Monocyclic Quinone Structure-Activity Patterns: Synthesis of Catalytic Inhibitors of Topoisomerase II with Potent Anti-Proliferative Activity

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Abstract: The monocyclic 1,4-benzoquinone, HU-331, the direct oxidation product of cannabidiol, inhibits the catalytic activity of topoisomerase II but without inducing DNA strand breaks or generating free radicals, and unlike many fused-ring quinones exhibits minimal cardiotoxicity. Thus, monocyclic quinones have potential as anti-cancer agents, and investigation of the structural origins of their biological activity is warranted. New syntheses of cannabidiol and (±)-HU-331 are here reported. Integrated synthetic protocols afforded a wide range of polysubstituted resorcinol derivatives; many of the corresponding novel 2-hydroxy-1,4-benzoquinone derivatives are potent inhibitors of the catalytic activity of topoisomerase II, some more so than HU-331, whose monoterpane unit replaced by a 3-cycloalkyl unit conferred increased anti-proliferative properties in cell lines with IC₅₀ values extending below 1 mM, and greater stability in solution than HU-331. The principal pharmacophore of quinones related to HU-331 was identified. Selected monocyclic quinones show potential for the development of new anti-cancer agents.

Introduction
Quinones, especially derivatives of 1,4-benzoquinone, represent an important group of potent therapeutic agents,[1-4] especially as antibiotics and anticancer agents, including anthracyclines such as doxorubicin and daunorubicin,[5] as
well as other streptomyces-derived compounds such as streptonigrin and mitomycin C,[6] and synthetic quinonoids such as mitoxantrone and epirubicin.[7,8] The mechanisms by which anthracyclines and many fused-ring quinones act are complex, multiple pathways usually being involved, including intercalation between base pairs, resulting in the inhibition of RNA and DNA synthesis in cancer cells.[9,10] Secondly, quinones may act as inhibitors of the DNA topoisomerase II enzyme, preventing the relaxation of supercoiled DNA, thereby blocking transcription and replication.[11] Thirdly, they may generate haeme-mediated oxygen radicals that damage DNA, proteins and cell membranes,[12] although not necessarily selectively. Fourthly, anthracyclines have recently been shown to cause histone eviction from open chromatin structures, leading to activation of the DNA damage response or apoptosis of cancer cells.[13]

A major therapeutic goal is the advancement of new quinonoid compounds that display anti-neoplastic activity as well as reduced toxicity. However, anthracycline usage is often limited by cardiotoxicity,[4] which also applies to a number of other classes of fused-ring quinones. In particular, chemotherapy using anthracyclines causes cardiomyocyte injury, left ventricular dysfunction, and at the upper levels of dosage required for cancer therapy carries a high risk of heart failure and death.[11] Since such cardiotoxicity might arise through undesired intercalation into normal DNA, or through non-selective damage via quinonoid oxy radicals, we considered whether monocyclic quinones (whose lack of ring fusion would be expected to show little DNA intercalation) might confer reduced toxicity compared to their fused-ring congeners, while retaining anti-cancer potency. Indeed, a number of monocyclic 1,4-benzoquinone derivatives (Fig. 1) show promise including ardisianone (inducing anti-proliferation and apoptosis of several cancer cell lines, including PC-3, DU-145 and human hormone-refractory prostate cancer),[14] isoquin, primin,[15] and HU-395, a dihydro derivative of HU-331.[17] Embelin, another important monocyclic quinone natural product, is an analgesic which also possesses anti-fertility, anti-inflammatory and anti-tumour properties.[16]

The monocyclic quinone HU-331 is of particular interest,[19-21] since it showed no impairment of heart function,[20] in contrast to doxorubicin cardiotoxicity that was severe in a mouse model, even at only one-third of the concentration of HU-331 administered. Quinone cardiotoxicity is generally thought to arise by generation of reactive oxygen species (ROS), however, in that respect, as well as others, HU-331 shows patterns of biological activity that are atypical of many quinones. The lack of cardiac toxicity upon administration of HU-331 is consistent with findings that observed no production of ROS,[22] as well as other studies which showed that ROS were produced, but were not the cause of cell death.[23] Additionally, in various studies, cell death induced by HU-331 could not be attributed to DNA strand breaks, caspase activation, apoptosis, or cell cycle arrest.[24] Although the main mode of action of HU-331 is thought to be catalytic inhibition of DNA topoisomerase II,[20,24] the unusual properties of HU-331 have not been fully accounted for, especially in terms of its molecular structure; despite being of cannabinoid origin, HU-331 does not target cannabinoid receptors.[20] Consequently, of central importance is to gain a detailed understanding of the biological and potentially therapeutic effects as a function of substitution patterns around the quinone ring; as shown in the present study, the extent and nature of substitution can have profound effects.

The close structural relationship (Scheme 3) of HU-331 to cannabidiol (CBD), from which HU-331 can be obtained solely by oxidation,[21] invites the question as to whether cannabinoids can possess anti-cancer activity. Recently, cannabinoids have been shown to inhibit tumour growth and angiogenesis, and to induce apoptosis,[25-27] notably in hormone-sensitive prostate cancers.[7] Cannabidiol can prevent metastasis of breast cancer, possibly by suppressing the activity of the gene Id-1.[28] A recent study concluded that 5-hydroxycannabinol and HU-331 inhibit both the α- and β-forms of topoisomerase II.[29] Δ9-Tetrahydrocannabinol inhibits the growth and metastasis of lung cancer cell lines and prevents the growth of Lewis lung adenocarcinoma by inhibiting DNA synthesis.[25,30] Cannabinoids comprise a diverse class of compounds that interact with a number of...
molecular targets including the endocannabinoid-1 and -2 receptors and the vanilloid receptor,\textsuperscript{[31]} although their anti-cancer mechanisms are not fully understood.

In the search for new anti-cancer agents containing a quinone ring, cannabinoid quinones were considered to be a suitable class since the structural motif affords in vitro and in vivo anti-cancer potency, drug-likeness and, especially, as is the case for HU-331, the potential for little or no cardiac toxicity or myelotoxicity, but with inhibition of angiogenesis.\textsuperscript{[32]} However, HU-331 is too chemically unstable, especially in solution, to be used for cancer therapy. Consequently, aims of the present studies were to identify the anti-cancer pharmacophore latent in HU-331 through the study of structurally related monocyclic quinones, and to synthesise and identify more potent derivatives than HU-331, preferably with improved stability and other drug-like properties.

Results and Discussion

Chemistry

The initial objective was to devise a set of integrated synthetic protocols to cover a wide range of polysubstituted resorcinol derivatives, thereby affording by oxidation novel 2-hydroxy-1,4-benzoquinone derivatives encompassing maximised chemical space. Access to alkyl, cycloalkyl, cycloalkenyl and aryl substituents, among others, was required. The main synthetic strategies to substituted 1,4-benzoquinones are depicted in Scheme 1, in which the last step has in common the oxidation of a substituted resorcinol derivative 8 by Frémy's salt (dipotassium nitrosodisulfonate),\textsuperscript{[33]} in an aqueous buffer of KH$_2$PO$_4$,\textsuperscript{[34]} giving the desired corresponding quinone 9, a means of accessing the quinones which was found in the present study to be both convenient and widely applicable.

Scheme 1. Synthetic routes to monocyclic 2-hydroxy-1,4-benzoquinones. Reagents and conditions: (a) BBr$_3$ (3 equiv.), CH$_2$Cl$_2$, -78 °C, 10 min then 20 °C, 1 h; (b) n-BuLi (1.1 equiv.), THF, 0 °C then cyclic ketone, 0 °C, 30 min then 20 °C, 18 h; (c) TFA, CH$_2$Cl$_2$; (d) TFA, CH$_2$Cl$_2$, then Et$_3$SiH; (e) Frémy's salt (3.5-10 equiv.), KH$_2$PO$_4$aq, acetone; (f) cycloalkenol and CSA (0.1 equiv.), CH$_2$Cl$_2$ or BF$_3$,Et$_2$O, CH$_2$Cl$_2$; (g) n-BuLi, THF then alkyl iodide.

Where feasible, Friedel-Crafts alkylation at the 2-position of resorcinols 2 was the most succinct and hence the preferred route, and was successful with various allylic cycloalkenols. To explore the viability of the Friedel-Crafts route as a general strategy, olivetol was reacted with 1-methylocyclohex-2-en-1-ol,\textsuperscript{[35]} in the presence of BF$_3$,Et$_2$O as previously described;\textsuperscript{[36]} in our hands, only a very low yield of 8p was obtained (Table 1, entry 1). Although a freshly supplied sample of alumina (entry 2) afforded an improved result, 8p was accompanied by its cyclised derivative (Table 1), as an inseparable mixture. Increasing the equivalents of allylic alcohol (1:1 reactants, entry 3) and lowering the amount of Lewis acid gave a high yield of the undesired cyclised compound A which was formed in lower yield when the reaction was carried out at 20 °C (entry 4) but still as a 1:1 mixture of products. Evidently, the basic alumina found in other cannabinoid studies to suppress acidity,\textsuperscript{[36]} and hence minimise subsequent cyclisation, did not prove effective in this pilot reaction.

The overall inconsistency of these results using BF$_3$,Et$_2$O to perform related condensations to give cannabinoids has also been reported by Kassiou’s group,\textsuperscript{[37]} although in their work generally satisfactory results were obtained using only a 0.1% solution of that reagent. In the event, a Bronsted acid approach using 10-camphorsulfonic acid, (CSA),\textsuperscript{[38]} proved much more successful in the present work. In our parallel study of the condensation of the isopropyl cyclohexenol 14 (1.2 equiv.) with olivetol in the presence of CSA (0.1 equiv.) reactant stoichiometry greatly influenced the outcome, an appreciable quantity of olivetol remaining from an initial 1:1 stoichiometry, whereas 8a was obtained in 35% yield using 1.2 equivalents of 14, and in 59% yield using 1.2 equivalents of 14, no olivetol being detected in the latter case. When a 1:1.5 stoichiometry of 1-methylocyclohex-2-en-1-ol: olivetol was used in the presence of CSA (0.1 equiv.) at 20 °C no olivetol remained, and the required cannabinoid adduct 8p was isolated in 45% yield (Table 1, entry 5). This condensation protocol (General Procedure D) was found to be effective in fourteen cases of this study, isolation being usually straightforward, and cyclised by-products being absent for reaction periods of 3-6 h. The resulting resorcinol...
derivatives were oxidised with Frémy’s salt to give a focused set of closely related congeners of HU-331 (Table 2).

In cases where an aromatic alkylation route (i.e. 2 to 8) using a Brønsted acid or a Lewis acid was not efficient, cycloalkenyl derivatives 5 which were then reduced with CF₃CO₂H-Et₃SiH. More usually, and preferably, the sequential dehydration-reduction of alcohols 4 was accomplished in one overall step by addition of TFA followed by Et₃SiH. The resulting methyl ethers 7 were demethylated using BBrs in dichloromethane to give the corresponding resorcinol derivatives 8 which were oxidised to the target quinones 9. The 2,5-dimethylresorcinol derivative 3 was prepared by metalation of 1,3-dimethoxy-5-methylbenzene with n-BuLi and alkylation with MeI to give 1,3-dimethoxy-2,5-dimethylbenzene in 78% yield,[39,40] followed by demethylation of the latter in 88% yield using BBrs.

In order to obtain a full range of diversely substituted resorcinols, several synthetic methods were required (Scheme 2). Suzuki reaction of the boronic acid 10 with a cycloalkenyl trifluoromethanesulfonate (‘triflate’) afforded the 5-substituted cycloalkenyl derivatives 11 and 12 which were reduced to the corresponding 5-substituted cycloalkyl derivatives 1d and 1f using ammonium formate and 5% Pd-C in ethanol, and TFA-Et₃SiH in CH₂Cl₂, respectively. Demethylation of those and other substituted resorcinols was found to be generally satisfactory using BBrs, using a procedure of addition at -78 °C (stirred for 10 min) followed by warming to 20 °C over 1 h.[41] The 5-(2-indenyl) resorcinol ether 16, prepared in 59% yield by Suzuki coupling of boronic acid 10 with 2-bromo-1H-indene, was reduced to the indanyl derivative 1e using ammonium formate and 5% Pd-C in ethanol.

5-Substituted resorcinols usually underwent Friedel-Crafts alkylation mainly or exclusively at the 2-position, as might be expected on grounds of a combination of electronic effects of the two ortho-phenolic OH groups, together with steric hindrance of approach to the 4- and 6-positions provided by the 5-substituent. However, the only isolated product of the condensation of resorcinol with 14 was the 4-substituted derivative 8j (18%). Also, in the condensation of 13 with 14, an appreciable quantity of the 4-substituted product 8l was isolated. Those results are consistent with observations on the condensation of methylcyclohex-2-en-1-ol with orcinol (5-methylresorcinol) in CH₂Cl₂ in the presence of BF₃·Et₂O, in which the main product was that of alkylation at the 4-position.[42]

Conversely, from olivetol (larger C-5 substituent) the main product was the result of solely 2-alkylation (e.g. Scheme 4).

With the principal reactions for the synthesis of HU-331 analogs now in hand, consideration was given to a total

Table 1. Condensation of 1-methylcyclohex-2-en-1-ol with olivetol (1 equiv.) in CH₂Cl₂.[a]

<table>
<thead>
<tr>
<th>Entry</th>
<th>Alkyl alcohol (eq.)</th>
<th>Reagent</th>
<th>Additive</th>
<th>T (°C)</th>
<th>Time (s)</th>
<th>8p (%)</th>
<th>A (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.8</td>
<td>BF₃·Et₂O (2.5 eq.)</td>
<td>Al₂O₃ (20 eq.)</td>
<td>reflux</td>
<td>10</td>
<td>&lt; 5:0</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>0.8</td>
<td>BF₃·Et₂O (2.5 eq.)</td>
<td>Al₂O₃ (20 eq.)[b]</td>
<td>reflux</td>
<td>10</td>
<td>25:25</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>1.0</td>
<td>BF₃·Et₂O (1.6 eq.)</td>
<td>Al₂O₃ (20 eq.)[b]</td>
<td>reflux</td>
<td>10</td>
<td>0:89</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>1.0</td>
<td>BF₃·Et₂O (1.6 eq.)</td>
<td>Al₂O₃ (20 eq.)</td>
<td>20</td>
<td>5 min</td>
<td>31:31</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>1.5</td>
<td>CSA (0.1 eq.)</td>
<td>-</td>
<td>20</td>
<td>6 h[d]</td>
<td>45:0</td>
<td></td>
</tr>
</tbody>
</table>

[a] Percentage of conversion was determined from the ¹H NMR spectra of the crude products. [b] For entries 1-3, a suspension of alumina in solution of BF₃·Et₂O in dry CH₂Cl₂ was stirred for 15 min, then heated at reflux for 1 min (entries 1 and 2) or 5 min (entry 3), and then the reactants added. After the reaction time stated the mixture was quenched with saturated aqueous NaHCO₃. [c] A new bottle of alumina was used. [d] 1-Methylcyclohex-2-en-1-ol was added dropwise over 3 h via a syringe pump; then the mixture was stirred for an additional 3 h.
synthesis of the cannabinoid HU-331. The natural product HU-331 has usually been prepared by aerial oxidation of cannabidiol (CBD), obtained from plant sources.\(^\text{[21]}\) A pure source of HU-331 was required as a benchmark against which to measure biological activity of the novel quinones here reported. Accordingly, a synthetic route was sought, preferably using the optimised Friedel-Crafts alkylation route described above. In our hands, a direct approach to (±)-CBD by the condensation of olivetol with isopiperitenol in the presence of CSA afforded an inseparable mixture of (±)-cannabidiol (CBD) and (±)-\(\Delta^2\)-tetrahydrocannabinol (THC), the latter formed by acid-catalysed cyclisation of the former. Instead, olivetol was condensed with the diol 19 using the Friedel-Crafts conditions optimised as above; pleasingly, the desired triol 20 was obtained in 73% yield (Scheme 3). Although this triol has been reported to cyclise to THC and its derivatives,\(^\text{[43]}\) to the best of our knowledge it has not been used to synthesise CBD. Initially, protection of the phenolic OH groups by tosylation and also by triflation, with concomitant elimination of the carbino was attempted, but in neither case could efficient O-deprotection be achieved. However, mesylation of 20 in the presence of Et\(\text{N}\) also induced the desired elimination to the isopropenyl group, and subsequent removal of the aromatic mesylate groups using MeLi afforded (±)-CBD in satisfactory yield.\(^\text{[44]}\) Notably, oxidation of (±)-CBD using Frémy’s salt afforded (±)-HU-331 in 92% yield. This use of Frémy’s salt compares very favourably with the yield of 20% achieved in the early oxygenation of cannabidiol in aqueous 5% KOH to give HU-331.\(^\text{[21]}\)

\[
\text{Scheme 2. Regioselective synthesis of substituted resorcinol derivatives: Reagents and conditions: (a) For 1d: cycloalkeny] triflate (0.9 equiv.), Ph_3P (1 mol%), Pd(OAc)_2 (1 mol%), K_2CO_3 (3 equiv.), 1:1 dimethoxyethane: water; (b) HCO_2NH_2 (10 equiv.), 5% Pd-C, EIOH; (c) TFA (1.0 equiv.), CH_2Cl_2, then Et_3SiH; (d) BBr_3, 3 equiv., CH_2Cl_2, -78 °C, 10 min then 20 °C, 1 h; (e) 14 (1.5 equiv.), CSA (0.1 equiv.), CH_2Cl_2; (f) trimethylsilylacetylene, (Ph_3P)_2PdCl, then TBAF, THF, 26% (2 steps); (g) H_2, Pd-C, MeOH, 59%; (h) PhBH(OH)_2, (Ph_3P)_2Pd, Na_2CO_3, aq., 1,2-dimethoxyethane, 69%.
\]

\[
\text{10} \quad \text{11} \quad \text{12} \quad \text{13} \quad \text{14} \quad \text{15} \quad \text{16} \quad \text{17} \quad \text{18} \quad \text{19} \quad \text{20} \quad \text{21}
\]
Scheme 3. A convergent synthesis of (±)-cannabidiol and (±)-HU-331 via the triol 20. Reagents and conditions: (a) 19 (1.5 equiv.), CSA (0.1 equiv.), CH₂Cl₂, 20 °C, 3 h; (b) MsCl (6.0 equiv.), Et₃N (10 equiv.), CH₂Cl₂; 0 °C, 1 h, then 20 °C, 16 h; (c) MeLi (23 equiv.), THF, 0 °C, 1 h; (d) Frémy’s salt (5.5 equiv.), KH₂PO₄ (2.3 equiv.), aq. acetone, 20 °C, 16 h.

Table 2. Inhibition of Topoisomerase II by Substituted Monocyclic Quinones.^[a,b]^
<table>
<thead>
<tr>
<th>Entry</th>
<th>Compound</th>
<th>Structure</th>
<th>Topo II Inhibition (%)</th>
<th>Entry</th>
<th>Compound</th>
<th>Structure</th>
<th>Topo II Inhibition (%)</th>
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</thead>
<tbody>
<tr>
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<td><img src="image1" alt="Etoposide Structure" /></td>
<td>45 ± 2</td>
<td>13</td>
<td>9j</td>
<td><img src="image2" alt="9j Structure" /></td>
<td>37 ± 2</td>
</tr>
<tr>
<td>2</td>
<td>(±)-HU-331 (commercial)</td>
<td><img src="image3" alt="HU-331 Structure" /></td>
<td>86 ± 3</td>
<td>14</td>
<td>9k</td>
<td><img src="image4" alt="9k Structure" /></td>
<td>44 ± 2</td>
</tr>
<tr>
<td>3</td>
<td>(±)-HU-331 (synthetic)</td>
<td><img src="image5" alt="HU-331 Structure" /></td>
<td>84 ± 3</td>
<td>15</td>
<td>9l</td>
<td><img src="image6" alt="9l Structure" /></td>
<td>62 ± 3</td>
</tr>
<tr>
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<tr>
<td>5</td>
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<td><img src="image9" alt="9b Structure" /></td>
<td>&lt;5</td>
<td>17</td>
<td>9n</td>
<td><img src="image10" alt="9n Structure" /></td>
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<td>12 ± 1</td>
<td>18</td>
<td>9o</td>
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<td>&lt;5</td>
<td>19</td>
<td>9p</td>
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<td>&lt;5</td>
<td>20</td>
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<td>9</td>
<td>9f</td>
<td><img src="image17" alt="9f Structure" /></td>
<td>&lt;5</td>
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<td>9r</td>
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<td>11</td>
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<td>23</td>
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<td>12</td>
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<td><img src="image23" alt="9i Structure" /></td>
<td>&lt;5</td>
<td>24</td>
<td>9u</td>
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<td>97 ± 4</td>
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</table>

[a] Inhibition at 80 µM concentration of test compound. [b] Average of triplicate runs ±SEM.
Regioselectivity of the Friedel-Crafts condensation reactions involving resorcinol and its derivatives was established by NMR spectroscopy. For example, reaction of resorcinol at the 4-position giving 9j was established from the singlet in the $^1$H NMR spectrum for the isolated C-3 hydrogen atom at δ 6.09 and the four-bond coupling constant of 0.7 Hz for the C-6 hydrogen atom at δ 6.57. Consideration of the number of lines in the $^{13}$C NMR spectra of the resorcinol derivatives 8 was also often diagnostic of the position of condensation (i.e. C2- versus C-4).

In the $^1$H NMR spectrum of the quinones 9 the transdialxial coupling constant of approx 11.5-12.0 Hz confirmed the trans configuration of the isopropyl (or isopropenyl) substituent with the 1,4-benzoquinonyl group. Although the relative configuration of the methyl group in 8h and 9h could not be deduced from NMR data owing to overlapping multiplets in the relevant regions, only one diastereoisomer was detected, and by analogy with the hydroboration-oxidation of (+)-2-carene to give (-)-2-isocarolanol (92%),[45] hydroboration on the face opposite to the bulky isopropyl group would be expected, affording the triequatorially substituted stereoisomer 8h, and hence 9h.

**In vitro** inhibition of topoisomerase II by monocyclic quinones

Several studies have confirmed that the monocyclic quinone HU-331 inhibits the relaxation activity of topoisomerase II,[19,20,24,46] and is highly selective for this isozyme. Both the commercial HU-331 and our synthetic samples of HU-331 (concentration range tested from 0.3-100 mM) were found not to increase the amount of nicked DNA in a topoisomerase I DNA cleavage assay, (data not shown) in accord with previous studies.[20] Equally, no evidence was found for the intercalation of HU-331 (concentration range tested from 0.03-100 mM), as assessed by a DNA intercalation assay using ethidium bromide as the control. For the topoisomerase II DNA relaxation assay (Figs. 2-4) a reliable system involved preincubation of the test compounds with human topoisomerase II (Affymetrix) for 10 min, and a relaxation buffer containing ATP (10 mM) and MgCl$_2$ (0.1 M), giving full inhibition between 27-270 mM and some inhibition at 0.27 mM. First, under those conditions, the same concentrations of commercial HU-331 and synthetic (±)-HU-331 were shown to perform in the same way in the topoisomerase IIα assay (Fig. 2). Secondly, a suitable benchmark concentration of 80 mM was established for the evaluation of the series of monocyclic quinones (Table 2). At that concentration, significant amounts of both supercoiled plasmid DNA and relaxed topoisomeric DNA were observed in the histograms and the corresponding pixel intensities of ethidium bromide fluorescence could be accurately measured using ImageJ software (Fig. 3).

**Quinone structure-activity relationship based on topoisomerase IIα inhibition**

In our hands, synthetic racemic HU-331 and commercial HU-331 were also found to be equipotent in the topoisomerase IIα DNA relaxation assay (Table 2). Additionally, enantiomers of HU-331 have been reported to be equipotent against a variety of cancer cell lines.[20] As with a set of bisdioxopiperazine catalytic inhibitors of topoisomerase II (i.e. inhibitors of one or more steps in the enzyme-DNA ligation/deligation/strand cleavage cycle) whose enantiomers showed equipotency in respect of both enzyme inhibition and cellular cytotoxicity,[46] a similar explanation can be advanced for the quinone HU-331, namely that the site of binding to topoisomerase IIα is either sufficiently large or sufficiently flexible to accommodate each enantiomer equally well.
Of the trisubstituted HU-331 analogs 9a and 9c-9h, in which the terpenoid unit had been retained (or hydrogenated in the cases of 9a and 9h) each of this series of compounds showed some inhibition, but none greater than or equal to HU-331 (Table 2). Saturation of the isopropenyl group of HU-331 lowered the inhibition (61% for 9a, c.w. 86% for HU-331). Complete saturation of the terpenoid unit lowered the inhibition further (53% for 9h). However, removal of the isopropenyl group in HU-331 resulted in only a small loss of inhibition (79% for 9p), suggesting that the isopropenyl group is not essential for potent inhibition of topoisomerase IIα. Although the fully reduced terpenoid unit in 9h showed the lowest potency within the series of terpenoid derivatives studied, this compound illustrated that a 2-cyclohexyl ring can be compatible with good inhibition. Following this observation, the subset of saturated cycloalkyl quinones 9m-9o and 9q was synthesised; not only was good inhibition, consistent with a proposed mechanism involving conjugate addition of a topoisomerase II residue to an unsubstituted position on the quinone ring, residues Cys392 and Cys405 have been identified as significant for addition to p-benzoquinone. In contrast, quinones 9k and 9l showed moderate and good potency, respectively, compared with HU-331. If the novel topoisomerase IIα inhibitors 9j, 9k and 9l, which all possess a 5-substituent, act in a way similar to the parent compound HU-331, conjugate addition presumably occurs at C-3. This requirement for an unsubstituted (Michael acceptor) site on the quinone ring was further probed by examining the effect of 9d and 9b at 200 mM on topoisomerase IIα (Fig. 4); at this higher concentration, 9d showed complete inhibition, but without the formation of observable topoisomers. Conversely, the tetrasubstituted quinone 9b showed no inhibition. Those observations are consistent with covalent adduct formation being a principal mode by which monocyclic quinones including HU-331 can act on human topoisomerase IIα, and which recently has been proposed to involve interference of the quinone with the N-terminal domain, thereby preventing the N-terminal clamp from capturing DNA. The fully substituted quinone ring of 9b retards or prevents adduct formation.

Figure 4. Human topoisomerase II-induced relaxation of supercoiled plasmid DNA in the presence of (a) the trisubstituted quinone 9d and (b) the tetrasubstituted quinone 9b.

The above survey of topoisomerase IIα inhibition as a function of variations in substitution suggested further investigation of 3-cycloalkyl derivatives of 2-hydroxy-6-n-pentyl-1,4-benzoquinone, particularly since 9o had been shown to have much greater stability than HU-331. Initially, and to identify a reference for acyclic substituents, quinone 9s was prepared and evaluated; its considerable potency (58%) was diminished upon O-methylation (31%
for 9t), confirming the need for the unsubstituted hydroxy group. Introduction of a cyclohexyl group at C-3 afforded the most potent compound in the quinone series, showing 97% inhibition at 80 µM, and hence that an n-pentyl chain is not a requirement for potent inhibition of topoisomerase IIα. Variation of the ring size of the C-3 cycloalkyl substituent showed that cyclopentyl conferred the greatest enzyme inhibition (95% for 9n), being more potent than HU-331 (86%). The cycloheptyl congener 9q was the least potent (49% inhibition) of the set 5m-9r. The cyclohexenyl derivative 9p (78% inhibition) was not significantly more potent than the 3-cycloalkyl quinone 9o (76%).

The cyclohexyl ring present in the corresponding compound 9o confers markedly greater stability as well as crystallinity, an observation which helped focus the search for potency a set of quinones 9 that contained a simplified and saturated 3-cycloalkyl unit, compared to the terpenoid unit present in HU-331. Since compounds 9n, 9o and 9u were more potent inhibitors of topoisomerase IIα than HU-331, as well as being considerably more stable, they were tested against a number of cancer cell lines to determine whether the potent enzyme assay results would be accompanied by antiproliferation in whole cell assays.

Cytotoxic effects of monocyclic quinones

Inhibitory activity against proliferation of the prostate cancer cell line DU-145 was measured using an XTT assay with continuous exposure for 72 h (Table 3); compared to HU-331, four compounds were considerably more potent inhibitors. Of the compounds assayed that contain a terpenoid-like substituent, the isopropyl derivative 9a and the corresponding system 9h with a saturated terpenoid system were of similar potency, both being more potent than HU-331, implying that a more saturated terpenoid skeleton can be an improvement. Compared to 9a, conformational restriction of the 6-substituent to cyclopentyl, as in 9d, further increased the inhibitory potency two-fold. Of similar or greater potency against DU-145 cell proliferation was the 3-cyclohexyl derivative 9o. The other derivatives showed medium to low inhibitory activity, despite 9p and 9u showing, respectively, similar and greater potency than HU-331 in the topoisomerase IIα assay. This can be accounted for by a lack of sufficient lipophilicity in the 6-substituent (9s-9w), although that is partly compensated for by the 3-cyclohexyl substituent in 9u. That 9w was by far the least active compound evaluated against DU-145 cell line proliferation is consistent with the absence of any lipophilic group at both C-3 and C-6 (hence very low potency), despite the presence at C-6 in 9w of the complete terpenoid group that is found in HU-331. The main factor accounting for the difference in cellular inhibitory potency is probably the difficulty of the much more polar 9w in traversing the DU-145 cell membrane, as expected from the markedly different lipophilicities (HU-331 cLogP = 6.14 versus 9w cLogP = 3.51). However, the lower potency of 2-methoxy compounds compared to their 2-hydroxy analogs, for both enzyme inhibition and DU-145 cell line anti-proliferation cannot be accounted for in terms of lipophilicities (9s cLogP = 0.99 and 9t cLogP = 1.15), and is ascribed to the hydroxy group acting as a hydrogen bond donor.

A selection of the more potent compounds was then evaluated for inhibition of proliferation of Jurkat human acute T cell leukaemia and Raji human lymphoma cell lines (Table 3). Comparison of 9n with 9o shows that 3-cyclohexyl ring confers nearly an eight-fold greater inhibitory activity across the three cell lines studied than a 3-cyclopentyl ring. The 3-cyclohexyl derivative 9u also

<table>
<thead>
<tr>
<th>Cell line: Compound</th>
<th>DU-145</th>
<th>Jurkat</th>
<th>Raji</th>
</tr>
</thead>
<tbody>
<tr>
<td>(+)-HU-331</td>
<td>9.15±0.57</td>
<td>1.30±0.20</td>
<td>0.61±0.05</td>
</tr>
<tr>
<td>9a</td>
<td>5.45±0.26</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9d</td>
<td>2.4±0.60</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9h</td>
<td>5.2±0.20</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9n</td>
<td>13.7±0.60</td>
<td>2.37±0.10</td>
<td>5.27±0.25</td>
</tr>
<tr>
<td>9o</td>
<td>2.05±0.10</td>
<td>0.29±0.02</td>
<td>0.68±0.03</td>
</tr>
<tr>
<td>9p</td>
<td>20.1±4.4</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9s</td>
<td>27.8±4.3</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9t</td>
<td>40.6±2.9</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9u</td>
<td>12.8±0.52</td>
<td>0.42±0.02</td>
<td>1.45±0.06</td>
</tr>
<tr>
<td>9v</td>
<td>19.0±0.23</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>9w</td>
<td>57.0±15.5</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

[a] Data from cell viability (ATP) assay for compounds 9n, 9o and 9u; all other data for DU-145 from XTT assay. Average of triplicate runs ±SEM. [b] ND = not determined.
showed good-to-excellent potency. The most potent compound was identified as 9o, showing significantly improved potency in two of the three cell lines studied, including submicromolar potency in two of the cell lines. Simplification of chemical structure by removal of metabolically active unsaturation led to greater overall potency than HU-331 as well as greatly increased in vitro stability.

Some differences were observed in the relative inhibition of topoisomerase IIα versus inhibition of cell line proliferation. For example, a terpenoid isopropenyl group (HU-331, 86%) conferred greater potency than the corresponding isopropyl group (9a, 61%), whereas in the DU-145 cell line assay the isopropenyl group in HU-331, 86%) conferred 1.7-fold less potency than the isopropyl group in 9a. Also, absence of the isopropenyl group in HU-331 (86% inhibition of topoisomerase IIα) gave only a small loss in enzyme inhibition (79% for 9p), whereas compared to HU-331 inhibition of DU-145 cell proliferation by 9p was more than halved. A third difference was seen for 9u, the most potent inhibitor of all the novel quinones tested (and more potent than HU-331), but which was less potent than HU-331, 9a, 9d, 9h and 9o in the DU-145 cell line assay. Fourthly, the results of the topoisomerase IIα assay showed that decreasing the 3-cycloalkyl ring size from 7 to 6 to 5 increased potency (9n having the second-highest inhibition of 95% at 80 μM of all compounds tested in the enzyme assay). Conversely, in the cell line assays, decreasing the ring size was detrimental to inhibition of proliferation, the 3-cyclopentyl derivative 9n exhibiting over 6-fold less potency than the 3-cyclohexyl derivative 9o. Thus, while the greater requirement for lipophilicity in whole cell assays compared to an in vitro enzyme assay must be considered, and is expected on account of cell membrane permeability, overall the relative potency of whole cell inhibitory activity still shows some significant differences with the relative potency of the topoisomerase IIα assay, and may indicate that inhibition of topoisomerase IIα (e.g., for 9o) is not the only relevant molecular target;[50] further studies are indicated.

Conclusions

New syntheses of cannabidiol and (+)-HU-331 are reported herein. General synthetic approaches to a wide variety of substituted derivatives of 1,4-benzoquinone have been described, the shortest employing a Friedel-Crafts alkylation of a resorcinol derivative followed by oxidation to the quinone using Frémy’s salt. New derivatives of 1,4-benzoquinone have been identified as more potent than HU-331, a potent natural product anti-cancer agent showing no significant cardiotoxicity. Novel potent inhibitors of human topoisomerase IIα and of proliferating cancer cell lines were identified, including 9a, 9d, 9h and especially 9o which showed a greater than four-fold inhibition of the DU-145 prostate cancer cell line than HU-331, as well as submicromolar inhibitory potency against both Jurkat human acute T cell leukaemia and Raji human lymphoma cell lines. The simplified analogs, including 9o, showed much greater in vitro stability compared to HU-331, ascribed to the absence of chemically reactive groups (which in HU-331 are likely sites of metabolism).

Anti-cancer properties here identified were correlated with structure, which indicated clear directions for further optimisation, especially variation in 6-alkyl substituents; 6-aryl and 3-(4-substituted)cyclohexyl derivatives may also be promising areas of chemical space. This preliminary structure-activity profile identified for these quinones should facilitate the discovery of new, drug-like monocyclic quinone anti-cancer agents with low cardiotoxicity (partly attributable to minimal ROS production), unlike many fused quinone systems, especially the clinically important anthracyclines whose use is constrained by severe and dose-limiting cardiotoxicity.

![Figure 5. Correlation of substituted hydroxy-1,4-benzoquinones with anti-cancer properties identified.](image-url)
multiple cell lines, its crystallinity and its greatly increased stability compared to HU-331, compound 9o is suitable as a probe to elucidate uncharted quinone chemical biology, and also as the basis for next-generation antiproliferative hydroxyquinones possessing minimal cardiotoxicity compared to existing clinical quinone anti-cancer agents.

**Experimental section**

**Instruments and Methods**

Reactants and reagents were used as received. Commercial HU-331 was purchased from Abcam. Compound homogeneity was monitored by ascending thin-layer chromatography, performed on Merck 0.2 mm aluminium-backed silica gel 60 F\textsubscript{254} plates and visualised using ultraviolet irradiation (254 nm) or by using aqueous alkaline potassium permanganate. Flash column chromatography was performed using Merck silica gel 60 of particle size 33-70 mm. Evaporation refers to the removal of solvent under reduced pressure. Melting points were determined using an Optimet apparatus and are uncorrected. Infrared (IR) spectra were recorded on a Bruker alpha FT-IR spectrometer using neat samples. \textsuperscript{1}H and \textsuperscript{13}C NMR spectra (respective operating frequencies of 500 MHz and 125 MHz) were obtained using a Bruker AM-500 spectrometer in a variety of solvents and a residual non-deuterated solvent peak as the internal reference: CDCl\textsubscript{3} (\textsuperscript{1}H = 7.26 ppm, \textsuperscript{13}C = 77.0 ppm), DMSO (\textsuperscript{1}H = 2.50 ppm, \textsuperscript{13}C = 39.5 ppm) CD\textsubscript{3}OD (\textsuperscript{1}H = 3.31 ppm, \textsuperscript{13}C = 49.0 ppm). Mass spectra were obtained on a Fisons VG70-SE mass spectrometer or Thermo Finnigan MAT900xp instrument using electrospray ionisation (ESI), electron ionization (EI) or chemical ionization (CI).

**Supplementary data**

Preparation and spectral data of compounds are given in the Supporting Information.

**General procedure A (demethylation of 1,3-dimethoxyarenes):** To a stirring solution of the 1,3-dimethoxyarene (0.27 M, 1.0 equiv., approx. 0.4 mmol scale) in dichloromethane under nitrogen and cooled in a dry ice-acetone bath was added dropwise boron tribromide (3.0 equiv.) at -78 °C over 2 min. Stirring was continued at -78 °C for a further 10 min. The dry ice-acetone bath was then removed and the mixture allowed to warm to 20 °C over 1 h. Saturated aqueous sodium hydrogen carbonate (20 mL) was then added and the mixture was stirred at 20 °C for a further 10 min.\textsuperscript{[41]} The mixture was extracted with diethyl ether (3 x 20 mL) and the combined organic layers were washed with saturated aqueous sodium hydrogen carbonate then with brine. The combined organic layers were dried (MgSO\textsubscript{4}), filtered and the solvent was evaporated. The residue was purified by column chromatography, typically using 10-30% volume of ethyl acetate in hexane as the eluent to give the resorcinol derivative (details provided for each compound in Supporting Information).

**General procedure B (metalation of a 1,3-dimethoxyarene, addition to a ketone and dehydration of the carbinol):** To a stirring solution of the 1,3-dimethoxy-5-substituted benzene (0.2 M, 1.0 equiv., approx. 1.5 mmol scale) in dry tetrahydrofuran at 0 °C under nitrogen was added n-butyllithium (1.6 M, 1.1 equiv.) dropwise over 5 min. Stirring was continued at 0 °C for a further 1 h. A solution of the cyclic ketone (1.8 M, 1.5 mol equiv.) in dry tetrahydrofuran was then added dropwise over 10 min. The mixture was stirred for an additional 30 min at 0 °C, then at 20 °C for 18 h. Saturated aqueous ammonium chloride (20 mL) was then added. The mixture was then extracted with diethyl ether (2 x 20 mL) and the combined organic layers were washed with water, dried (Na\textsubscript{2}SO\textsubscript{4}), filtered and the solvent was evaporated. The residue was purified by column chromatography, typically using 10-25% volume of ethyl acetate in hexane as the eluent to give the tertiary alcohol (details provided for each compound in Supporting Information).

**General procedure C (reductive elimination of carbinols, and reduction of alkenes to the corresponding aryl cycloalkanes):** To a stirring solution of the tertiary alcohol or the alkene (0.36 M, 1.0 equiv.) in dichloromethane at 20 °C was added trifluoroacetic acid (5.5 equiv.), whereupon the colourless solution immediately turned dark red. Stirring was continued at 20 °C for 15 min, then triethylsilane (2.5 mol equiv.) was added dropwise over 5 min, and the mixture stirred a further 1.5 h. Saturated aqueous sodium hydrogen
carbonate (20 mL) was then added. The organic layer was separated, washed with water (10 mL), dried (Na₂SO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography, typically using 0-5% by volume of ethyl acetate in hexane as the eluent to give the aryl cycloalkane (details provided for each compound in Supporting Information).

**General Procedure D (condensation of resorcinol derivatives with cyclic allylic alcohols):** To a stirring solution of the resorcinol derivative (0.20 M, 1.0 equiv., approx. 0.5 mmol scale) and camphorsulfonic acid (0.10 mol equiv.) in dichloromethane was added dropwise a solution of the cyclic allylic alcohol (0.20 M, 1.5 equiv.) in dichloromethane via a syringe pump at 20 °C over 3 h. The solution was stirred for a further 3 h at 20 °C then saturated aqueous sodium hydrogen carbonate (10 mL) was added. The mixture was extracted with dichloromethane (3 x 20 mL) and the combined organic layers were washed with water, then with brine. The combined organic layers were dried (Na₂SO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography (details provided in Supporting Information: usually using 5% by volume of ethyl acetate in hexane for cannabinoids) to give the cycloalkenyl resorcinol derivative.

**Representative general procedure E (Oxidative dearomatisation of resorcinol derivatives using Frémy's salt):** The detailed procedures and quantities given in the Supporting Information for specific reactions should be used, especially the quantity of Frémy’s salt. To a stirring solution of the resorcinol derivative (0.10 mmol) in acetone (5.0 mL) at 20 °C was added dropwise over 2 min a pre-mixed buffered aqueous solution containing potassium dihydrogen orthophosphate (0.24 mmol, 0.060 M, 4.0 mL) and Frémy’s salt (dipotassium nitrosodisulfonate, 0.268 g, 1.0 mmol; CAUTION!). Only small quantities of Frémy’s salt were used, and under the conditions described did not lead to a rapid exotherm. However, the use of Frémy’s salt represents an explosion hazard, and suitable protection during handling and use should be taken. After stirring at 20 °C until the reaction was complete by TLC (typically 3-4 h), the mixture was extracted with diethyl ether (3 x 10 mL). The combined organic layers were dried (MgSO₄), filtered and evaporated. In most cases the hydroxyquinone product was of high purity, as assessed by ¹H and ¹³C NMR spectroscopy and TLC, but could be further purified by column chromatography, typically using 2%-10% by volume of ethyl acetate in hexane.

**Topoisomerase I DNA cleavage assay**

Human recombinant topoisomerase I (Affymetrix) was supplied in a storage buffer (15 mM sodium phosphate, pH 7.1, 700 mM NaCl, 0.1 mM EDTA, 0.5 mM DTT, 50% glycerol) at a concentration of 20 units/µL and was stored at -20 °C. Inhibitor solutions were prepared by serial dilution of a 10 mM stock solution in DMSO with relaxation buffer (20 mM Tris-HCl pH 7.5, 200 mM NaCl, 0.25 mM EDTA, 5% glycerol). To each Eppendorf tube was added supercoiled plasmid DNA (pBR322, 0.3 µg) followed by different concentrations of HU-331 (0.3 µM to 1 mM). Human recombinant topoisomerase I (2U) was added to each mixture, making the final volume up to 30 µL. The mixtures were incubated together at 37 °C for 45 min. The reactions were then terminated by addition of the stopping buffers SDS and proteinase K to a final concentration of 0.25% and 250 µg/mL respectively. The reaction mixtures were incubated at 50 °C for 30 min then analysed by electrophoresis on 1% agarose gel with 0.5 µg/mL ethidium bromide in a 10% Tris-borate EDTA buffer. The different forms of DNA were then separated by electrophoresis at 90-140 V conducted at room temperature for approximately 2 h. Gels were photographed under UV irradiation using a UV transilluminator coupled with a camera.

**DNA Intercalation assay**

A literature procedure, reported by Peixoto and co-workers[51] was used to investigate the intercalation activity of HU-331 compared to ethidium bromide based on the topoisomerase I relaxation assay. Inhibitor concentrations were made by serial dilution of stock solutions (10 mg/mL for ethidium bromide and 10 mM for HU-331) with topoisomerase I relaxation buffer (20 mM Tris-HCl pH 7.5, 200 mM NaCl, 0.25 mM EDTA, 5% glycerol). To each Eppendorf tube was added supercoiled plasmid DNA (pBR322, 0.3 µg) followed by different concentrations of ethidium bromide (0.03–3 µM) and HU-
331 (0.03–100 μM). Human recombinant topoisomerase I (5 U) was added to each mixture, and the final volume made up to 30 μL using dilution buffer. The mixtures were incubated together at 37 °C for 45 min. Reactions were then terminated by addition of the stopping buffers SDS and proteinase K to a final concentration of 0.25% and 250 μg/mL respectively. The reaction mixtures were incubated at 50 °C for 30 min then analysed by electrophoresis on 1% agarose gel in a 10% Tris-borate EDTA buffer. The different forms of DNA were then separated by electrophoresis at 150 V conducted at room temperature for approximately 2 h. The gels were visualised by staining with ethidium bromide (0.5 μg/mL) for 30 min then de-staining in water for 15 min. The gels were photographed under UV irradiation using a UV transilluminator coupled with a camera.

**Topoisomerase II relaxation assay**

The procedure was similar to the topoisomerase I relaxation assay but with slight modifications.[52] Each inhibitor was tested at one concentration (80 μM) by diluting the stock concentrations with topoisomerase II dilution buffer (10 mM sodium phosphate pH 7.1, 50 mM NaCl, 0.2 mM DTT, 0.1 mM EDTA, 10% glycerol and 0.5 mg/mL BSA. To each Eppendorf tube was added human recombinant topoisomerase II (2 U) followed by reaction buffer (100 mM Tris-HCl (pH 7.9), 500 mM KCl, 500 mM NaCl, 50 mM MgCl₂, 1 mM EDTA, 0.15 mg/mL BSA and 10 mM ATP) diluted tenfold. To this mixture was added the diluted inhibitor to give a final concentration of 80 μM. The mixtures were incubated for at room temperature for 10 min then the supercoiled DNA substrate (0.3 μg) was added, making the final volume up to 30 μL. The mixture was then incubated at 37 °C for 30 min prior to reaction termination with the stopping buffers SDS and proteinase K to give a final concentration of 0.25% and 250 μg/mL respectively. The mixture was then analysed on 1% agarose gel in a 10% Tris-borate EDTA buffer. The different forms of DNA were then separated by electrophoresis at 150 V conducted at room temperature for approximately 2 h. Gels were visualised by staining with ethidium bromide (0.5 μg/mL) for 30 min then de-staining in water for 15 min. The gels were photographed under UV irradiation using a UV transilluminator coupled with a camera. For quantitation, the integrated pixel intensities of the ethidium bromide fluorescence were determined using ImageJ software. Background fluorescence (mainly from open circular ‘nicked’ DNA) was subtracted from all fluorescence values to give normalised intensities, the percentage inhibition being given by 100 – 100 x (normalised intensity for relaxed DNA formed).

**Cell proliferation XTT assay**

A procedure for an MTT assay was used with the following adaptations.[53] DU-145 cells were suspended in RPMI 1640 medium, supplemented with fetal calf serum (40 mL) and L-glutamine (5 mL) at 37 °C in a humidified atmosphere of 5% CO₂ and 95% air. Aliquots (200 μL) of suspensions of cancer cells were dispensed into wells of 96-well tissue culture plates at densities of 4000 cells/well. After incubation for 24 h various concentrations of the 2-hydroxy-1,4-benzoquinone derivative were added and cell viability determined after 72 h using 2,3-bis-(2-methoxy-4-nitro-5-sulphophenyl)-2H-tetrazolium-5-carboxanilide (XTT) in the presence of phenazine methosulfate (PMS). Absorbance was measured against a background control as a blank (0.5% DMSO in growth medium) using a scanning multiwell spectrophotometer (Wallac Victor 1420 multilabel counter) at 450 nm. IC₅₀ values for various compounds were then calculated using OriginPro 9 by plotting the Log of the concentrations against the mean percentage inhibition, and using a non-linear curve fit algorithm.

**Cell Viability Assay**

Cell lines were purchased from American Type Culture Collection (Manassas, VA). Cells of the DU-145 prostate cancer cell line were grown in EMEM medium, whereas those from Raji human lymphoma and Jurkat human acute T cell leukaemia cell line were grown in RPMI-1640 medium. All culture media were supplemented with 10% fetal bovine serum, penicillin (100 μg/mL) and streptomycin (100 μg/mL). Cultures were maintained at 37 °C in a humidified atmosphere of 5% CO₂ and 95% air.

Test compounds and the reference compound staurosporine (Sigma-Aldrich) were prepared in DMSO as 10 mM stock solutions which then were diluted with 10-fold and three-fold dilutions in a source plate. A solution of each test compound (0.25 mL) or of staurosporine (0.025 mL) was delivered from the source plate to each
well of the 384-well cell culture plates by Echo 550. Culture medium (25 µL) containing one thousand cells from the DU-145 prostate cancer cell line, or the Raji human lymphoma cell line, or the Jurkat human acute T cell leukaemia cell line was added to the wells of the cell culture plates. The cells were incubated with the compounds at 37 °C for 72 h under an atmosphere of 5% CO₂ and 95% air. To each well was added Cell Titer Glo 2.0 reagent (25 µL) (Promega, Madison, WI), then the contents were mixed on an orbital shaker for 2 min prior to incubation at room temperature for 10 min to stabilise the luminescent signal. Luminescence was recorded by an Envision 2104 Multilabel Reader (PerkinElmer, Santa Clara, CA). The number of viable cells in culture was determined based on quantitation of the ATP present in each culture well. IC₅₀ curves were plotted and IC₅₀ values were calculated using the GraphPad Prism 4 program based on a sigmoidal dose-response equation.

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Conflict of interest statement
The authors declare no conflict of interests.

Keywords: anti-cancer agents . cannabidiol . Frémy’s salt . HU-331 . monocyclic quinones . structure-activity relationships .

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FAX: +44(0)20 7679 7463.

References
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Many potent anti-cancer and antibiotic agents contain a fused 1,4-benzoquinone ring, but also show dose-limiting cardiotoxicity. Noting the lack of cardiotoxicity of the monocyclic quinone HU-331, new monocyclic quinones were synthesised to determine their anti-cancer properties and identify a provisional pharmacophore. We describe a crystalline monocyclic quinone that like HU-331 is also a topoisomerase IIα inhibitor but showed greater potency against DU-145 and Jurkat cell lines, as well as having much greater stability in solution than HU-331.