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A nitroalkane-based approach to one-pot threecomponent synthesis of isocryptolepine and its analogs with potent anti-cancer activities†

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A second generation polyphosphoric acid-mediated one-pot three-component synthesis of indoloquinoline scaffold is developed. This improved version of the process involves electrophilically activated nitroalkanes for the installation of strategic C-C and C-N bonds and ring C assembly. This modification allows the elimination of unnecessary solvent change operations and all steps are carried out in a true, uninterrupted one-pot manner. A further improvement involves the possibility to install an *ortho*-amino group *in situ*. A synthetic application of this method is showcased by the concise synthesis of an isocryptolepine alkaloid and its synthetic analogs with potent anticancer activities.

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Introduction

Isocryptolepine (cryptosanguinolentine, 1), one of the four types of indologuinoline alkaloids isolated from West African plant Cryptolepis sanguinolenta, has been investigated as a promising antimalarial and anticancer agent (Fig. 1),2 and has been the target of numerous synthetic efforts.3 All these synthetic routes required multistep linear sequences with the isolation and purification of intermediates. Recently, we disclosed a highly efficient metal-free three-component synthesis of isocryptolepine and its bioactive analogs involving a sequential assembly of the indoloquinoline core in one-pot with a single isolation operation.4 This synthetic approach was inspired by the 2009 Kundu total synthesis, featuring Fischer indolization followed by an oxidative Pictet-Spengler type reaction to assemble ring C.5 Our featured protocol (Generation 1) involved the formation of hydrazones 4 from hydrazines 2 and orthoaminoacetophenones 3 (step A), followed by polyphosphoric acid (PPA)-assisted Fischer indolization (step B) to furnish indole 5 (Scheme 1).4 Subsequent acid-mediated acylation of ortho-aniline substituent at C-2 (step C) with an appropriate

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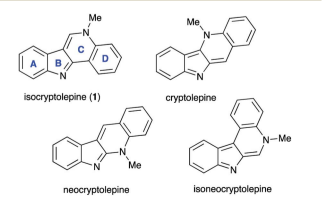


Fig. 1 Natural indoloquinoline alkaloids isolated from *Cryptolepis* sanguinolenta.

carbonyl compound equivalent (carboxylic acid, nitrile or triazine) afforded species 6, that underwent intramolecular Vilsmeier reaction (step D) to provide the indologuinoline core of the alkaloid (Scheme 1).4 Remarkably, this transformation is carried out in a one-pot fashion requiring a single isolation operation at the end of the sequence, leading to an efficient synthesis of an anticancer drug candidate library for SAR studies. However, this synthetic protocol depended on the availability of ortho-aminoacetophenones 3 and carbonyl compound equivalents (acylating agents used in step C: carboxylic acids, nitriles or 1,3,5-triazines). We also discovered an alternative protocol involving unusual cyclization with nitroalkenes accompanied by extrusion of HCN (Scheme 2).6 Still, these previously developed methods did not provide any synthetically meaningful results in reactions of acylating agents bearing functional group as substituent R⁵. In certain cases,

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Paper

$$R^{1}$$
 R^{2}
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these issues rendered synthesis of some types of compounds for our SAR studies impossible or cost prohibitive. Herein, we disclose a further modified synthetic protocol (Generation 2) addressing these problems, and results of our second series of screening for antiproliferative activities that allowed to identify highly potent lead structures.

Results and discussion

It should be pointed out that cyclization of intermediate 8 shown in Scheme 2 involves an unusual electrophilic component formed via phosphorylation of a nitroalkane in nitronate form. Mechanistically, this process is related to the classical Nef reaction,7 employing nucleophilic amine species instead of water. We have discovered a number of innovative one-pot transformations involving nitrogen8 or carbon9 nucleophilebased versions of the Nef reaction and wondered if nitroalkane species 9, electrophilically activated by PPA, could serve as an alternative acylating agent to trigger cyclization of the in

situ generated intermediate 5 into indoloquinoline species 1. Indeed, a nucleophilic attack of indole 5 across the C=N bond of the phosphorylated nitronate 10 would afford (azanediylbis(oxy))bis(phosphonic) species 11, which after 1,2-elimination of ortho-phosphoric acid would produce O-phosphorylated oxime 12 (Scheme 3). Intramolecular nucleophilic attack of aniline moiety would produce cyclic aminal 13, which after 1,2elimination of hydroxylamine O-phosphonate would aromatize into product 1 (Scheme 3). Alternatively, nitronate 10 can be attacked by the nucleophilic aniline moiety in 5 to form an Ophosphorylated N'-hydroxyacetimidamide species that can be viewed as a direct analog of intermediate 6 (with X = NOPO(OH)2). This intermediate can undergo a subsequent Vilsmeier-type intramolecular attack involving the nucleophilic C-3 center of indole moiety to produce indologuinoline 1 in a way similar to the one described in Scheme 1.

To test this idea, we subjected 2-phenyl-1H-indole (5aa) to the reaction with excess nitromethane (9a) (ca. 5 equiv.) in PPA (80% P₂O₅). Gratifyingly, the reaction proceeded smoothly at 100 °C forming the desired indologuinoline 1aaa as the sole product, which was isolated in 92% yield (Scheme 4). Large excess of nitromethane was used in this case to offset its high volatility at high temperature, and probably its amount could be reduced upon scale up or in the case when the process is carried out in a closed pressurized reactor. For this study, however, to simplify the set up and reduce the reagent cost, we preferred to use an open vessel and excess of nitromethane. Reactions with nitroethane (9b, ca. 3 equiv.) and 1-nitrohexane (9c, 1.2 equiv.) also proceeded uneventfully at 110 °C affording the corresponding tetracyclic products 1aab and 1aac in 91 and 88% yields, respectively (Scheme 4). Expectedly, lower

Scheme 4

volatility of nitroalkane 9c allowed a nearly stoichiometric amount of this reagent to be used.

Also, we tested if the featured method is compatible with the ester functionality. To this end, reaction of 5aa in PPA was carried at $130~^{\circ}\text{C}$ in the presence of nitroacetic ester (9e, 1.2 equiv.). We were pleased to find that the corresponding functionalized indoloquinoline 1aae was obtained smoothly in high yield (Scheme 4).

With this reaction protocol in hand, we proceeded with the development of a one-pot method allowing for an *in situ*

assembly of indole moiety 5 *via* Fischer indolization of readily available arylhydrazines 2 and *ortho*-aminoacetophenones 3. To this end, phenylhydrazine 2a was first heated with an equimolar amount of acetophenone 3a for a short period of time without solvent to assemble hydrazine 12. The addition of PPA and heating at 70 °C triggers the Fischer indolization, which proceeded within 3 h to afford indole 5aa. Nitroalkanes could be then added directly to this mixtures and the temperature of the mixture should be raised (100 °C for nitromethane (9a), 110 °C for longer linear nitroalkanes (9b, c), 130 °C for ethyl

nitroacetate (9e), 140 °C for α -nitrotoluene (9d)). The optimal excess amounts of the nitroalkanes are shown in Scheme 4. We were pleased to find that via this one-pot protocol indoloquinoline products were obtained in high yields, which were only slightly lower than those obtained from indole precursor 5a (Scheme 4).

Next, we evaluated the possibility of employing N-methyl protected aminoacetophenone 3b as one of the key reactants. If successful, the application of this starting material would provide access to derivatives of 5H-indolo[3,2-c]quinoline, including the natural alkaloid isocryptolepine. To our delight, a three component one-pot reaction between hydrazine 2a, acetophenone 3b, and nitromethane (9a) afforded isocryptolepine (1aba) in high yield (Scheme 4). The replacement of nitromethane with nitroethane (9b) or 1-nitrohexane (9c) allowed for the efficient preparation of homologs 1abb and 1acc, respectively (Scheme 4). To further elaborate on the newly developed method we decided to synthesize a small library of isocryptolepine analogs by varying hydrazine and acetophenone reagents, capitalizing mostly on the use of nitromethane to take advantage of its low cost. Also, taking into account the results of our preliminary SAR studies, we concentrated on the preparation of structures bearing alkoxy substituents in ring D, as they demonstrated the highest antiproliferative potency. Reactions involving p-tolylhydrazine 2b proceeded smoothly affording the corresponding indologuinolines 1bba, 1bbb, and 1bbd in good yields. It is important to mention that straight after aqueous work up of the reaction mixtures all of these products were sufficiently pure for all practical purposes.

The employment of *meta*-substituted arylhydrazines **2c** and **2d** proved to be more challenging as the Fischer indolization reaction in this case suffered from the lack of regioselectivity. When carried out at elevated temperatures (above 90 °C) it afforded nearly equimolar mixtures of 4- and 6-substituted indole intermediates, which were further converted into

a mixture of 9- and 7-substituted indoloquinolines. The problem was solved by carrying out the indolization at lower temperatures (below 70 °C), which led to 6-substituted indoles predominantly (the typical product ratio was about 9: 1 for both hydrazines 2c and 2d). The subsequent reaction with nitroalkanes resulted in the formation of 9-methyl- (1cea, 1cfa) or 9-chloro- (1daa, 1dda, 1dea) substituted indoloquinolines along with minor amounts of 7-substituted isomers. This impurity was removed by preparative column chromatography, which lowered the yields of purified materials (Scheme 4).

In continuation of our studies of isocryptolepine and its analogs as potential anticancer agents, we evaluated the synthesized compounds against a panel of six cancer cell lines that included the human A549 non-small cell lung cancer (NSCLC), MCF-7 breast carcinoma, U373 glioblastoma, HS683 oligodendroglioma, SKMEL-28 melanoma as well as murine B16F10 melanoma using cryptolepine as control (Table 1). The A549, U373 and SKMEL-28 cell lines represent in vitro cell models of the corresponding aggressive cancer types, i.e. NSCLCs, glioma and melanoma because they have been shown to display various degrees of resistance to pro-apoptotic insults.10 It was found that methylated compounds, including isocryptolepine (1aba) itself, in general had higher activities than their demethylated counterparts. A number of such compounds were equally or more potent than the parent alkaloid itself. These included 1abb, 1aea, 1bbb, 1cea, 1dea. It also appeared that alkoxy groups in the quinoline part of the molecule, such as in compounds 1aea, 1cea, 1cfa, 1dea were beneficial for activity. Of note however, demethylated analog 1cfa, which contains bridged alkoxy groups in the quinoline part of the molecule showed nanomolar antiproliferative activity, more than two orders of magnitude more potent than that of isocryptolepine (1aba). Synthesis of a methylated analogue of 1cfa, which is expected to have even higher potency is underway.

Table 1 50% in vitro growth inhibitory concentration (IC $_{50}$, μ M a) determined by the colorimetric MTT assay

| Compound | Carcinoma | | Glioma | | Melanoma | | Mean + |
|-------------------|-----------|-------|--------|-------|----------|--------|---------------|
| | A549 | MCF-7 | U373n | HS638 | SKMEL-28 | B16F10 | SEM |
| 1aaa ^b | 8 | 27 | 16 | 4 | 24 | 7 | 14 ± 4 |
| 1aab | 3 | 4 | 6 | 3 | 6 | 3 | 4.0 ± 0.5 |
| 1aac | 17 | 27 | 24 | 11 | 22 | 4 | 17 ± 4 |
| 1aae | 4 | 5 | 5 | 4 | 4 | 3 | 4.0 ± 0.3 |
| 1aba b | 0.6 | 1.0 | 1.3 | 0.4 | 0.9 | 0.7 | 0.8 ± 0.1 |
| 1abb | 0.5 | 1.3 | 3.3 | 0.5 | 1.0 | 0.7 | 1.2 ± 0.4 |
| 1aea ^b | 0.4 | 0.5 | 0.6 | 0.3 | 0.4 | 0.3 | 0.4 ± 0.05 |
| 1bba | 3 | 4 | 4 | 1 | 4 | 2 | 3.0 ± 0.4 |
| 1bbb | 0.4 | 0.6 | 1.2 | 0.4 | 0.7 | 0.3 | 0.6 ± 0.1 |
| 1bbd | 4 | 6 | 4 | 4 | 4 | 3 | 4.1 ± 0.4 |
| 1cea | 0.4 | 0.7 | 1.6 | 0.4 | 0.5 | 0.4 | 0.7 ± 0.2 |
| 1cfa | 0.008 | 0.05 | 0.27 | 0.01 | 0.007 | 0.008 | 0.06 ± 0.04 |
| 1daa | 5 | 6 | 2 | 4 | 6 | 3 | 4.0 ± 0.7 |
| 1dda | 2 | 3 | 3 | 1 | 2 | 0.6 | 2.0 ± 0.4 |
| 1dea | 0.04 | 0.18 | 0.24 | 0.06 | 0.65 | 0.21 | 0.2 ± 0.1 |

^a Mean concentration required to reduce the cell viability by 50% after a 72 h treatment relative to a control as determined by MTT assay. Each experiment was performed once in sextuplicate. ^b These data were previously reported in ref. 4.

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Conclusion

In conclusion, we designed and developed an improved one-pot assembly of 5*H*- and 11*H*-indolo[3,2-*c*]quinoline skeleton. The method involved an *in situ* formation of hydrazones, followed by a PPA-assisted Fischer indolization reaction and intramolecular cycloaddition of electrophilically activated nitroalkanes. Using this protocol, we synthesized the natural alkaloid isocryptolepine and a small library of its analogs, which were evaluated for anti-proliferative activity against six cancer cell lines. Several analogs demonstrated activity higher than that of the natural alkaloid, with one compound (1cfa) having significant activity at nanomolar concentrations.

Experimental part

General information

¹H and ¹³C NMR spectra were recorded on a Bruker Avance-III spectrometer (400 or 100 MHz, respectively) equipped with a BBO probe in $CDCl_3$ or $DMSO-d_6$, using TMS as internal standard. High-resolution mass spectra were registered with a Bruker Maxis spectrometer (electrospray ionization, in MeCN solution, using HCO2Na-HCO2H for calibration). Melting points were measured with a Stuart smp30 apparatus. All reactions were performed in oven-dried 5 mL round-bottomed flasks open to the atmosphere, employing overhead stirring. The reaction progress and purity of isolated compounds were controlled by TLC on Silufol UV-254 plates, with EtOAc as eluent. All reagents and solvents were purchased from commercial vendors and used as received. Indoloquinoline products 1aaa, 11 1aab, 12 1aac, 4 1aba, 10 1abb, 11 1abc, 4 1aca, 13 1ada, 3g 1aea, 3g 1bba, 4 1bbb, 12 1bbd6 are known compounds and their physical and spectral properties matched those previously reported in literature.

Preparation of indoloquinolines *via* three-component one-pot protocol (general procedure A)

Reaction vessel was charged with arylhydrazine 2 (1.0 mmol) and o-aminoacetophenone 3 (1.0 mmol), and the mixture was heated at 120 °C for 10 min. Then, the mixture was cooled down to ca. 70 °C, and polyphosphoric acid (80% PPA, 2.0 g) was added. The mixture was stirred at the same temperature for 3 h, then the nitroalkane of choice was added in the listed excess (5.0 mmol (305 mg, 368 μL) of nitromethane (9a), 3.0 mmol (225 mg, 214 μL) of nitroethane (9b), 1.2 mmol (157 mg, 167 μL) of 1-nitrohexane (9c), 1.4 mmol (192 mg, 160 μ L) of α -nitrotoluene (9d), or 1.2 mmol (160 mg, 133 μ L) of ethyl α -nitroacetate (9e)). The reaction mixture was stirred at the appropriate temperature (100 °C for reactions with 9a, 110 °C for 9b, c, 130 °C for **9e**, and 140 °C for **9d**) for 5 h, then poured into water (50 mL) and neutralized with 25% aqueous ammonia. The formed precipitate was filtered out and dried on air. Usually such material was sufficiently pure for all practical application, but for analytical purposes it could be additionally purified by flash column chromatography on silica gel, eluting with

mixture acetone/benzene (1:3) doped with aqueous ammonia (1 drop of per 10 mL of eluent).

Preparation of indoloquinolines from indole (general procedure B)

The reaction vessel was charged with 2-phenyl-1*H*-indole (5aa) (193 mg, 1.0 mmol) and polyphosphoric acid (80% PPA, 1.5 g). The mixture was stirred at 70 °C, and nitroalkane was added according to the specifications described in general procedure A. Temperature regimen, post-reaction work up, isolation and purification of the products was performed in the same manner as described in general procedure A.

Ethyl 11*H*-indolo[3,2-*c*]quinoline-6-carboxylate (1aae)

Yield 78% (general procedure A), 89% (general procedure B). Light-yellow crystalline solid, mp 91–92 °C (acetone–hexane), $R_{\rm f}$ 0.46 (Hex/EtOAc/EtOH 7 : 2 : 1). ¹H NMR (400 MHz, CDCl₃) δ 11.39 (s, 1H), 8.63 (d, J = 7.8 Hz, 1H), 8.37 (d, J = 8.1 Hz, 1H), 8.25 (d, J = 8.4 Hz, 1H), 7.63 (d, J = 8.0 Hz, 1H), 7.58 (t, J = 7.6 Hz, 1H), 7.50–7.42 (m, J = 15.6, 7.8 Hz, 2H), 7.36 (t, J = 7.6 Hz, 1H), 4.61 (q, J = 7.1 Hz, 2H), 1.39 (t, J = 7.1 Hz, 3H); ¹³C NMR (101 MHz, CDCl₃) δ 166.6, 144.0, 143.4, 142.6, 139.6, 129.4, 129.3, 127.6, 126.8, 124.2, 121.8, 121.6, 121.5, 117.4, 113.5, 111.8, 62.7, 14.4; IR (KBr, film, cm⁻¹): 2930, 2855, 2368, 1730, 1565, 1501, 1460, 1355, 1310, 1238, 1193, 1081, 1017, 744; HRMS (ES TOF) calcd for $C_{18}H_{15}N_2O_2$ (M + H)⁺ 291.1128, found 291.1127 (0.4 ppm).

2,3-Dimethoxy-5,9-dimethyl-5*H*-indolo[3,2-*c*]quinoline (1cea)

Yield 58% (*via* general procedure A). Yellow crystalline solid, $R_{\rm f}$ 0.20 (acetone/benzene/ammonia 8 : 8 : 1), mp 252–253 °C (CH₂Cl₂–benzene–hexane), Lit mp 252–253 °C.^{3g} ¹H NMR (400 MHz, DMSO) δ 9.14 (s, 1H), 7.95–7.88 (m, 2H), 7.48 (s, 1H), 7.25 (s, 1H), 7.05 (d, J = 7.8 Hz, 1H), 4.19 (s, 3H), 3.98 (s, 3H), 3.96 (s, 3H); ¹³C NMR (101 MHz, DMSO) δ 151.5, 149.9, 148.2, 136.5, 135.4, 131.1 (2C), 121.9, 121.4, 119.4, 116.5, 114.9, 113.6, 103.0, 99.2, 56.1, 55.8, 42.8, 21.9; HRMS (ES TOF) calcd for C₁₉H₁₉N₂O₂ (M + H)⁺ 307.1441, found 307.1443 (0.8 ppm).

10-Methyl-2,3-dihydro-12*H*-[1,4]dioxino[2,3-*g*]indolo[3,2-*c*] quinoline (1cfa)

Yield 51% (*via* general procedure A). Colorless crystalline solid, $R_{\rm f}$ 0.60 (acetone/benzene/ammonia 8 : 8 : 1). Decomposed without melting at 380 °C (DMF-dioxane). Lit decomp. 380 °C. ^{3g} ¹H NMR (400 MHz, DMSO) δ 12.29 (s, 1H), 9.33 (s, 1H), 8.09 (d, J = 8.0 Hz, 1H), 7.90 (s, 1H), 7.50 (c, 1H), 7.47 (c, 1H), 7.12 (dd, J = 8.0, 0.7 Hz, 1H), 4.39–4.41 (m, 4H), 2.51 (s, 3H); ¹³C NMR (101 MHz, DMSO) δ 145.2, 143.4, 142.8, 141.3, 139.3, 139.0, 134.8, 122.0, 119.9, 119.6, 114.4, 113.2, 112.3, 111.6, 106.9, 64.4, 64.2, 21.7; HRMS (ES TOF) calcd for $C_{18}H_{15}N_2O_2$ (M + H)⁺ 291.1128, found 291.1126 (0.7 ppm).

9-Chloro-11H-indolo[3,2-c]quinoline (1daa)

Yield 63% (*via* general procedure A). Colorless crystalline solid, $R_{\rm f}$ 0.60 (acetone/benzene/ammonia 8 : 8 : 1), mp 325 °C with sublimation (dioxane), Lit mp 325 °C. ^{3g} ¹H NMR (400 MHz,

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DMSO) δ 12.89 (br. s, 1H), 9.61 (s, 1H), 8.52 (dd, J = 8.0, 1.2 Hz, 1H), 8.35 (d, J = 8.4 Hz, 1H), 8.15 (dd, J = 8.3, 0.6 Hz, 1H), 7.70–7.79 (m, 3H), 7.38 (dd, J = 8.4, 1.9 Hz, 1H); 13 C NMR (101 MHz, DMSO) δ 145.6, 144.9, 140.4, 139.4, 130.0, 129.7, 128.4, 126.0, 122.2, 121.6, 121.0, 120.8, 117.0, 113.9, 111.6; HRMS (ES TOF) calcd for $C_{15}H_{10}ClN_2$ (M + H) $^+$ 253.0527, found 253.0528 (0.6 ppm).

9-Chloro-2,3-dimethoxy-5-methyl-5*H*-indolo[3,2-*c*]quinoline (1dea)

Yield 61% (*via* general procedure A). Yellow crystalline solid, $R_{\rm f}$ 0.39 (acetone/benzene/ammonia 8 : 8 : 1), mp 270–271 °C (dioxane–EtOH), Lit mp 270–271 °C.^{3g} ¹H NMR (400 MHz, DMSO) δ 9.17 (s, 1H), 8.04–7.98 (m, 2H), 7.70 (d, J = 1.8 Hz, 1H), 7.31 (s, 1H), 7.16 (dd, J = 8.2, 1.9 Hz, 1H), 4.20 (s, 3H), 4.00 (s, 2H), 3.99 (s, 3H); ¹³C NMR (101 MHz, DMSO) δ 155.5, 153.7, 151.2, 148.0, 136.8, 131.0, 129.5, 124.3, 120.6, 118.7, 117.3, 115.1, 115.0, 103.3, 99.5, 56.0, 55.9, 42.5; HRMS (ES TOF) calcd for $C_{18}H_{16}ClN_2O_2$ (M + H)⁺ 327.0895, found 327.0886 (2.8 ppm).

9-Chloro-2,3-dimethoxy-11H-indolo[3,2-c]quinoline (1dda)

Yield 54% (via general procedure A). Off-white crystalline solid, mp 309–310 °C (acetone/hexane), Lit mp 309–310 °C. 3g $R_{\rm f}$ 0.55 (acetone/benzene/ammonia 8 : 8 : 1). ¹H NMR (400 MHz, DMSO) δ 12.53 (s, 1H), 9.39 (s, 1H), 8.26 (d, J = 8.4 Hz, 1H), 7.91 (s, 1H), 7.71 (d, J = 1.6 Hz, 1H), 7.53 (s, 1H), 7.32 (dd, J = 8.4, 1.5 Hz, 1H), 3.99 (s, 3H), 3.95 (s, 3H); ¹³C NMR (101 MHz, DMSO) δ 150.9, 148.8, 142.2, 142.2, 140.3, 139.3, 129.6, 121.3, 121.0, 120.5, 113.1, 111.3, 111.0, 109.2, 101.2, 55.8, 55.6; HRMS (ES TOF) calcd for $C_{17}H_{14}ClN_2O_2$ (M + H)⁺ 313.0738, found 313.0732 (1.9 ppm).

Cancer cell growth inhibition

Cell lines used to evaluate the growth inhibitory effects of the compounds were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA), the European Collection of Cell Culture (ECACC, Salisbury, UK) and the Deutsche Sammlung von Mikroorganismen und Zellkulturen (DSMZ, Braunschweig, Germany). The human cell lines breast carcinoma MCF-7 (DSMZ ACC115), oligodendroglioma Hs683 (ATCC HTB138), non-small cell lung cancer A549 (DSMZ ACC107), glioblastoma U373 cells (ECACC 08061901), melanoma SKMEL-28 (ATCC HTB72) and the murine melanoma B16F10 (ATCC CRL-6475) cells were cultured in RPMI supplemented with 10% FBS, 4 mM glutamine, 100 μg mL⁻¹ gentamicin, and 200 units per mL to 200 μg mL⁻¹ penicillin-streptomycin. The cell lines were cultured in flasks, maintained and grown at 37 °C, 95% humidity and 5% CO2. The evaluation of the antiproliferative effects of the compounds on these cell lines was performed through the colorimetric assay MTT.14 Firstly, cells were trypsinized and seeded in 96 well plates. After 24 h, cells were treated with the compounds at different concentrations, ranging from 10 nM to 100 μ M or left untreated for 72 h. The compounds' dilutions in culture medium were prepared from a stock solution in DMSO (10 mM). Estimation of cell viability was performed by means of the MTT - (3-(4,5-dimethylthiazol-2yl)-2,5-diphenyl tetrazolium bromide) (Sigma, Bornem, Belgium) mitochondrial reduction into formazan by living cells. The optical density of the untreated control after 72 h was normalized as 100% of viable cells allowing determination of the concentration that reduced their global growth by 50% after 72 h of treatment.

Conflicts of interest

There are no conflicts to declare.

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