Combretazet-3 a novel synthetic cis-stable combretastatin A-4-azetidinone hybrid with enhanced stability and therapeutic efficacy in colon cancer

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**Abstract.** In recent years an extensive series of synthetic combretastatin A-4 (CA-4)-azetidinone (β-lactam) hybrids were designed and synthesised with a view to improve the stability, therapeutic efficacy and aqueous solubility of CA-4. Lead compounds containing a 3,4,5-trimethoxy aromatic ring at position 1 and a variety of substitution patterns at positions 3 and 4 of the β-lactam ring were screened in three adenocarcinoma-derived colon cancer cell lines (CT-26, Caco-2 and the CA-4 resistant cell line, HT-29). In both CT-26 and Caco-2 cells all β-lactam analogues analysed displayed potent therapeutic efficacy within the nanomolar range. Substitution of the ethylene bridge of CA-4 with the β-lactam ring together with the aforementioned aryl substitutions improved the therapeutic efficacy of CA-4 up to 300-fold in the combretastatin refractory HT-29 cells. The lead compound combretazet-3 (CAZ-3); chemical name [4-(3-hydroxy-4-methoxyphenyl)-3-(4-hydroxyphenyl)-1-(3,4,5-trimethoxyphenyl)azetidin-2-one] demonstrated improved chemical stability together with enhanced therapeutic efficacy as compared with CA-4 whilst maintaining the natural biological properties of CA-4. Furthermore, CAZ-3 demonstrated significant tumour inhibition in a murine model of colon cancer. Our results suggest that combretastatin-azetidinone hybrids represent an effective novel therapy for the treatment of combretastatin resistant carcinomas.

**Introduction**

Microtubules are a principle component of the cytoskeleton and play a key role in numerous biological functions including cell division and organelle transport. The pivotal role of tubulin in both the formation of the mitotic spindle and chromosomal separation prompted the surge in the development of both natural and synthetic microtubule targeting agents (MTAs). One such naturally occurring drug, combretastatin A-4 (CA-4; Fig. 1) was originally described by Pettit et al (1). The structure of CA-4 proved readily amenable to chemical manipulation to improve the stability, solubility and therapeutic index of this class of MTAs. Over the past two decades a vast array of synthetic CA-4 analogues were designed and synthesised with many surpassing the stability, solubility and therapeutic efficacy of the parent compound (2). The clinical success of the synthetic prodruk of CA-4, combretastatin-A4 phosphate (CA-4P) in the treatment of anaplastic thyroid carcinoma (www.clinicaltrials.gov) (3) has maintained an active interest in the chemical manipulation of CA-4 with the view to further enhancing the therapeutic efficacy of this lead compound. Furthermore, amino acid containing prodrugs of CA-4 are also undergoing clinical evaluation including AVE8062 (www.clinicaltrials.gov) (4). Structural modifications of CA-4 can be divided into three areas, those involving the manipulation of either ring A, ring B or those involving the substitution of the double bond (ethylene bridge structure) connecting both rings (Fig. 1). These A and B substituted aromatic rings fit into the A and B pockets of the colchicine binding site on tubulin. Data collated from numerous structural activity relationship (SAR) studies confirm that a non-planar cis conformation is essential for the tubulin binding properties of CA-4. Furthermore, the majority of studies suggest that the 3,4,5-trimethoxy-substituted aromatic A-ring should be conserved to maintain maximum anticancer activity. However, contrary to this, a recent study conducted by Beale et al (5) showed that substitution of the larger meta-methoxy groups of triazole CA-4 derivatives with smaller halogen atoms yielded more potent CA-4/CA-1 analogues. Several independent studies have demonstrated that the therapeutic activity of CA-4 can also be enhanced by the strategic modification of ring B (6). Apart from strategies to improve the potency of CA-4 other main areas of research focused on methods to overcome the solubility issues of CA-4 and also to prevent the undesired conversion into the inactive trans isomer (Fig. 1). Modification of the phenolic group on
ring B forming either a phosphate or an amino acid ester was demonstrated to be an effective method of improving the solubility of CA-4 whilst retaining optimum biological activity. Double bond isomerisation can be prevented by the strategic inclusion of various types of heterocyclic rings in place of the usual ethylene bridge structure of CA-4 (7).

In recent years our group has designed and synthesised an extensive series of azetidinone (ß-lactam) CA-4 analogues with a view to overcoming double bond isomerisation by substituting the ethylene bridge structure for a 1,4-diaryl-2-azetidinone ring. The rigid ß-lactam ring scaffold allows a similar spatial arrangement between the two aromatic rings as observed in the non-planar cis-conformation of CA-4 while permanently preventing the undesired conversion to the inactive trans-conformation (8). Further studies demonstrated that the inclusion of an aromatic ring at position 3 of the ß-lactam significantly improved the potency of the series. Hence, a ß-lactam substituted at position 3 with a phenolic ring soon became the core structure for future designs (9). Solubility issues of the CA-4-azetidinone analogues were addressed by esterification of the 3-OH group of ring B with phosphates and amino acids (unpublished data).

However, despite the significant advances made in recent years in terms of improving the solubility and stability of CA-4, the lack of therapeutic efficacy as a single agent and the emergence of resistance to CA-4 has somewhat hindered the clinical and commercial success of this compound. We recently reported that CA-432, a lead combretastatin-azetidinone hybrid was 10-fold more potent than CA-4 in CA-4 refractory HT-29 cells, suggesting a possible functional advantage of the ethylene bridge-azetidinone substitution (10). In this study, we screened selected combretastatin-azetidinone hybrids (hereafter referred to as combretazets) to further characterise the structure activity relationship of these compounds in the CA-4 resistant HT-29 cells. The stability and therapeutic potential of a lead compound combretazet-3 (CAZ-3) as a single agent in the murine CT-26 colon cancer model was evaluated.

Materials and methods

Compounds. CA-4 and bafilomycin A1 were purchased from Sigma-Aldrich (Poole, Dorset, UK). 1,4-Diaryl-2-azetidinone analogues were synthesised as previously described by Carr et al (8) CAZ-1, CAZ-2 and CAZ-3 (9), CAZ-4 (7), CAZ-5 (11), CAZ-2P, CAZ-3P, CAZ-4P, CAZ-6 and CAZ-7 (unpublished data). All general reagents unless stated otherwise were purchased from Sigma. Bafilomycin A1 was dissolved in DMSO. CA-4 and all analogues were prepared as a 10-mM stock in ethanol and stored at -20°C.

Cell culture. CT-26 cells are a chemically (N-nitroso-N-methylurethane) induced, undifferentiated murine colon carcinoma fibroblast cell line originating from BALB/c mice. HT-29 and Caco-2 cells originate from a human adenocarcinoma of the colon and were originally obtained from the European Collection of Cell Cultures. All cells were grown in DMEM Glutamax media. CT-26 and HT-29 media were supplemented with 10% foetal bovine serum (FBS) and Caco-2 were cultured with 20% FBS. Both CT-26 and Caco-2 media were supplemented with 1% non-essential amino acids (NEAA). All media contained 100 U/ml penicillin and 100 µg/ml streptomycin. Cells were maintained at 37°C in 5% CO₂ in a humidified incubator. Cell culture materials were supplied from Gibco, Invitrogen Corp. (Grand Island, NY, USA). All cells were sub-cultured 3 times/week by trypsinisation.

Alamar blue cell viability assay. Cell proliferation was analysed using the Alamar Blue assay (Invitrogen Corp.) according to the manufacturer's instructions. Cells were seeded at a density of 5x10⁴ cells/well (CT-26) or 1x10⁵ cells/well (Caco-2, HT-29) in triplicate in 96-well plates. After 24 h, cells were then treated with either medium alone, vehicle [1% ethanol (v/v)] or with serial dilutions of CA-4 or combretazets. After 72 h, Alamar Blue [10% (v/v)] was added to each well and plates were incubated for 3-5 h at 37°C in the dark. Fluorescence was read using a 96-well fluorimeter with excitation at 530 nm and emission at 590 nm. Results were expressed as percentage viability relative to vehicle control (100%). Dose response curves were plotted and IC₅₀ values (concentration of drug resulting in 50% reduction in cell survival) were obtained using the commercial software package Prism (GraphPad Software, Inc., La Jolla, CA, USA). Experiments were performed in triplicate on at least three separate occasions.

Cell cycle detection. After treatment, cells were collected then centrifuged at 800 x g for 10 min and fixed with 70% ethanol overnight at -20°C. The ethanol was removed by centrifugation at 800 x g for 10 min. The cells were then stained in PBS containing 0.5 mg/ml RNase A and 0.15 mg/ml propidium iodide and then incubated for 30 min in the dark at 37°C. Cell cycle distribution was analysed by flow cytometry at 488 nm using the FACSCalibur flow cytometer (Becton-Dickinson, San Jose, CA). All data were recorded and analysed using the CellQuest Software (Becton-Dickinson).

Plasma and pH stability studies. Peripheral blood was collected from healthy donors with informed consent and was made anonymous prior to use. The plasma was separated by Ficoll-gradient and diluted (1:9) with PBS pH 7.4 and warmed to 37°C. The pH stability study was carried out in PBS pH 3.0. Test compounds (1.5 mg/ml) were dissolved in acetonitrile at time t=0. Plasma solution containing test compound (250 µl) was added to 2% (w/v) ZnSO₄ solution in acetonitrile with either medium alone, vehicle [1% ethanol (v/v)] or with serial dilutions of CA-4 or combretazets and incubated for 3 min at 9,500 x g before injection onto an HPLC column (Varian Pursuit XRs C18 reverse phase 250x4.6 mm chromatography column) to determine the stability using a Waters 2487 Dual Wavelength Absorbance Detector, a Waters 1525 Binary HPLC Pump, a Waters In-Line Degasser AF and a Waters 717 plus Autosampler. Samples were detected using a wavelength of 254 nm. All samples were analysed using acetonitrile (60%):water (40%) with 0.1% (v/v) trifluoroacetic acid over 10 min and a flow rate of 1 ml/min to evaluate the percentage decline in the predetermined peak area for each compound. The retention times for CA-4 and CAZ-3 were 5.9 and 3.7 min, respectively. The percentage recovery was calculated using the following formula: [plasma or pH 3.0 peak area/mean aqueous peak area] x 100].
**Human microsomal stability study.** Microsomal stability was determined using pooled human liver microsomes (The UK Human Tissue Bank, Leicester, UK). Ethical approval was obtained from south Cheshire Local Research Ethics Committee (Chester, UK). Test compound (3 µM) together with microsome protein (0.5 mg/ml), 1 mM NADPH in 0.1 M phosphate buffer pH 7.4 was incubated for 0, 5, 15, 30 and 45 min. The negative control did not contain NADPH. The samples were quenched with methanol and the protein was precipitated by centrifugation for 20 min at 1,100 x g at 4°C. Supernatants were then analysed by LC/MS. The In peak area ratio (compound peak area/internal standard peak area) was plotted against time and the slope of the line determined to give the elimination rate constant \( K = (-1)^{\text{slope}} \). The half life \( t_{1/2} \) and the \textit{in vitro} intrinsic clearance (CL\textit{int} µl/min/mg protein) were calculated by the following equations; \( t_{1/2} = 0.693/K; \) \( \text{CL}_{\text{int}} = V \times (0.693)/t_{1/2} \) where \( V \), incubation volume in µl/mg microsomal protein.

**Quantification of AVOs with acridine orange staining using flow cytometry.** Autophagy is characterised by the formation and promotion of acidic vesicular organelles (AVOs). The formation of acidic compartments was quantified by flow cytometric analysis of acridine orange stained cells (10). Acridine orange stains the cytoplasm green and the nucleus a dim red, whereas acidic compartments fluoresce bright red. The intensity of the red fluorescence is proportional to the amount of acidity. Following treatment, cells were stained with acridine orange 1 µg/ml for 15 min at 37°C. Bafilomycin A1 (5 nM) was dissolved in DMSO and added to the cells 45 min prior to the addition of acridine orange. Cells were then trypsinised and collected in phenol-red free medium. Green (510-530 nm) and red (650 nm) fluorescence emission from \( 10^4 \) cells illuminated with blue (488 nm) excitation light was measured with a CyAn ADP Flow Cytometry Analyzer (Beckman Coulter, Nyon, Switzerland). The red:green fluorescence ratio for individual cells was calculated using FlowJo software (Tree Star, Inc., San Carlos, CA).

**In vivo studies.** Tumour growth was initiated by subcutaneous injection of a CT-26 cell suspension (10⁶ cells) into the right flank of 6-8 week old female Balb/c mice. The experiments were conducted on Day 7 when tumours reached a maximum diameter range of 3.6-6.3 mm. Mice were randomly divided into two groups of five. The treatment group received a single intraperitoneal (i.p.) injection of 40 mg/kg CAZ-3 dissolved in ethanol:cremophore:PBS [10%:10%:80% (v/v)] and the control group received one i.p. injection of vehicle only. Tumour growth was measured every second day with a sterile vernier callipers. The long (L) and short (S) axes were recorded and tumour volume (V) was calculated using the following equation \( V = (S \times L)/2 \). Mice were culled by CO₂ asphyxiation at the experimental end point. Ethical approval was obtained from the Research Ethical Approval Committee, Trinity College Dublin. The study was performed under the license number: B100/4275 granted by Department of Health and Children, Hawkins House, Dublin 2, Ireland.

**Results**

The effects of CA-4 and selected combretazets on the viability of colon cancer-derived cell lines. The synthetic combretazets were designed and synthesised with a view to improve the stability, therapeutic efficacy and aqueous solubility of the parent compound CA-4, hence many in the series contained a phenolic group, phosphate ester or an amino group (Fig. 1).
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All combretazets analysed were effective in the nanomolar range in drug-sensitive CT-26 and Caco-2 cells and were more potent than CA-4 in CA-4 refractory HT-29 cells (Table I). Compound CAZ-2 is identical to CA-4 with the exception of the azetidinone-ethylene bridge substitution and demonstrated an 8-fold increase in activity in HT-29 cells confirming a functional advantage of the ethylene bridge-azetidinone substitution in overcoming combretastatin resistance. Compounds containing a B-ring meta-hydroxy group (CAZ-1, CAZ-2 and CAZ-4) or a phosphate (CAZ-2P, CAZ-3P and CAZ-4P) conjugate were the least active of the series in the combretastatin refractory HT-29 cells. Deletion (CAZ-5) or substitution of the B-ring meta-hydroxy group with an amine conjugated amino acid (CAZ-6 and CAZ-7) significantly increased activity of the series with IC₅₀ values in the nanomolar range in combretastatin refractory HT-29 cells. However, CAZ-3 with a B-ring meta-hydroxy group was the exception to this observation. CAZ-3 was more potent than CA-4 in all three adenocarcinoma-derived cell lines tested. Hence, CAZ-3 was selected for further biological analysis.

The effects of CA-4 and selected combretazets on the cell cycle in CT-26 cells. CT-26 cells were selected for further analysis given that all combretazets were effective in the nanomolar range in this cell line. The effects of CA-4, CAZ-3 and its corresponding prodrug CAZ-3P on the cell cycle and cell death were assessed by flow cytometric analysis of propidium iodide stained CT-26 cells. The percentage of cell death was estimated by the quantification of the pre-G₁ peak. As shown in Fig. 2, all compounds tested produced an early G₂M cell cycle arrest at 8 h followed by a significant time-dependent increase in cell death.

Induction of autophagy by CAZ-3. Our recent findings demonstrated that CA-4 and CAZ-2 (CA-432) induced autophagy in adenocarcinoma-derived colon cells as confirmed by acridine orange staining of vesicle formation, electron microscopy and increased expression of LC3-II (10). Hence, the effect of

Table I. Evaluation of CA-4 and selected combretazets in adenocarcinoma-derived colon cancer cells.

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC₅₀ (nM)</th>
<th>RI</th>
<th>Compound</th>
<th>IC₅₀ (nM)</th>
<th>RI</th>
<th>Compound</th>
<th>IC₅₀ (nM)</th>
<th>RI</th>
</tr>
</thead>
<tbody>
<tr>
<td>CA-4</td>
<td>5.7</td>
<td>1.0</td>
<td>CAZ-1</td>
<td>11.8</td>
<td>-2.1</td>
<td>CAZ-2</td>
<td>13.46</td>
<td>-2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CAZ-1</td>
<td>11.8</td>
<td>-2.1</td>
<td>CAZ-2</td>
<td>13.46</td>
<td>-2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CAZ-2</td>
<td>13.46</td>
<td>-2.4</td>
<td>CAZ-3</td>
<td>13.46</td>
<td>-2.4</td>
</tr>
</tbody>
</table>

Cells were exposed to multiple concentrations of the indicated compound for 72 h. Cell viability was assessed using the Alamar Blue assay and respective IC₅₀ values were calculated. ND, not determined. RI, relative inhibition, compared to CA-4. The most active analogue, CAZ-3, is highlighted in bold.

Table II. Calculation of intrinsic clearance values (CLᵢₐᵢ) for CA-4 and CAZ-3 in human microsomes.

<table>
<thead>
<tr>
<th>Compound</th>
<th>CLᵢₐᵢ (µl/min/mg protein)</th>
<th>t₁/₂ (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CA-4</td>
<td>157.0±18.9</td>
<td>8.83</td>
</tr>
<tr>
<td>CAZ-3</td>
<td>76.7±21.4</td>
<td>18.1</td>
</tr>
</tbody>
</table>

Compounds (3 µM) were incubated in the presence of NADPH (1 mM) and human liver microsomes (0.5 mg/ml) and intrinsic clearance values were calculated as described in Materials and methods. Clearance values are expressed as (µl/min/mg) of microsomal protein. Values represent the mean ± SEM of at least triplicate determinations.

Table III. Suppression of tumour growth by CAZ-3.

<table>
<thead>
<tr>
<th>Group</th>
<th>Tumour volume (Day 7)</th>
<th>Tumour volume (Day 11)</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>37.08±8.326</td>
<td>263.2±62.13</td>
<td>0/5</td>
</tr>
<tr>
<td>CAZ-3</td>
<td>42.45±11.45</td>
<td>77.62±19.24</td>
<td>1/5</td>
</tr>
</tbody>
</table>

Single IP NS <0.05

40 mg/kg

BALB/c mice (n=10) were inoculated subcutaneously with 10⁶ CT-26 cells. On Day 7 mice received a single i.p. injection of vehicle (ethanol:cremophore:PBS; 10%:10%:80%) or 40 mg/kg CAZ-3. Tumour growth was measured every second day using a sterile vernier callipers. Values represent the mean ± SEM (Student's t-test; *P<0.05; NS, not significant).

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Induction of autophagy by CAZ-3. Our recent findings demonstrated that CA-4 and CAZ-2 (CA-432) induced autophagy in adenocarcinoma-derived colon cells as confirmed by acridine orange staining of vesicle formation, electron microscopy and increased expression of LC3-II (10). Hence, the effect of
CAZ-3 on autophagic vesicle formation was evaluated by flow cytometric analysis of acridine orange stained cells. Fig. 3 indicates that like other combretastatins CAZ-3 also induced autophagy in adenocarcinoma cells. Furthermore, CAZ-3 also induced autophagy in HT-29 and Caco-2 adenocarcinoma-derived colon cancer cells (data not shown). Numerous studies have demonstrated a dependence of the acidification of cellular organelles on the vacuolar H$^{+}$ ATPase using the specific inhibitor bafilomycin A1. Similarly, pretreatment of CT-26 cells with bafilomycin A1 significantly inhibited CAZ-3 induced autophagy (Fig. 3).

CAZ-3 is more stable than CA-4 in both human plasma and microsomes. The stability of CA-4 and CAZ-3 in acidic media and in human plasma was next determined by HPLC. Both compounds were stable in plasma at physiological pH 7.3 for up to 5 h (Fig. 4A) and remained stable up to 24 h (data not shown). Under acidic conditions (pH 3.0) CAZ-3 is more stable than CA-4 (Fig. 4B). These findings are in agreement with other studies demonstrating instability of CA-4 in acidic media (12). Microsome stability was determined using pooled human liver microsomes (Table II). The β-lactam bridge improved the metabolic stability of CAZ-3 as compared with CA-4 by doubling the in vitro clearance time.

CAZ-3 significantly inhibited the growth of CT-26 cells grafted to mice. To study the effects of CAZ-3 on tumour growth we selected the CT-26 murine model of colon carcinoma, a model frequently used to test the efficacy of CA-4 and its synthetic analogues (13). Furthermore, experimental models involving xenografts of human tumours in a mouse host may lack some of the critical tumour host interactions. In the antitumour efficacy experiment mice received a single IP injection of 40 mg/kg on Day 7 when tumours were on average 5 mm in diameter. On Day 7 there was no significant difference between control and CAZ-3 treated groups. By Days 9 (data not shown) and
11 (Table III), CAZ-3 significantly inhibited tumour growth. Both a rough coat and diarrhoea were observed in 100% of CAZ-3 treated mice.

Discussion

Several water soluble CA-4 analogues including CA-4P (ZYBRESTAT), CA-1P (OXi4503) and AC7700 (AVE 8062) are currently undergoing clinical trials as vascular targeting agents (www.clinicaltrials.gov) (14,15). However, these compounds all contain the isomerisable olefinic bond which may hinder the continued clinical success of the compounds. To date there is no cis-stable CA-4 analogue undergoing clinical trials and hence there is a demand for pre-clinical data on potent cis-restricted CA-4 analogues. The combretazets are a novel class of synthetic combretastatin and azetidinone (β-lactam) hybrids that function through a combretastatin-like mechanism. Overall the combretastatins and the combretazets are structurally and functionally similar. Both classes exhibit a similar spatial arrangement between the two phenyl A and B rings but differ in the bridge structure connecting the rings. The strategic ethylene bridge-azetidinone substitution produced cis-stable analogues with improved chemical stability and ease of synthesis. Extensive biochemical analysis demonstrated that the ethylene bridge-azetidinone substitution did not influence the biological properties of CA-4 (9,10,16,17). In more detail, both classes of drugs inhibit the polymerisation of tubulin inducing a range of cellular responses including; G2/M cell cycle arrest, autophagy, mitotic catastrophe, caspase-dependent cell death and caspase-independent cell death. In this report, we
demonstrate that further substitutions to the aromatic ring at position 3 of the azetidinone yielded a superior compound with enhanced stability and potency against combretastatin refactory adenocarcinoma-derived cells and tumours without altering the biological properties of CA-4. The lead compound CAZ-3 inhibited the polymerisation of tubulin (9), induced a G2M cell cycle arrest and a time-dependent increase in cell death (sub-G1) in the colon adenocarcinoma-derived CT-26 cells in a similar manner to its phosphate prodrug counterpart (CAZ-3P) and CA-4. As recently observed with CA-4 (10), CAZ-3 also induced autophagy in CT-26 adenocarcinoma cells. Autophagy is a highly regulated self-catabolic process which can facilitate a prolonged cell survival in spite of adverse stress by generating energy via lysosomal degradation of cytoplasmic constituents (18). Furthermore, we have previously demonstrated that manipulation of autophagy can enhance the therapeutic potential of CAZ-2 (10).

The adenocarcinoma-derived HT-29 cells are inherently resistant to CA-4. The mechanism of innate resistance remains undefined. Recent studies rule out multidrug resistance protein-1 (MRP-1) mediated resistance to CA-4 in HT-29 cells (12). MRP-1 is a member of the ATP-binding cassette family of polytopic membrane transporters and is responsible for conferring resistance to a broad range of chemotherapeutic drugs (19). However, the authors demonstrate a role for MRP-1 in mediating resistance to some oxazole CA-4 derivatives (12). Data obtained from SARs from numerous independent studies on CA-4 analogues provide some insight into the possibility of structural modification of CA-4 as a means of overcoming CA-4 resistance. In this report we demonstrate that a substitution of the ethylene bridge with a β-lactam ring (CAZ-1) increased activity in HT-29 cells compared to CA-4 by 8-fold. This finding would suggest that cis-trans isomerisation is not solely responsible for CA-4 resistance in HT-29 cells but may contribute in part to CA-4 resistance in these cells. This mechanism of resistance may be overcome by synthetic analogues featuring an ethylene bridge substitution with various types of heterocyclic rings yielding stable analogues which do not isomerise. We also report that deletion or substitution of the B-ring meta-hydroxy group with an amine conjugated amino acid in conjunction with introduction of a 3-position aromatic ring significantly enhances the activity of the series by up to 300-fold in HT-29 cells as compared to CA-4. These findings are in agreement with a recent report by Schobert et al (12) demonstrating improved activity of oxazole bridged CA-4 analogues by substitution of the B-ring phenolic group with H, fluoro or amino groups. Also, substitution of the ethylene bridge with a sulfone group coupled with the substitution of B-ring with a 5-amino-6-methoxyquinoline moiety yielded a novel compound with activity in the low nanomolar range in HT-29 cells (IC50=16 nM) (20). Ring B 4-ethoxyphenyl 1,5-diaryl substituted 1,2,3,4-tetrazoles also displayed potent activity in HT-29 cells (21). Taken together these findings highlight the potential of B-ring meta-hydroxy group substitutions or deletions in cis-stable analogues of CA-4 as a means of overcoming innate resistance to CA-4. However, our lead compound CAZ-3, a cis-restricted CA-4 analogue with a B-ring meta-hydroxy group demonstrated potent nanomolar activity in HT-29 cells. Molecular modelling studies highlighted a novel site of interaction between the para-phenolic 3-position of compound CAZ-3 with the colchicine binding site of tubulin (9). This unique tubulin binding characteristic is shared with compound CAZ-5 via the B-ring ethoxy group but not with CA-4 and the other listed β-lactams (11). The additional hydrophobic contact of CAZ-3 and CAZ-5 with tubulin may facilitate the observed potent antiproliferative effects observed in the CA-4 resistant HT-29 cells and offer a novel means of overcoming CA-4 resistance.

Based on promising in vitro data we proceeded to evaluate the therapeutic efficacy of CAZ-3 in the mouse CT-26 model of colon adenocarcinoma. As a single agent CA-4 failed to reduce the growth of CT-26 tumours in vivo (13). Here we report that a single injection of CAZ-3 (40 mg/kg) significantly inhibited the growth of CT-26 tumours in vivo. Furthermore, CAZ-3 reduced the tumour levels to 34% of control untreated tumour size. This value is below the TC50 value of 42% which is defined as the minimum level of activity required by the National Cancer Institute criteria. However, despite an excellent tumour response rate, a single i.p. injection of CAZ-3 at 40 mg/kg gave adverse side effects such as rough coat, loss of appetite and diarrhoea along with a mortality rate of 1/5. The single maximum tolerated dose for CA-4P was 360 mg/m2 in rats and 100 mg/m2 in dogs (22). Serious diarrhoea has been reported elsewhere in animals given an i.p. injection of CA-4 at 100 mg/kg (23). CA-4P (100 mg/kg) had a mortality rate of 25% in rats (http://arno.unimaas.nl/show.cgi?f=7247). In 90% of patients CA-4P is well tolerated at 52 mg/m2 (22). Given that CAZ-3 is intrinsically more stable than CA-4 and has a slower intrinsic clearance time, CAZ-3 may be more active in vivo as well as in vitro and thus may require significantly lower dosing rates. Optimising the dosing schedule and/or administration route may yield more favourable results.

In conclusion, we have presented preclinical data on a novel series of cis-stable combretastatin analogues. We demonstrate that our lead compound CAZ-3 is effective against CA-4 resistant colon cancer-derived cells in vitro and in vivo. Further studies are warranted to characterise the metabolites of CAZ-3 and optimise dosing schedules with a view to improving the therapeutic potential of this novel class of cis-stable vascular targeting agents.

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References


