast strains, indicative of their cytotoxicity and potential antitumor activity [2]. Furthermore Kaul and co-workers isolated a known cytotoxic quinine Irisoquin which demonstrated cytotoxic properties [3]. In previous reports Muhammad et al. [4] evaluated cytotoxic and antioxidant activities of alkylated benzoquinones from the leaves of Maesa Lanceolata. Moreover, another important 2,5-dihydroxy-3-alkyl-1,4-benzoquinone is Embelin known for a very long time in Indian and Chinese pharmacopoeias. It is the major constituent in the extracts of various parts of the shrub Embelia Ribes. Embelin and its derivatives possess analgesic, anti-inflammatory, antioxidant, antitumor and antifertility properties [5-7].

Important results have been described with HU-331, that exhibited potent and selective cytotoxicity against several tumorigenic cell lines such as Burkitt's lymphoma, glioblastoma, breast, prostate and lung cancer [8]. Recent findings described that this derivative is strongly antiangiogenic at concentration as low as 300 nM by directly inducing apoptosis of vascular endothelial cells [9-11].

As a part of our research program devoted to the preparation and evaluation of new antiproliferative agents, [12-14] the para-quinone of cannabinol HU-331 (1) was selected as biologically validated starting point for compound library development, in order to evaluate the structural requirements important for biological activity and in particular the role of the substituents linked to the quinone nucleus.

We prepared compounds analogues whose structure closely resembles the natural compound, thus the 2-hydroxy 1,4 benzoquinone core was not changed.

**Methods**

**Chemistry**

Compounds I-V (series I) retain the same hydroxy-1,4-benzoquinone core of lead, modifications were carried out on the alkyl chain that was elongated and shifted, cycloalkenyl substituent in position 2 of HU-331 was removed (compound V) or replaced by a cyclohexyl (I and II) or by a cyclohexylmethyl moiety (III and IV).

On the other hand, to evaluate the influence of position of alkyl chain and hydroxy group on the 1,4-benzoquinone nucleus, compounds VI, VII and VIII were prepared. In parallel, we studied the variation of the cytotoxicity in a series of 2,5-dihydroxy-3-alkyl-1,4-benzoquinone system (series II). These compounds (IX-XIV) are characterized by a butyl or hexyl chain in position 3 of quinone ring which is 2,5 disubstituted with hydroxy or methoxy groups (Figure 1).

Compounds I and II were prepared starting from commercially available 1,3-dimethoxybenzene (2a) and 1-hexyl-2,4-dimethoxybenzene (2b) that were easily prepared according to procedure described by Kikuchi and co-workers [12-14]. Condensation of cyclohexanone with 2a-b gave the tertiary alcohols 3a-b in 70% and 80% yields respectively.

In order to remove their hydroxyl groups, 3a-b were submitted to the Barton-McCombie procedure, an extremely useful method with widespread application in synthetic organic chemistry. Compounds obtained were oxidized into quinoid compounds I (65% yield) and II (60% yield). Deprotection and final oxidation in air under basic conditions, led to the formation of the desired quinones III and IV in 55% and 60% yield, respectively.

The choice of the effective oxidizing agent and conditions used in the last step of both the synthetic schemes, derived from a set of experiments applied to the oxidation of the commercially available model compound 4-hexylresorcinol 10 to V, as shown in Table 1.

For example the conversion of 10 to V was tried in the presence of Salcomine, a coordination complex derived from the salen ligand and cobalt.

This catalyst is known to catalyze the oxidation of 2,6-disubstituted phenols by dioxygen but in our case a complete conversion of the starting material 10 in o-hydroxyquinone was observed (Entry 1-2).

In another attempt, using a catalytic amount of ethanol in air, a solution of 5% of KOH as base and

![Figure 1 Development of compounds of general formula A and B.](image-url)
petroleum ether as solvent (Entry 3), only a little quantity of starting material was converted into quinone V. For these reasons Fremy’s salt (disodium nitrosodisulfonate) was tried as oxidizing agent, under several conditions (Entry 4–7). However, in the presence of Na₂CO₃ as base and a mixture of H₂O/THF as solvent, was obtained the best results (Entry 7) [15].

Ag(I)-promoted Suzuki–Miyaura cross-coupling of 1-bromo-2,3,5-trimethoxy benzene (11) and hexylboronic acid pinacol ester furnished intermediate 12 in 20% yield. Deprotection and simultaneous oxidation to 2-hydroxy-5-benzoquinone (VI) was achieved treating 12 with an excess of BBr₃. Oxidation with ammonium cerium nitrate (CAN) in a mixture of acetonitrile and water allowed to obtain 2-methoxy derivative VII. Compound VIII was synthesized starting from 1,3-dimethoxy benzene (2a) which was subjected to a ortho-metalation reaction in presence of n-BuLi. Coupling reaction of 1,2,4,5-tetramethoxybenzene with two different alkyl iodides, in presence of n-BuLi and hexamethylphosphoramide (HMPA) furnished intermediates 15 and 16 in moderate yields (42% and 37% respectively). CAN-mediated oxidative reaction provided a mixture of 2,5-dimethoxy (IX and XII) and 2-hydroxy-5-methoxy derivatives (X and XIII). Treatment of IX and XII with 2M NaOH allowed to obtain 2,5-dihydroxy derivatives XI and XIV in good yields (72% and 89%).

**General procedures**

All reagents were analytical grade and purchased from Sigma-Aldrich (Milano-Italy). Flash chromatography was performed on Carlo Erba silica gel 60 (230÷400 mesh; Carlo Erba, Milan, Italy). TLC was carried out using plates coated with silica gel 60F 254 nm purchased from Merck (Darmstadt, Germany). Melting points were determined in open capillary tubes on a Electrothermal 9100. ¹H and ¹³C NMR spectra were registered on a Bruker AC 300. Chemical shifts are reported in ppm. All target compounds were assessed for purity by analytical high performance liquid chromatography (HPLC), using an Agilent 1100 series, UV detector monitoring at 254 nm, HP Chem Station software, and a Waters C18 RP-column (5 μm, 300 mm × 3.9 mm). All target compounds were found to be >95% purity. MS spectrometry analysis ESI-MS was carried out on a Finnigan LC Decaion trap instrument. Microanalyses were carried out on Carlo Erba 1106 elemental analyzer.

**Biological studies**

**Cell culture**

Our experimental models consist of several cell lines derived from human cancers of different histogenesis. The cells were grown in RPMI or DMEM supplemented with heat inactivated 10% FBS, 20 mM HEPES, 100 U/ml penicillin, 100 μg/ml streptomycin, 1% L-glutamine in a humidified atmosphere of 95% air/5% CO₂ at 37°C [16].

Analysis of cell proliferation was performed in the presence of all derivatives on all cell lines seeded in 96-well plates at the different densities depending on the cell type. Pancreas cancer cell lines ( BXPC3, Panc-1) were plated to the average density of 3,600 cells/ well. Prostate cancer cell lines (DU145, PC3, LNCAP) were plated to the average density of 2,000 cells/ well. Melanoma cell lines (COLO38, A375, M14) were plated to the average density of 1,800 cells/ well. Renal cancer cell lines (A498, RXF393, SN12C, 769P) and glioblastoma cell lines ( LN229, U87 MG, U373 MG) were plated to the average density of 1,900 cells/ well. Breast cancer cell lines (CG5, MCF-7, MDA-MB 231, MDA-MB 468, MDA-MB 436 ) were plated to the average density of 3,100 cells/ well. After 24 h incubation at 37°C, the cells were treated with increasing concentrations of compounds (0,037-50 μM). Cells were incubated under these conditions for 72 h.

**MTT bioassay**

Human cancer cells (3 × 10³) were plated in 96-well culture plates in 90 μL of culture medium and incubated at 37°C in humidified atmosphere of 5% CO₂. The day after, 10 μL aliquot of serial dilutions of compounds (1–50 μM) was added to the cells and incubated for 72 h. The cell viability was assessed with MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide] method [17]. After 72 h of treatment with derivatives MTT solution 5 mg/ml in PBS was added to each well. The plates were then incubated at 37°C for an additional 4 h to allow MTT to form formazan crystals by reacting with metabolically active cells. The formazan crystals were solubilized in a 1N isopropanol/HCl 10% solution at 37°C, on a shaking table for 20 min. The absorbance values of the solution in each well were measured at 570 nm using a micro plate reader.
Cell viability was determined by the formula:

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\text{cell viability (\%)} = \frac{(\text{absorbance of the treated wells} - \text{absorbance of the blank control wells})}{(\text{absorbance of the negative control wells} - \text{absorbance of the blank control wells})} \times 100\%
\]

as previously reported [18]. All MTT experiments were performed in quadruplicated and repeated at least three times. Data are as mean ± standard deviation (SD). Each IC\(_{50}\) mean value was obtained from four independent experiments.

**Apoptotic cell death**

Hypodiploid DNA was analysed using the method of propidium iodide (PI) staining and flow cytometry as described [19]. At the end of incubation with HU compounds, with or without pretreatment with pan-caspase inhibitor Z-vad-fmk, purchased from BD Pharmingen (BD Bioscience, Bedford, USA), cells were washed in phosphate-buffered saline (PBS) and resuspended in 500 μL of a solution containing 0.1% sodium citrate, 0.1% Triton X-100 and 50 μg/ml propidium iodide (Sigma-Aldrich, Italy). After incubation at 4°C for 30 minutes in the dark, cell nuclei were analyzed with Becton Dickinson FACScan flow cytometer using the Cells Quest program. Cellular debris was excluded from analysis by raising the forward scatter threshold, and the DNA content of the nuclei was registered on logarithmic scale. The percentage of the cells in the hypodiploid region was calculated [20].

**Western blotting analysis**

Total intracellular proteins were extracted from the cells by membrane disruption in lysis buffer 50 mM Tris-HCl, 1% Na-deoxycholate, 1% SDS and 0.5% IGEPA Let (All from Sigma-Aldrich, Gallarate, Italy) containing protease and phosphatase inhibitors (1mM PMSF, 1 μg/mL leupeptin, 1μg/mL pepstatin, 1 μg/mL aprotinin, 1 μM Na\(_3\)PO\(_4\), 1 μM NaF; all from Sigma-Aldrich, Gallarate, Italy) on ice for 20 min. The cell lysate was then centrifugated at 10,000 \(\times g\) at 4°C for 15 min. The supernatant was collected as protein extract. Protein content was estimated according to Biorad protein assay (BIO-RAD, Milan, Italy) and the samples either analysed immediately or stored at -80°C. Total protein (30 μg) samples were loaded into a 10-12% acrylamide gels and separated by SDS-PAGE in denaturing conditions at 150 V.

The separated proteins were then transferred electrophoretically (100 mA per blot 90 min; Trans Blot Semi-Dry, BIO-RAD) to nitrocellulose paper (Immobilon-NC, Millipore, Bedford, USA) soaked in transfer buffer (25 mM Tris, 192 mM glycine, Sigma-Aldrich) and 20% methanol vol/vol (Carlo Erba, Milan, Italy) [21]. Non specific binding was blocked by incubation of the blots in 5% no fat dry-milk powder (BIO-RAD) in TBS/0.1% Tween (25 mM Tris; 150 mM NaCl; 0.1% Tween vol/vol, Sigma-Aldrich) for 60 min. After washing, the blots were incubated overnight at 4°C with the following primary antibodies: mouse monoclonal anti-PARP? (diluted 1:1,000) and anti-XIAP (all from Santa-Cruz Biotechnology, Santa Cruz, CA). After incubation with the primary antibodies and washing in TBS/0.1% Tween, the appropriate secondary antibody, either anti-mouse (diluted 1:5,000), or anti-rabbit (diluted 1:5,000) (both from Sigma-Aldrich, Italy) was added for 1h at room temperature. Immunoreactive protein bands were detected by chemiluminescence using enhanced chemiluminescence reagents (ECL) and exposed to Hyperfilm (both from Amersham Biosciences, Italy). The blots were then scanned and analysed (Gel-Doc 2000, BIO-RAD).

**Flow cytometric estimation of intracellular redox state**

The effect of all compounds on intracellular reactive oxygen species (ROS) was evaluated by measuring dichlorofluorescein (DCF) fluorescence [22]. Cells were incubated in presence and absence of compounds. At the end of incubation time, cells were washed and resuspended (2 × 105 cells/ml) in Hank’s balanced salt solution (HBSS) cointaining 10 μM 2′,7′-dichlorodihydrofluorescein diacetate (DCFH-DA). Following a further 20 min incubation at 37°C, DCF fluorescence was monitored by flow cytometry (FL1-H channel). In order to estimate the antioxidant potential of the compounds, control and treated cells were exposed to 300 μM of the oxidant tert-butylydroperoxide (t-BOOH) for 30 min at 37°C before DCFH-DA loading.

**Topoisomerase I-Mediated DNA cleavage reactions**

Human recombinant Top1 was purified from Baculovirus as previously described [23]. DNA cleavage reactions were performed using a 22-bp DNA oligonucleotide with a prominent Topoisomerase I cleavage site. Single-stranded oligonucleotide was labeled according to the manufacturers’ instructions by using terminal deoxynucleotidyltransferase (USB Corporation, Cleveland, OHIO) that adds a single labeled cordycepin molecule (γ-32P, 5000 Ci/mmol, PerkinElmer Life and Analytical Sciences, MA) to the 3’ terminus. Unincorporated nucleotides were removed by QIAquick Nucleotide Removal Kit (Qiagen, Hilden, Germany). The duplex DNA oligonucleotide was annealed by addition of an equal concentration of the
complementary strand, heated to 95°C and slow cooled to room temperature. For the Topoisomerase I cleavage reaction, DNA oligonucleotides were reacted for 20 min at 25°C with a 12 ng/mL solution of human Topoisomerase I and the desired amount of drugs, in 10 mM Tris–HCl pH 7.5, 50 mM KCl, 5 mM MgCl₂, 0.1 mM EDTA and 15 μg/mL bovine serum albumin. Reactions were stopped by adding 0.5% SDS and formamide containing 0.25% bromophenol blue and xylene cyanol, heated at 95°C for 5 min and chilled on ice. Reaction products were separated in 20% polyacrylamide denaturing sequencing gels. Dried gels were visualized using a B40 Storm phosphor imager (Amersham Biosciences, GE Healthcare, UK).

**Topoisomerase II-Mediated DNA cleavage reactions**

DNA was purchased from Invitrogen Corporation (Carlsbad, CA). It represents a portion of SV40 sequence, in particular from position 3449 to 3538, that contains prominent topoisomerase II cleavage sites [24]. DNA was purified on denaturing 20% polyacrylamide gel, recovered by soaking gel slices in water and then ethanol precipitated. Single-stranded DNA was 5′-labeled using T4 polynucleotide kinase (New England Biolabs, Ipswich, MA) with [γ-32P]ATP (3000 μCi/mmol, PerkinElmer Life and Analytical Sciences, MA) according to the manufacturers’ instructions. Unincorporated nucleotides were removed by QIAquick Nucleotide Removal Kit (Qiagen, Hilden, Germany). The duplex DNA was annealed by addition of an equal concentration of the complementary strand, heated to 95°C and slow cooling to room temperature. For the topoisomerase II cleavage reaction, DNA fragments were reacted for 30 min at 37°C with 30 units/sample of human topoisomerase II, purified as previously described, [25] and the desired amount of drugs, in 40 mM Tris–HCl, pH 7.5, 80 mM KCl, 10 mM MgCl₂, 5 mM DTT, 1 mM ATP, and 15 μg/mL bovine serum albumin. Reactions were stopped by adding 1% SDS and 0.2 mg/mL proteinase K and incubated at 42°C for 45 minutes. Samples were then ethanol precipitated, resuspended in 10 μL of formamide containing 0.25% bromophenol blue and xylene cyanol, heated at 95°C for minutes and chilled on ice. Reaction products were separated in 20% polyacrylamide denaturing sequencing gels. Dried gels were visualized using a B40 Storm phosphor imager (Amersham Biosciences, GE Healthcare, UK).

**Statistical analysis**

All results are shown as mean ± SEM of three experiments performed in triplicate. The optical density of the protein bands detected by Western blotting was normalized on β-actin levels. Statistical comparison between groups were made using ANOVA followed by Bonferroni parametric test. Differences were considered significant if P < 0.05.

**Results and discussion**

**Biological evaluation**

For the evaluation of cytotoxicity, five different human cancer cell lines were utilized: M14 and A375 (melanoma cell lines), MCF-7 (human breast cancer cell), PC3 (human prostate cancer cell line), A498 (human renal cancer cell line). The survival percentage was determined after a period of 72 h at screening concentrations from 50 to 1 μM, using the survival percentage obtained from the cells treated only with the solvent (DMSO at 0.5%) as a reference. Our experiments confirmed the cytotoxic activity of HU-331. Most of the compounds displayed moderate cytotoxicity against cancer cell lines in relatively lower micromolar concentrations when compared to the standard. Among the compounds, derivatives V, IX, XII and XIII showed significant cytotoxicity in most of the cell lines, displaying similar or slightly weaker potency than positive control. Compound V can be considered the most interesting compound that showed a good anticancer activity against all tumor cell lines and was more potent than HU-331 against M14 (7 μM vs 15 μM). The structure-activity relationship studies regarding the first series of compounds revealed that the n-hexyl chain in position 5 of the hydroxy-quinone ring was fundamental for the anticancer activity (compounds II, IV and V), in fact compounds I and III, which lacked of the alkyl chain, were completely inactive. At the same time, the change of position of alkyl chain was clearly detrimental (VI, VII and VIII vs V). No relevant influence on the activity was observed if a methylene spacer was inserted between cyclohexyl and 1,4-benzoquinone ring (IV vs II). As concern for compounds of series II, the 5-methoxy-1,4-benzoquinone derivatives IX, XII and XIII were the most active compounds of the series, while compound X was slightly active only against M14 cell line. Considering the importance of the length of the side chain, the hexyl chain seemed to be more favorable for the activity but further investigations will be done. 2,5-dihydroxy-1,4-benzoquinone derivatives XI and XIV were completely inactive against all tumor cell lines at the concentration of 100 μM. Structure-activity relationship evaluations, comparing compounds of first and second series, demonstrated that the introduction of a methoxy (XII) or hydroxy (XIII) group on the 1,4-benzoquinone ring of compound VII caused a strong improvement in the cytotoxicity against almost tumor cell lines, except A498. On the contrary, if another hydroxy group was inserted on the quinone core of compound VI, no improvement of activity was recorded (compound XIV). Having identified, from the first screening, the most cytotoxic compound against all tumor cell lines, we have carried out a screening on other solid tumor cell lines to confirm the cytotoxic activity of this molecule. Moreover, we investigated the molecular mechanisms underlying the antiproliferative activity in comparison with the natural compound HU-331 on M14 cells. However all data are reported in Table 2. The MTT
viability assay showed that compound V has good antiproliferative properties against all tested solid human cancer cell lines (Table 3).

**In vitro cytotoxicity**

The cytotoxicity of HU-100-V was evaluated on different cell lines derived from different tumors. In all treatments the cells were plated in 96-multiwell in the presence of increasing concentrations of the drug, (0.37 to 50 μM). The MTT viability assay showed that HU 100-V decreased the viability of most of the cell lines tested in a time- and dose-dependent manner for which they are achieved good values of IC50 (concentration inhibiting 50% of growth). Especially, prostate cancer DU-145, pancreas cancer BX-PC3 [26,27], renal cancer RXF393 and glioblastoma cancer LN229 cells have proved to be the most sensitive to this treatment, with IC50 values of less than 20 micromolar (Table 3 and Figure 2).

**Apoptotic cell death**

To ascertain whether loss of cell viability was mediated by effects on apoptosis we directly analyzed the effects

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**Table 2 Effects of HU compounds on proliferation of several cancer cell lines**

| Cpd | R1  | R2  | R3  | R4  | M14     | MCF-7     | PC3     | A498    | A375    |
|-----|-----|-----|-----|-----|---------|-----------|---------|---------|---------|
| I   | H   | H   | H   |     | >100    | >100      | >100    | >100    | >100    |
| II  | n-hexyl | H | H   |     | 23 ± 0.12 | 28.13 ± 0.07 | 41 ± 0.20 | 34.91 ± 3.82 | >100    |
| III | H   | H   | H   |     | >100    | >100      | >100    | >100    | >100    |
| IV  | n-hexyl | H | H   |     | 45.6 ± 0.20 | 37.3 ± 0.34 | 38 ± 0.12 | **28.8 ± 0.04** | 30.7 ± 0.12 |
| V   | n-hexyl | H | H   | H   | 7.0 ± 0.10 | **18.7 ± 0.06** | **24.3 ± 0.20** | 19.8 ± 0.02 | **12.9 ± 0.06** |
| VI  | H   | n-hexyl | H | H   | -        | >100      | >100    | >100    | >100    |
| VII | H   | n-hexyl | CH3 | H   | -        | >100      | >100    | >100    | >100    |
| VIII| H   | H   | CH3 | n-hexyl | -        | >100      | >100    | >100    | >100    |
| IX  | -CH3 | n-butyl | CH3 | H   | 24.5 ± 0.15 | **12 ± 0.03** | **17.9 ± 0.20** | 51 ± 0.02 | **17.6 ± 0.05** |
| X   | H   | n-butyl | CH3 | H   | 35 ± 0.64 | >100      | >100    | >100    | >100    |
| XI  | H   | n-butyl | H   | H   | >100    | >100      | >100    | >100    | >100    |
| XII | -CH3 | n-hexyl | CH3 | H   | **10.7 ± 0.15** | **16.2 ± 0.03** | **18.8 ± 0.03** | >100    | **21.0 ± 0.04** |
| XIII| H   | n-hexyl | CH3 | H   | **14.1 ± 0.15** | **13.9 ± 0.04** | **20.1 ± 0.20** | >100    | **18.1 ± 0.04** |
| XIV | H   | n-hexyl | H   | H   | >100    | >100      | >100    | >100    | >100    |
| H331|      |      |      |      | 15.0 ± 0.09 | 24.5 ± 0.15 | 32.0 ± 0.15 | 34.6 ± 0.23 | 21.8 ± 0.03 |

Cell viability was assessed through MTT assay. Data represent the mean ± SD values of three independent determinations performed in triplicate. A375, M14, human melanoma cells; MCF-7, human breast cancer cells; PC3, Human prostate cancer cell line, A498, Human renal cancer cell line.

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**Table 3 Cytotoxic activity of compound V in solid human cancer cell lines**

| Cell lines | IC50 (μM) |
|------------|-----------|
| Prostate   | LN-CAP    |
|            | DU-145    | 15.2  |
|            | BX-PC3    | 19.8  |
| Renal      | SN12C     | 31.6  |
|            | RXF393    | 23.6  |
|            | 769P      | 34.6  |
| Glioblastoma| LN229    | **18.2** |
|            | U373MG    | 23.6  |
|            | U87MG     | 30.8  |
| Breast     | CG-5      | 34.6  |
|            | MDA-MB 231| 33.6  |
|            | MDA-MB 468| 41.2  |
|            | MDA-MB 436| 40.1  |
of either V or HU-331 on apoptosis of M14 cells by using PI-staining of DNA fragmentation after cell permeabilization. Cells were treated with different concentrations (1–10 μM) of V and HU-331 for 24 and 72 hours and then the population of sub-G1 cells (hypodiploid nuclei) was determined. Compound V induced apoptosis of M14 cells in a concentration-dependent manner with 40% of cell death at 10 μM after 72 h, whereas a small pro-apoptotic effect was observed with 10 μM HU-331 (Figure 3). These results showed that the cytotoxic effect of V is dependent by an apoptotic mechanism that is more significant than HU-331 effect on M14 cells.

Caspases involvement
To investigate the involvement of caspases in the mechanism of apoptosis induced by compounds, we pretreated the cells with a pan-caspase inhibitor Z-VAD-fmk for 30 min before to add V and HU-331. Results in Figure 4 show that apoptosis induced by V in presence of the inhibitor was significantly reduced indicating the involvement of caspases in the apoptotic mechanism in M14 cells.

Cell cycle analyses
The cell cycle is divided into four phases, i.e. sub-G1, G1, S and G2. The sub-G1 represents the apoptotic cells whereas the G1 represents the first phase of cell cycle in which cell prepares itself for DNA synthesis. The cell cycle assay was performed through tagging the DNA with the PI dye as explained in “Materials and Methods”. M14 cells were plated in 6-well tissue culture plates. The cells were induced with compound V (10 μM) and the standard HU-331 (10 μM) and analyzed on a FACScan instrument using CELLQuestPRO software after time intervals of 24 h and 72 h. Cell cycle phases were compared in treated and untreated samples. No effect for either HU-331 or V was observed on cell cycle distribution of melanoma cells (data not shown).

Intracellular pathway involvement
Evasion from apoptosis is one of the hallmarks of human cancers contributing to tumor formation and treatment...
resistance. The alterations in apoptosis signaling pathway often occur in drug-resistant cancer cells. In particular, defective apoptosis signaling may be caused by an increase in content of anti-apoptotic molecules and/or by a decreased content or impaired function of pro-apoptotic proteins. Thus, identification of novel substances for overcoming the drug resistance has gained much attention in cancer therapy. The drug resistance of cancer cells is a complex phenomenon comprising different intracellular processes. It was described for doxorubicin that short-term-treated CEM cells gradually developed drug resistance. In particular, caspases activation, and XIAP and PARP cleavage were blocked. Thereafter, we evaluated the effect of the active apoptotic concentrations on expression of X-linked Inhibitor of Apoptosis Protein (XIAP) and Poly (ADP-ribose) polymerase (PARP) proteins. Cells were treated with V and HU-331 at 10 μM for 24 h and then the expression of XIAP and cleavage of PARP were analyzed by western blotting. Results in Figure 5 show that apoptotic effect of V was due to PARP cleavage that leads to inactivation of this protein, importantly involved in DNA repair. No effect on PARP cleavage was observed with HU-331 treatment. We also showed (Figure 6) that V was able to abolish XIAP protein levels whereas a little effect was observed in reduction of XIAP expression after HU-331 treatment.

**ROS involvement**

The quinoid anticancer agents undergo enzymatic reduction via one or two electrons to give the corresponding semiquinone radical or hydroquinone. Under aerobic conditions the semiquinone radical anion can give its extra electron to molecular oxygen to give the parent quinone and superoxide radical anion. This reaction sequence, is known as redox-cycling, and it continues until the system becomes anaerobic. Both the semiquinone and the superoxide radical anion can generate the hydroxyl radical, which is the cause of DNA strand breaks [28]. HU-331 induced cell death of human cancer cell lines is not mediated by reactive oxygen intermediates/species, as exposure to HU-331 failed to elicit the generation of reactive oxygen species. To assess the involvement of free radicals in V-mediated cell death we measured the production of reactive oxygen species (ROS) after exposure of compound at different times (5–120 min) by FACS analysis of DCFH-DA fluorescence intensity. Treatment with V increased intracellular ROS levels at early time point after 5 minutes of treatment with maximal effect after 30 min (Figure 5A), while we can confirm that the effect of HU-331 was very poor on ROS intracellular production (Figure 5B).

**Topoisomerase inhibition**

To determine topol catalytic activity, assays were carried out with supercoiled pBR322 DNA as the substrate according to protocol. Camptothecin (CPT) was the...
reference compound for Top1-mediated DNA cleavage reactions. Test derivatives did not increase DNA cleavage levels. Similar results were obtained with Top2. Mitonafide was the reference compound for Top2-mediated DNA cleavage reactions. The findings show that none of the assessed compounds are poisons of human Top2, thus their cellular effects can likely be due to a molecular mechanism different from topoisomerase poisoning (data not shown).

**Conclusion**

Natural benzoquinone compounds are a rich source for modern, molecular targeted-specific drug discovery [29]. Over the years, a great amount of efforts have been spent to isolate individual compounds and screen for anti-cancer activity. Previous research has demonstrated that HU-331 in vivo was more active and less toxic than doxorubicin [10,11] and thus represents a promising lead compound for designing a new class of anti-cancer treatments. The aim of this study was to check the cytotoxicity of novel synthetic 1,4 benzoquinone compounds. The new derivatives together with the natural lead were tested for their anti-proliferative activity against five cancer cell lines. The general trend on which the design of these structure is based has proven to be valid in obtaining new interesting compounds. In particular, 2-hexyl-5-hydroxycyclohexa-2,5-diene-1,4-dione (V) resulted the best synthesized compound; therefore, it was further subjected to downstream apoptotic analysis. Our study demonstrated a time-dependent pro-apoptotic activity of compound V. We determined that cell death of M14 induced by V is mediated by caspases activation and poly-
(ADP-ribose)-polymerase (PARP) protein cleavage. In addition we showed that HU-331 does not elicit the production of ROS while apoptosis induced by compound V could be activated by production of ROS observed after 30 min of treatment in M14 cells. We can suppose that the quinone structure permits to this compound to act as electron acceptor of free radicals that may kill cancer cells. To better define the possible mechanism of action of compounds, we also examined their dose-dependent effect on topoisomerases, as HU-331 has been proposed to be a catalytic inhibitor of topoisomerase II. We tested their ability to directly inhibit topoisomerases in cleavage assays demonstrating that our derivatives are not able to poison the nuclear enzymes. To conclude, the analyses of the present study have revealed that the synthesized quinone V has the potential to induce apoptosis in M14 cancer cell line in vitro and it is very important to note that this compound additionally has the ability to inhibit the expression of the antiapoptotic protein XIAP, a regulatory protein that suppresses apoptosis cell death by binding the caspase proteins [30,31]. On the light of interesting pharmacological results, a more extensive medicinal chemistry program has been engaged to consolidate the series and identify lead candidates for the design of more potent antitumor agents based on 2-hydroxyquinone skeleton which in turn should afford a better understanding of biological mechanisms regulating apoptosis.

Abbreviations
ROS: Reactive oxygen species; HRPC: Hormone-refractory prostate cancer cells; HMPA: Hexamethylphosphoramide; XIAP: X-linked inhibitor of apoptosis protein; PARP: Poly (ADP-ribose) polymerase; CPT: Camptothecin; DCF: Dichlorofluorescein.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RF and MC carried out the design of the experiments and drafted the manuscript. CP, MF, AP and MC participated in the experiments of cell culture and molecular biology. JM, AM, AG and GC, participated in statistical analysis and interpretation. ALG and MDR participated in the design of the experiments. All authors read and approved the final manuscript.

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