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Letter

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# Synthesis, Biological Evaluation of Fluorescent 23-Hydroxybetulinic Acid Probes and Their Cellular Localization Studies

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ABSTRACT: 23-Hydroxybetulinic acid (23-HBA) is a complex lupane triterpenoid, which has attracted increasing attention as an anticancer agent. However, its detailed mechanism of anticancer action remains elusive so far. To reveal its anticancer mode of action, a series of fluorescent 23-HBA derivatives conjugated with coumarin dyes were designed, synthesized and evaluated for their anti-proliferative activities. Subcellular localization and uptake profile studies of representative fluorescent 23-HBA probe 26c were performed in B16F10 cells and the results suggested that probe 26c was rapidly taken up into B10F10 cells in a dose-dependent manner and mitochondrion was the main site of its accumulation. Further mode of action studies implied that the mitochondrial pathway was involved in 23-HBA-mediated apoptosis. Together, our results provided new clues for revealing the molecular mechanism of natural product 23-HBA for its further development into an anti-tumor agent.

KEYWORDS: betulinic acid, fluorescent probe, anti-tumor, mechanism, mitochondrion

Natural products represent an enormous source of pharmacologically active compounds, and are often used as the starting point in modern drug discovery.<sup>1</sup> 23-Hydroxybetulinic acid (23-HBA, 1, Fig. 1) is a natural lupane-type pentacyclic triterpene isolated from the root of traditional Chinese herb *Pulsatilla chinensis*.<sup>2</sup> In recent years, 23-HBA has been attracting considerable attention due to its remarkable antineoplastic activities.<sup>3-6</sup> Several reports have indicated that the anticancer mechanisms of 23-HBA or its derivatives may be associated with depolarization of the mitochondrial membrane potential (MMP) and subsequent cell apoptosis.<sup>7-8</sup> However, the detailed molecular mechanism and targets underlying the antitumor activities of 23-HBA is still not fully understood.

In recent years, much progress has been made in establishing methods for the identification of the cellular targets or mode of action (MOA) of small molecules,<sup>9</sup> but current approaches mostly require lysis or rely on indirect assays. Ideal strategies that are capable of sensitive detection of biological processes in living systems (e.g., live cells, tissues, or animals) are required for the MOA studies and target proteins identification.<sup>10</sup>

Fluorescence-based cell-imaging strategies are highly useful in monitoring biological processes in living cells owing to their practicality and high sensitivity.<sup>11-12</sup> In recent years, fluorescently labelled probes of bioactive small molecules have become powerful tools to explore biological phenomena and provide insights into the mechanism of actions of these small molecules.<sup>13-16</sup> With the help of fluorescently labelled probes, this approach makes it applicable in imaging of drug uptake and subcellular distribution.<sup>17</sup> Besides, it can make the drug-target complex visible by using in-cell imaging, or in-gel



23-Hydroxybetulinic acid (1) Fluorescent 23-hydroxybetulinic acid probe fluorescence scanning after the sodium dodecyl sulfatepolyacrylamide gel electrophoresis (SDS-PAGE) separation.<sup>18</sup>

Figure 1. Overall strategy of fluorescent 23-HBA probe for subcellular localization and mechanism studies.

In our previous studies, we have prepared hundreds of derivatives of 23-HBA to develop novel potential anticancer therapeutics.<sup>3-6</sup> However, the detailed anticancer mechanism, mode of action, and intracellular target of 23-HBA remain elusive. To better understand the biological mechanism and illuminate mode of action of 23-HBA, Soural et al. employed a solid-phase synthetic approach to prepare biotinylated 23-HBA, <sup>19</sup> however, no fluorescent 23-HBA probes have been reported so far to the best of our knowledge.

As our continuous interests in developing novel fluorescent probes of natural products for mechanism of action studies<sup>14</sup>, we herein report the synthesis, biological activities, and labeling performance of fluorescent 23-HBA probes. We



mechanism of 23-HBA, and ultimately accelerate its development as an antitumor agent.



Various conjugation strategies have been developed to construct fluorescent probes of natural products, and the most practical method is attaching a small-molecule fluorescent probe to the tolerated position of natural products with appropriate linkers.<sup>20-21</sup> In this work, coumarin was selected as the fluorescent probe due to its water-solubility, small size, photostability and nontoxicity.<sup>22-23</sup> Various linkers were employed because linker type and length may often affect the anti-proliferative activities of probes and their interactions with target proteins. And above all, fluorescent probes must be attached to the compounds at a position that does not interfere with target protein binding.<sup>24</sup> We recently synthesized hundreds of derivatives of 23-HBA and found that the C-23 hydroxy on 23-HBA can be conjugated with a bulky group without affecting its ability to inhibit cancer cell growth, and therefore could be used to attach fluorescent probes.

Fluorescent N, N-diethyl-7-aminocoumarin dyes were synthesized according to the route depicted in Scheme 1. Briefly, 4-(diethylamino)salicylaldehyde was reacted with diethyl malonate in the presence of catalytic piperidine to yield coumarin-3-carboxylic acid ethyl ester (4). The ester was then hydrolyzed with sodium hydroxide to obtain acid 5. Compound 6 was prepared by the reaction of 5 with Nhydroxysuccinimide in CH<sub>2</sub>Cl<sub>2</sub>. Derivatives 7a-b were synthesized by the reaction of compound 6 with corresponding amino acids while 8a-b were prepared from the corresponding mono-N-boc-diamines. Following deprotection of amino group with trifluoroacetic acid afforded 9a-b. Treated with succinic anhydride in the presence of DMAP, compounds 9a**b** were converted to acids **10a-b**. On the other hand, compound 5 afforded alcohols 13a-c upon reaction with corresponding glycols, which were reacted with succinic anhydride to yield subsequent compounds 14a-c.

Compounds 12 and 17 were synthesized by reacting with corresponding ethanolamines following the steps described above. In addition, in order to understand the impact of water-

soluble ethylene glycol linkers on anti-proliferative activities of target compounds, **23** was synthesized from diglycolamine.

23-HBA was treated with BnBr in DMF to yield 28-benzyl-23-hydroxybetulinic ester (24, Scheme 2). Compounds 7a-b

were converted to acyl chloride and then reacted with ester 24 using DMF as a catalyst in dry  $CH_2Cl_2$  to give compounds 25a-b. Target compounds 26a-b was obtained by debenzylation of ester 25a-b with 10% Pd/C as catalyst under atmospheric pressure of hydrogen. Compounds 26c-j were prepared in a route similar to that for 26a-b.

#### Scheme 2. Synthesis of fluorescent 23-HBA probes 26a-j<sup>a</sup>



<sup>*a*</sup>Reagents and conditions: (a) BnBr,  $K_2CO_3$ , DMF, rt, 12 h; (b) Corresponding coumarin acids, oxalyl chloride, triethylamine, cat. DMF, CH<sub>2</sub>Cl<sub>2</sub>, rt, 0.5 h; (c) H<sub>2</sub>, 10% Pd/C, THF/MeOH, 50 °C, 1 h.

Ideally, the designed natural product fluorescent probes must retain the activity of this natural product. The *in vitro* anti-proliferative activities of newly synthesized fluorescent 23-HBA probes were evaluated on two cancer cell lines (B16F10 mice skin melanoma cells, MCF-7 human breast

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cancer cells) and one normal cell line (Helf human embryonic lung fibroblast cells) by MTT assay. The abilities of these new analogues to inhibit the growth of cells were summarized in Table 1. Most of the derivatives exhibited stronger antiproliferative activities than parent compound 23-HBA against three cell lines, especially on mice skin melanoma B16F10 cells. The anti-proliferative activities of 26a-i showed that linkers with appropriate length and the introduction of amido groups were favorable to their anticancer activities (Table 1). The most potent derivatives **26c** and **26d** with lower  $IC_{50}$ values on two tested tumor cell lines had diamine linkers while compounds 26e-g with ester linkers only had moderate antiproliferative activities ranging from 11.78 to  $35.52 \mu$ M. When compared with 26a-b, enhanced activity of 26j indicated that the introduction of water-soluble ethylene glycol linkers was beneficial to its anti-proliferative activity. It should be noted that coumarin dve (10a) exhibited no obvious toxicity against cancer cell lines. Given the potent anti-proliferative activity of compound 26c, it was finally selected as the candidate for further studies.

 Table 1. Anti-proliferative activities of fluorescent 23-HBA

 probes against different cell lines

Compounds	Cell lines $(IC_{50}, \mu M)^a$		
	B16F10	MCF-7	Helf
23-HBA	24.46±2.11	20.83±1.46	31.49±1.95
26a	11.27±0.92	$18.06 \pm 1.07$	$27.76 \pm 0.88$
26b	12.27±0.78	37.77±1.66	79.65±3.20
26c	4.07±0.05	6.26±0.27	11.70±0.85
26d	7.77±0.33	9.70±0.49	$17.58 \pm 1.21$
26e	11.78±0.56	19.67±2.43	23.81±2.08
26f	35.51±3.11	30.52±3.04	21.40±1.75
26g	29.99±3.44	24.23±2.51	19.38±1.38
26h	10.53±0.82	18.13±1.72	25.12±4.27
26i	11.50±1.81	19.90±2.06	24.03±1.53
26j	5.25±0.93	8.16±0.12	$10.85 \pm 1.05$
10a	>100	-	-
$ADM^b$	0.19±0.03	$0.15 \pm 0.02$	0.21±0.02
<sup>2</sup> Each data	represents mean	$\pm$ SD from the	ee independe

<sup>&</sup>quot;Each data represents mean  $\pm$  SD from three independent experiments; <sup>b</sup>ADM: Doxorubicin.

The fluorescent properties of compound 26c including the electronic excitation and emission spectra were evaluated firstly. The results showed that probe 26c exhibited significant differences between maximum excitation ( $\lambda_{exc}$  415 nm) and emission ( $\lambda_{em}$  475 nm) wavelength, namely stokes shift, which was suitable for the imaging studies (Supporting Information, Figure S1). Then, cell labeling property of probe 26c was investigated by incubating B16F10 cells with various concentrations of 26c in a time dependent manner. The results showed that probe 26c was taken up into B16F10 cells quickly, and distinguishable fluorescence could be detected as early as 15 min after incubation with 10  $\mu$ M 26c (Figure S2-5). To understand the detailed cellular uptake profile of compound **26c** in live B16F10 cells, quantitative research was carried out using flow cytometry. The results showed that after incubation with 10  $\mu$ M **26c**, the fluorescent intensity increased quickly at the beginning (0-15 min). Then, a slow growth in fluorescent intensity was observed during the following period of time (15-120 min). Subsequently, the fluorescent intensity grew rapidly (Figure S6). Interestingly, cells stained with naked

coumarin dye (10a) were used as a control and the results showed that no obvious fluorescence could be observed after 1 h staining with up to 50  $\mu$ M of 10a (Figure S7). Considering the poor cell permeability and cytotoxicity of naked coumarin dye (10a), the above results implied the good cell permeability of 23-HBA.



Figure 2. Subcellular colocalization of probe 26c with MitoTracker (mitochondria specific dye) in B16F10 cells. Images were collected using confocal microscopy with excitation of probe 26c at  $\lambda_{ex}$ = 415 nm and emission at  $\lambda_{em}$ = 475 nm, MitroTracker at  $\lambda_{ex}$ = 579 nm and emission at  $\lambda_{em}$ = 600 nm. Images are provided with amplification factor (a) enlarged view of representative cells, 60x, (b) 40x.

The search for cellular targets may well be regarded as a quest for a needle in a giant haystack, the traditional affinitybased proteomics ("pulldown") method often suffers from non-specific binders.<sup>25</sup> To minimize unspecific binding in affinity-based approaches, lysate pre-fractionation may be considered. Use of a particular subcellular fraction will reduce the sample complexity and enrich the target protein. Prerequisite to the fractionation is the knowledge about subcellular localization of the binding proteins which often can be obtained through colocalization study using fluorescently labeled compound. To understand the subcellular localization of 23-HBA in living cells, probe 26c and commercially available MitoTracker (Mitochondria specific dye) were employed for colocalization studies firstly. Taking into account the result of qualitative and quantitative research on cellular uptake, the minimum concentration of compound **26c** was optimized to 4  $\mu$ M and the most appropriate incubation time was 1 h. Cells were imaged with confocal laser scanning microscopy. The location of probe 26c was viewed as green fluorescent spots upon excitation at 415 nm (Fig. 2, left column, green), and the MitoTracker emitted red fluorescence at 579 nm (Fig. 2, right column, red). Overlapping the fluorescent images of probe 26c and MitoTracker showed an extensive orange image in the same area (Fig. 2, middle column, orange), revealing the colocalization of these two probes. This observations provided compelling evidence for the conclusion that mitochondrion is the main accumulation site of 23-HBA, which was consistent with reports revealing that 23-HBA's anti-tumor mechanism partially involved in effecting mitochondrial pathway. On the other hand, some non-mitochondrial regions in cells (near the nucleus) where 26c is also shown to accumulate are observed in Figure 2a, indicating the possibility of other proteins targeted by 23-HBA. To explain the specificity of the probe for mitochondria, comparative studies with lysosome and nucleus specific dyes are performed and results are shown in Figure 3. Overlapping of the fluorescent images arising from 26c and LysoTracker Red (lysosome specific dye) or DAPI (nucleus specific dye) didn't show obvious colocalization. As a result, fluorescent 23-HBA probes provided direct visual evidences that 23-HBA colocalized mainly with mitochondrion in living cell system, and following target identification performed with mitochondrion should be a priority.



**Figure 3.** B16F10 cells were costained with **26c** (4  $\mu$ M) and LysoTracker Red, and then analyzed under a confocal microscope. Nuclei were stained with DAPI. Blue: Nucleus; red: LysoTracker Red, green: **26c**. Micron bar: (a) 10  $\mu$ m; (b) 50  $\mu$ m.

Mitochondrion plays an essential role in intrinsic pathway of apoptosis. Thus, commercially available fluorescent probe JC-1 was used to measure the MMP of B16F10 cells affected by 23-HBA and **26c** using flow cytometry. As shown in Figure 4, B16F10 cells were incubated with different concentrations of 23-HBA (0, 7.5, 15, and 30  $\mu$ M) for 48 h prior to staining with the JC-1, the number of cells with collapsed MMPs in different groups were determined by flow cytometry analysis, yielding 1.79%, 10.57%, 22.85%, and 42.87% apoptotic cells, respectively. Similar decreased MMPs were also observed on B16F10 cells treated with 0, 1.5, 3.0, 6.0  $\mu$ M **26c**, this result suggests that fluorescent conjugated 23-HBA intracellularly behaves similar as the starting material 23-HBA.



**Figure 4.** Effect of compounds 23-HBA and **26c** on the mitochondrial membrane potentials of B16F10 cells. Cells were incubated with the indicated concentrations of compounds for 48 h prior to staining with JC-1.

The above results showed that 23-HBA-induced apoptosis was closely related to the mitochondrial pathway. The Bcl-2 family including pro-apoptotic proteins (e.g., Bax, Bad) and anti-apoptotic proteins (e.g., Bcl-2, Bcl-xL) has been identified as essential proteins in controlling the mitochondrial pathway. To confirm whether these proteins are involved in apoptosis induced by 23-HBA, the expression of Bax, Bcl-2, cytochrome *c*, caspase 3 and caspase 9 was examined by Western Blot analysis. As shown in Figure 5, the expression of Bax, cytochrome *c*, caspase 3 and caspase 9 began to increase after the treatment of 23-HBA or **26c** for 48 h, while the Bcl-2 expression decreased markedly. The above results suggested that mitochondrial death pathway was involved in 23-HBA-induced apoptosis.



**Figure 5**. Western Blotting analysis of compounds 23-HBA and **26c** on the expression of apoptosis-related proteins bax, bcl-2, cytochrome *c*, caspase 3 and caspase 9, GAPDH was used as the loading control. (a) B16F10 cells were treated with 23-HBA of various concentrations (0, 7.5, 15, and 30  $\mu$ M) and **26c** (0, 1.5, 3.0, and 6.0  $\mu$ M); (b) The density ratio of proteins to GAPDH was shown as relative expression, data was represented as mean  $\pm$  SD of three independent experiments. (c) The density ratio between Bcl-2 and Bax. \*\**P*<0.01, \*\*\**P*<0.001, control compared with treated cells.

In summary, in order to reveal the mechanism of anticancer action of 23-HBA, a series of fluorescent probes of 23-HBA were designed and synthesized by connecting 23-hydroxyl group of 23-HBA with coumarin dye through various linkers. The pharmacological activity, uptake profile and subcellular localization of fluorescent 23-HBA probes were investigated in living cells. Qualitative and quantitative research on cellular uptake indicated that probe 26c was rapidly taken up by B16F10 cells in a dose-dependent manner. Importantly, colocalization studies with organelle specific dyes revealed that mitochondrion is the main site of its accumulation and subsequent mode of action studies implied that the mitochondrial pathway was involved in the 23-HBA-mediated apoptosis. Overall, our findings indicate that the novel fluorescent probes of 23-HBA could be used for live cell imaging and further to elucidate the mechanisms of anticancer action of 23-HBA.

#### ASSOCIATED CONTENT

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#### **Supporting Information**

Experimental protocols and NMR spectra. This material is available free of charge via the Internet at <u>http://pubs.acs.org</u>.

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#### Notes

The authors declare no competing financial interest

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## ABBREVIATIONS

23-HBA, 23-Hydroxybetulinic acid; MOA, mode of action; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; EDCI, 1-(3-dimethylaminopropyl)-3ethylcarbodiimide hydrochloride; DMAP, 4dimethylaminopyridine; DMF, *N*, *N*-dimethylformamide; THF, tetrahydrofuran; MMP, mitochondrial membrane potential; JC-1, 5,5', 6,6'-tetrachloro-1,1',3,3'-tetraethylimidacarbocyanine iodide

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