Combination of HDE and BIIB021 efficiently inhibits cell proliferation and induces apoptosis via downregulating hTERT in myelodysplastic syndromes

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Abstract. Treatment for higher-risk patients with myelodysplastic syndrome (MDS) should aim to modify the disease course by avoiding progression to acute myeloid leukemia and improving survival. When a patient is not eligible for intensive chemotherapy and lacks a donor hematopoietic cell source, or for a patient in a poor economic situation, consideration can be given to the use of Chinese herbal medicine. Numerous plant extracts, such as camptothecin, vinblastine and paclitaxel, have been reported to display antitumor effects, serving as potential therapeutic strategies for cancer. In the present study, the ultra-performance liquid chromatography-tandem mass spectrometry system (Waters Corporation) was used to detect the main chemical components of HDE, CCK-8 assay to detect the effects of HDE and BIIB021 on the proliferation of SKM-1 cells; and designed hTERT-small interfering (si) RNAs to detect the effects of HDE and BIIB021 on SKM-1 cell apoptosis after HTERT gene knockdown. The present study investigated a newly extracted coumarin HDE, the active component in Oldenlandia diffusa Willd, which efficiently inhibited SKM-1 (MDS cell line) proliferation and induced apoptosis, as determined by performing Cell Counting Kit-8 and flow cytometry assays, respectively. The effect of HDE was associated with decreased telomerase activity. Moreover, heat shock protein 90 inhibitor BIIB021 significantly enhanced the antitumor effects of HDE on SKM-1 cells. In addition, SKM-1 cell apoptosis was increased in human telomerase reverse transcriptase (hTERT)-knockdown cells compared with the negative control group. Cell apoptosis in hTERT-knockdown SKM-1 cells was further enhanced following HDE, BIIB021 or combination treatment, as evidenced by increased levels of cleaved caspase 3, cleaved caspase 8 and cleaved poly ADP ribose polymerase. Collectively, the results indicated synergistic antitumor effects of HDE and BIIB021, providing a novel therapeutic combination for higher-risk MDS.

Introduction

Myelodysplastic syndromes (MDSs) are incurable malignant hematological diseases caused by malfunctioning hematopoietic stem cells (HSC) or CD34+ progenitor cells (1). A substantial proportion of MDSs arise in the setting of exposure to environmental or occupational toxins, including cytotoxic therapy for a prior malignancy or other disorder (2). Patients with MDS are classified as lower-risk or higher-risk according to different prognostic scoring systems (3,4). The incidence of MDS progression to acute leukemia is 3.3/100,000 (5). According to the revised International Prognostic Scoring System (2012), high-risk and extremely high-risk patients often display multiple morbid hematopoiesis and malignant clonal hyperplasia, which can easily transform into leukemia, leading to a median survival of 1.6 and 0.8 years, respectively (6). Due to its poor prognosis, the treatment of higher-risk MDS has received increasing attention. Stem cell transplantation remains the most effective treatment strategy for patients with higher-risk MDS, but only a few patients are eligible for this treatment strategy (3,7). The primary chemotherapy regimen for patients with higher-risk MDS is hypomethylating agents (HMAs) (8); however, as only ~50% of patients with higher-risk MDS respond to HMA treatment and the response is often transient, there is a need to improve the treatment strategy for higher-risk MDS.

On the genomic level, MDS is classified by losses and translocations involving certain key gene segments, with disruption of the normal structure and function of genes that control the balance of proliferation and differentiation of hematopoietic precursors (9). The evidence suggests that unidentified tumor
suppressor genes may serve important roles in the molecular mechanisms underlying MDS (10,11). Further molecular approaches to genetic lesions will aid with the identification of relevant tumor suppressor genes. Over the past few years, major signal transduction molecules, their genetic alteration molecules, cell cycle regulators, and transcription factors have been identified (9). In particular, transcription factors such as EVI-1, that regulate both hematopoietic stem cell proliferation and differentiation were identified (12). Disruption of signal transduction pathways involving these molecules results in ineffective multilineage hematopoiesis and bone marrow failure (13).

Activation of telomerase can lead to uncontrollable cell proliferation or even tumorigenesis, and ~90% of malignant and immortalized cells display abnormal telomerase activity (14). In contrast to normal cells, heat shock protein 90 (HSP90) is continuously activated in cancer cells (15) and highly expressed in hematological malignancies, especially in acute leukemia (16). HSP90 and the co-chaperone p23 bind to human telomerase reverse transcriptase (hTERT) to activate telomerase during tumorigenesis (17,18). HSP90 expression is higher in patients with higher-risk MDS compared with patients with lower-risk MDS, and is therefore positively correlated with the risk of MDS conversion to acute leukemia and poor prognosis (19,20). It was hypothesized that high HSP90 expression contributes to higher-risk MDS-induced leukemia via reactivation of hTERT; hence, inhibiting HSP90 activity and promoting substrate degradation with specific inhibitors should inhibit MDS cell proliferation and conversion. BIIB021, a derivative of the first sputum compound PU3, is a new generation terpenoid that is currently in clinical trials for the treatment of solid tumors and hematological malignancies (21,22). Therapeutically, higher-risk MDS is insensitive to common chemotherapy, and bone marrow transplantation is often hindered by human leukocyte antigen (HLA) mismatch, high costs, immune rejection and postoperative infections (23). Moreover, the use of demethylation drugs, such as 5-aza cytidine and 5-aza-2 deoxycytidine, is typically limited due to non-specific toxicity, potential carcinogenicity and drug resistance (24). Therefore, novel treatment strategies for higher-risk MDS with higher efficacy, reduced aggression, fewer side effects and lower cost are urgently needed.

Based on the difficulties and risks of higher-risk MDS in clinical treatment, the present study aimed to investigate the pathogenesis of the disease and combine the advantages of traditional Chinese medicine (for example, low toxicity and side effects) to evaluate its advantages for the treatment of higher-risk MDS. *Oldenlandia diffusa* Willd belongs to the hedgotis genus of rubiaceous family, which grows in the south of the Yangtze River in China (23). *Oldenlandia diffusa* Willd is a widely used Chinese herbal medicine that has been reported to display the following effects: lowering heat, detoxification, promoting blood circulation, clearing blood stasis and benefiting diuresis (23). Higher-risk and extreme higher-risk MDS are diagnosed as visceral dysfunction and Shengqi deficiency by the traditional Chinese medicine system (25). *Oldenlandia diffusa* Willd is a herbal prescription for the treatment of visceral dysfunction, and previous studies have indicated that that *Oldenlandia diffusa* Willd displays anti-inflammatory, antibacterial, immune-enhancing and antitumour effects (26-28).

It has been previously reported that the total coumarins of *Hedyotis diffusa* (TCHD) could inhibit SKM-1 cell (MDS cell line) proliferation (23). The present study aimed to preliminarily explore the potential use of HDE extracted from *Oldenlandia diffusa* Willd combined with HSP90 inhibitor BIIB021 for the treatment of MDS in vitro.

Materials and methods

*Plant material and reagents.* Authentic plant material of HDE was purchased from Zhejiang University of Traditional Chinese Medicine Chinese Medicine Decoction Pieces Co., Ltd. and identified by Dr Jianping Jiang at Zhejiang Chinese Medical University (voucher specimen: Jiang J.P., 141001, ZM). The dried and sliced HDE material was ground into fine powder before extraction. FBS and RPMI-1640 medium were purchased from Gibco (Thermo Fisher Scientific, Inc.). BIIB021 was purchased from Selleck Chemicals (cat. no. S1175). All other reagents were obtained from Sigma-Aldrich (Merck KGaA).

*Cell culture and preparation.* The SKM-1 cell line was purchased from The Cell Bank of Type Culture Collection of the Chinese Academy of Sciences. Cells were cultured with RPMI-1640 (Gibco; Thermo Fisher Scientific, Inc.) supplied with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) were cultured in RPMI-1640 (Gibco; Thermo Fisher Scientific, Inc.) and 100 U/ml penicillin-streptomycin under a humidified atmosphere of 5% CO₂ at 37°C and used at the logarithmic growth phase. Cells were digested and a cell suspension (8x10⁶ cells/ml) was prepared. Subsequently, 100 µl cell suspension was added to each well of a 96-well culture plate. Following culture for 24 h at 37°C with 5% CO₂, the drugs were diluted to 0, 25, 50, 100, 200 µg/ml concentrations and added to the SKM-1 cells with 100 µl per well. Following 48 h incubation for test, bone marrow-derived stem cells (BMSCs; cat. no. HUXMA-90011; Cyagen Biosciences, Inc.) were cultured in α-MEM supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc.) and 100 U/ml penicillin-streptomycin (HyClone; GE Healthcare Life Sciences) at 37°C with 5% CO₂ in a humidified atmosphere.

*Cytotoxicity assay.* The drugs were diluted to the required working concentrations in culture medium and 100 µl medium was added to each well. A negative control (NC) group (standard curve reference) and experiment control group were simultaneously set up. Cell cytotoxicity was assessed using the Cell Counting Kit-8 (CCK-8) assay (cat. no. 35003; Biosharp Life Sciences) according to the manufacturer's protocol. Following incubation for 48 h, 10 µl CCK-8 solution was added to each well and incubated for 2-3 h at 37°C. The optical density (OD) was measured at a wavelength of 450 nm using a microplate reader (BioTek Instruments, Inc.; EL-x800). Cell proliferation was calculated according to the following formula: proliferation (%) = (Adrug/Ablank)/(A_blank/A_blank) x 100, where Adrug represents the OD value of wells containing treated cells, A_blank represents the OD value of blank wells (no cells) and A_blank represents the OD value of wells containing untreated cells.

Small interfering (si)RNA transfection. siRNA sequences targeting hTERT were designed as follows: hTERT-siRNA-1728 sense, 5'-GGAAGAGUGUCUGGAGCAAGU-3' and antisense,
5'-UUGCUCCAGACACUCUCCCGG-3'; hTERT-siRNA-966 sense, 5'-CGGUGUACCGGAGACCAAGC-3' and antisense, 5'-UGGGUCUGCCGUACCCGGG-3'; and hTERT-siRNA-1464 sense, 5'-GGAAACACAGAGAGUUCACU-3' and antisense, 5'-AUGAACUCUCCGUAGGCUGUCCG-3'); control siRNA (NC) sense, 5'-UUCCUGCAAGGUUGCCUAGTT-3' and antisense 5'-ACGUAGACGACUGAGAATT-3'. SKM-1 cells (8x10^4 cells/well) were cultured in 24-well plates for at least 12 h. A total of 100 nM of each siRNA was transfected into SKM-1 cells using Lipofectamine 2000™ (Invitrogen; Thermo Fisher Scientific, Inc.) for up to 48 h.

**Real-time quantitative PCR analysis.** Total RNA was extracted from cells using the EasyPure™ Blood RNA kit (TransGen BioTech Co., Ltd.) according to the manufacturer's instructions. Total RNA was reverse transcribed to cDNA using the One Step Gscript Reverse Transcription System kit (Promega Corporation). The reaction mixture contained: 15 µl RNA, 1.2 µl random primer and 1.2 µl Oligo (dT). Following mixing and centrifugation at 1200 x g for 15 sec at room temperature, cDNA was preheated at 70°C for 5 min, placed on ice for at least 5 min and centrifuged at 1200 x g for at least 10 sec at room temperature. The second strand cDNA synthesis mixture contained: 6 µl 5X reaction buffer, 3.8 µl MgCl₂ (25 mM), 1.5 µl PCR nucleotide mix, 0.3 µl RNA enzyme inhibitor and 1 µl reverse transcriptase. The following thermocycling conditions were used: 25°C for 5 min, 42°C for 2 min and 60°C for 10 sec, followed by 40 amplification cycles.

The bands were quantified using Quantity One image analysis software (Bio‑Rad Laboratories, Inc.) with ACTB as the internal reference gene. Subsequently, cDNA was diluted using RNase‑free water up to 100 µl on ice. qPCR was performed using the following reaction mixture (20 µl): 5 µl cDNA, 1 µl primer (4 pmol/µl), 1 µl TaqMan Probe (6 pmol/µl; Genscript), 10 µl 2X Mix (Vazyme Biotech Co., Ltd.) and RNase‑free water. The following thermocycling conditions were used for qPCR: 37°C for 2 min, 95°C for 5 min, and denaturation at 95°C for 10 sec and 60°C for 30 sec, followed by 40 amplification cycles. The relative expression levels were analyzed according to the 2^ΔΔCT method (29). The sequences of primers and probes (NCBI Bank) were as follows: hTERT forward, 5'-GTGGTT TCTGTTGTGTTGTA-3' and reverse, 5'-GGAGTGGGAGGA GTGCTTGGG-3'; actin-β (ACTB) forward, 5'-GATGAGATTG CCTTCT-3' and reverse, 5'-TCACGGGATATGTTTAGAGG GCATGGCTTT-3' and reverse, 5'-ACGUAGACGACUGAGAATT-3' and TaqMan probe, 5'-6-FAM‑CATTGCC TGCTTACTC-3' and 6-FAM‑CATGGCC CTCACAGCACCATTGTA‑BHQ1-3'.

**Telomerase activity detection by TRAP-ELISA.** SKM-1 cell telomerase activity was assessed using the TELISA kit (cat. no. CSB-E08021h; Cusabio Technology LLC, USA) according to the manufacturer's instructions. Following treatment for 24 h, cells (1x10^5) were centrifuged at 4200 x g for 20 min at 4°C. The cell pellet was incubated with cold lysis buffer (200 µl) form the aforementioned kit, for 30 min on ice. The supernatant (2 µl) was used as the TRAP reaction template. Subsequently, 25 µl reaction mixture was added to the PCR reaction tube and the total volume was adjusted to 50 µl with DEPC-treated sterilized double distilled water to perform the TRAP reaction according to the following protocol: primer extension for 10 min at 25°C; telomerase inactivation for 5 min at 94°C; 30 cycles of denaturation for 30 sec at 94°C, annealing for 30 sec at 50°C and extension for 90 sec at 72°C; and extension for 10 min at 72°C. The amplified product (5 µl) and the denaturant (20 µl) were mixed and maintained at room temperature for 10 min. Subsequently, 225 µl hybridization solution (containing digoxin-labeled probe) was added, and after mixing, 100 µl mixture was added to an anti-biotin coated plate and incubated for 2 h at 37°C (300 rpm). Then peroxidase was added and developed with TMB substrate for 30 min at room temperature. Finally, 100 µl stop solution was added. The A value of each well (detection wavelength, 450 nm; reference wavelength, 690 nm) and the magnitude of the A value represented the level of telomerase activity.

**HDE extraction.** A total of 200 g Oldenlandia diffusa Wild was weighed. Subsequently, the plant was incubated with 5.4 180% ethanol for 30 min at room temperature, followed by reflux-extraction for 110 min at 80°C. After filtering, the filtrate was dried at 45°C to obtain Hedyotis diffusa. The extracts were suspended in 200 ml distilled water and separately re-extracted four times with an equal volume of petroleum ether, ethyl acetate or n-butanol. Subsequently, the extracts were combined. The extracts of Hedyotis diffusa were named HDP (petroleum ether extract), HDE (ethyl acetate extract), HBD (n-butanol extract) and HDW (water fraction). HDE (5 g) was suspended in 100 ml distilled water and HPD-300 macro-porous resin was used for dynamic adsorption at an adsorption flow rate of 0.5 ml/min (diameter:height = 1:10), and then eluted with water until the residue was colorless. Dynamic elution was performed with 60% ethanol at a flow rate of 1 ml/min. Finally, the eluate was collected and dried under reduced pressure.

**Chemical analysis of HDE.** The ultra-performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS) system (Waters Corporation) was used to determine the chemical constituents of HDE. The filtered sample (per 20 µl) was separated using an Inertsil C18 column (250x4.6 mm; 5 µm; Agilent Technologies Inc.) at 30°C, with a controlled flow rate of 1 ml/min at a set wavelength of 310 nm. The mobile phase was composed of (A) acetonitrile and (B) 0.05% methanoic acid and used for gradient elution of HDE at 0 min (85:15 v/v), 15-20 min (80:20 v/v) and 30-60 min (75:15 v/v). For mass spectrometry, ultra-high-purity argon (Ar) and high-purity nitrogen (N2) were used as the collision gas and the nebulizing gas, respectively. Ultra-high-purity argon (Ar) and high-purity nitrogen (N2) were used as the collision gas and the nebulizing gas, respectively. The ESI(-) source was operated under the following conditions: 2.0 kV source voltage, 38 psi nebulizer pressure, 350°C nitrogen gas temperature, with a controlled
flow rate of 1 ml/min, 100°C source temperature, 300°C capillary temperature, 500 l/h gas flow rate and full scanning mode. The CA ion (163 m/z) produced three primary fragments in MS/MS (119, 93 and 65 m/z) and the E-CSME ion (549 m/z) also produced three primary fragments in MS/MS (163, 119 and 89 m/z). Each chromatographic peak was identified by comparing the mass spectrum with the NIST mass spectral database (http://www.nist.gov/nist). For further verification, the retention time and mass data of each peak were compared with the reference standard compound. All chromatograms and mass spectra were analyzed using the Micromass MassLynx data system (MassLynx 4.1; Waters Corporation).

The total coumarin content of HDE was determined by spectrophotometry using a UV-vis spectrophotometer (Shimadzu Corporation) based on the method described by Wang et al (30). Briefly, 1.0 g dried HDE was dissolved in 100 ml 70% ethanol and soaked in the dark for 24 h at room temperature. After filtration, the absorbance of HDE solution at a wavelength of 310 nm was determined. The total coumarins were calculated using a standard curve prepared with reference standard p-coumaric acid (CA; Selleck Chemicals) and expressed in terms of mg of CA equivalents per g of dried HDE.

Flow cytometry analysis. SKM‑1 cells (8x10⁴ cells/well) in the logarithmic growth phase were seeded into 6-well plates. Cell apoptosis was assessed using the Apoptosis detection kit (Biouniquer Technology Co., Ltd.). Following drug treatment/transfection for 24 h, BMSCs and SKM‑1 cells were washed twice with pre-cooled PBS. Subsequently, 300 µl binding buffer was added to each well. Annexin V-FITC (5 µl) was added to each well, gently mixed and incubated in the dark for 15 min at room temperature. Subsequently, propidium iodide (10 µl) was added to each well, gently mixed and incubated for 10 min at room temperature in the dark. Stained cells were analyzed using a FACSCalibur (BD Biosciences) and CellQuest software (version 1.2; BD Biosciences).

Western blotting. Following treatment for 24 h, SKM‑1 cells and untreated BMSCs were harvested and total protein was extracted using 2X SDS lysis buffer (Cell Signaling Technology, Inc.). Total protein was quantified using the bicinchoninic acid assay (Sangon Biotech Co., Ltd.). A total of 20 µl protein samples was separated via 12% SDS-PAGE and transferred onto PVDF membranes. The membranes were blocked with 5% skimmed milk in TBST for 2 h at room temperature. Subsequently, the membranes were incubated with primary antibodies (1:1,000) overnight at 4°C. The primary antibodies included hTERT (cat. no. ab32020; Abcam), caspase-3 (cat. no. ab184787; Abcam), caspase-8 (cat. no. ab108333; Abcam), PARP (cat. no. ab139417; Abcam) and β-actin (cat. no. ab124964; Abcam). Following primary incubation, the membranes were incubated with a horseradish peroxidase-conjugated secondary antibody (1:5,000, Bio-Rad Laboratories, Inc.) at room temperature for 2 h. All antibodies were purchased from Cell Signaling Technology, Inc. β-actin was used as a loading control. Protein bands were visualized using Western blotting Luminol reagent (Biological Industries).

Statistical analysis. Statistical analyses were conducted using SPSS software (version 17.0; SPSS, Inc.). Comparisons between two groups were analyzed using unpaired Student’s t-tests. Comparisons among multiple groups were analyzed using one-way ANOVAs followed by the Dunnett's and Tukey's post hoc test with GraphPad Prism software (version 5.0; GraphPad Software, Inc.). P<0.05 was considered to indicate a statistically significant difference. Experiments were performed in triplicates.

Results

Chemical analysis of HDE. The UPLC-MS/MS method displayed high precision and sensitivity, with mass accuracy of <10 ppm (data not shown). The chromatogram (Fig. 1) displayed two distinct chromatographic peaks with good resolution and response in negative ion mode. Peak 1 and peak 2 were identified as CA and E-6-O-p-coumaroyl scandoside methyl ester (E-CSME), respectively, based on their MS and MS/MS spectra, as well as UV retention time, compared with the reference standard compounds. The CA ion (163 m/z) produced three primary fragments in MS/MS (119, 93 and 65 m/z), and the E-CSME ion (549 m/z) also produced three primary fragments in MS/MS (163, 119 and 89 m/z). The results indicated that the total coumarin content of HDE was 87.4%.

HDE combined with BIIB021 significantly inhibits SKM‑1 cell proliferation and telomerase activity. To determine the anti-MDS efficacy of extracted HDE and BIIB021, the inhibitory effects of the two drugs on SKM‑1 cell proliferation were assessed by performing the CCK-8 assay. The concentration ranged from 0 to 200 µg/ml for HDE and 0 to 400 nmol/l for BIIB021. SKM‑1 cell proliferation gradually decreased with increasing HDE and BIIB021 concentrations (Fig. 2A and B). The IC₅₀ of HDE was 98.03 µg/ml and the IC₅₀ of BIIB021 was 171.7 nmol/l in SKM‑1 cells, which were used for subsequent experiments.

Compared with the control group, HDE or BIIB021 treatment inhibited cell proliferation (P<0.01), whereas the inhibitory effect of the two drugs on cell proliferation was significantly increased in the HDE+BIIB021 combination group (Fig. 2C; P<0.01). To explore the mechanism underlying the inhibitory effects of HDE and BIIB021 on SKM‑1 cell proliferation, the telomerase activity in BMSCs and SKM‑1 cells was assessed. The results indicated that telomerase was highly activated in SKM‑1 cells compared with BMSC (Fig. 2D; P<0.01), but significantly inhibited by HDE, BIIB021 or combination treatment (Fig. 2D; P<0.01).

HSP90 expression. The expression of HSP90 in each group was assessed via RT-qPCR. HSP90 was highly expressed in SKM‑1 cells compared with BMSC (P=0.00016). HDE (P=0.0012), BIIB021 (P=0.0009) or combination (P=0.0005) treatment significantly reduced HSP90 expression levels in SKM‑1 cells compared with untreated SKM‑1 cells (Fig. 2E).

Decreased expression of hTERT and treatment with HDE+BIIB021 enhances the inhibitory effect and induces SKM‑1 cell apoptosis. siRNAs were synthesized to inhibit hTERT expression. The transfection efficiency of hTERT-siRNAs was
assessed, which indicated that hTERT-siRNA-966 significantly decreased hTERT expression compared with the control group (Fig. 3); therefore, hTERT-siRNA-966 was used for subsequent experiments. Subsequently, hTERT-siRNA-transfected SKM-1 cells were treated with HDE, BIIB021 or combination treatment, and cell apoptosis was assessed via flow cytometry (Fig. 4A). The results indicated that HDE or BIIB021 treatment significantly induced SKM-1 cell apoptosis compared with non-treated SKM-1 cells (Fig. 4B; P<0.01). hTERT knockdown combined with HDE or BIIB021 treatment significantly increased SKM-1 cell apoptosis compared with SKM-1 cells treated with HDE and hTERT knockdown combined with BIIB021 treatment also remarkably increased cell apoptosis when compared with SKM-1 cells treated with BIIB021 (Fig. 4B; P<0.01). Moreover, HDE+BIIB021 combination treatment-induced SKM-1 cell apoptosis was enhanced by hTERT knockdown (Fig. 4B; P<0.01).

hTERT knockdown-induced SKM-1 cell apoptosis is enhanced by treatment with single drug or combination of two drugs. To investigate the mechanism underlying HDE in promoting cancer cell apoptosis, treated BMSCs and SKM-1 cells were homogenized to measure the protein expression levels of hTERT, caspase 3, cleaved caspase 3, caspase 8, cleaved caspase 8, PARP and cleaved PARP by western blotting (Fig. 5). Compared with BMSCs, protein expression in SKM-1 cells was not notably altered. However, the expression levels of hTERT, caspase 3, cleaved caspase 3, cleaved caspase 8 and cleaved PARP were increased following HDE, BIIB021 or HDE+BIIB021 treatment compared with non-treated SKM-1 cells. Additionally, following hTERT knockdown, the expression levels of cleaved caspase 3, cleaved caspase 8 and cleaved PARP were slightly increased compared with the hTERT-siNC group. By contrast, there was no notable difference in protein expression levels among the BMSC, SKM-1 and hTERT-siNC groups. Furthermore, following HDE, BIIB021 or HDE+BIIB021 treatment, the expression levels of cleaved caspase 3, cleaved caspase 8 and cleaved PARP were increased in hTERT knockdown SKM-1 cells compared with hTERT-siNC-transfected SKM-1 cells. The results further
indicated that hTERT knockdown facilitated the antitumor effects of HDE and BIIB021.

Discussion

Ineffective hematopoiesis with peripheral cytopenia despite normo or hypercellular bone marrow is the hallmark of MDSs, especially in the low-risk subgroups. It is caused by increased secretion of inhibitory cytokines in the marrow; however, the underlying mechanisms are not completely understood (31). Immunoinhibitory or modulatory therapy aims to restore hematopoiesis by reversing inhibitory processes, and includes anti-thymocyte globulin (ATG), cyclosporin A (CSA),

![Figure 2. Effect of drug treatment on cell proliferation and telomerase activity in SKM-1 cells.](image)

![Figure 3. Transfection efficiency of hTERT siRNA.](image)

*P<0.05, **P<0.01.
direct TNF-inhibitors and lenalidomide (32). A simple scoring system, using age, duration of transfusion dependency and HLA-DR15 status, has been developed, which facilitates identification of patients with a high, intermediate or low probability of response to treatment with ATG (33). Other immunosuppressive agents that can also induce responses, such as CSA, have been combined with ATG, but have not yet been fully evaluated for their potential role in the treatment of MDS (32). Lenalidomide displays superior efficacy in the 5q-syndrome, but low efficacy in other patients with low-risk MDS (34). Additional clinical trials are required to define the potential of immunosuppressive and modulatory treatment in MDS (35,36). Thus, identifying safer and more effective anti-MDS strategies remains a major global health challenge. In our previous study, TCHD displayed proapoptotic effects in SKM-1 cells, and the underlying mechanism was associated with the inhibition of the PI3K/Akt signaling pathway and activation of caspase proteins (23). A further two new compounds, CA and E-CSME, were identified in TCHD (23). Also, the newly extracted compound of HDE efficiently inhibited leukemia cell proliferation and induced cancer cell apoptosis with our patented extraction technology (patent no. ZL201410324070.2).

Patients with higher-risk MDS with AML transformation display higher hTERT mRNA expression levels compared with healthy individuals and patients with low-risk MDS (37). hTERT-positive patients can easily develop leukemia (37), indicating that hTERT is positively related to the progression of MDS. Several natural drugs can inhibit the function of HSP90, such as benzoquinone ansamycin antibiotics, radicicol and neomycin, by competing with ATP for binding to the N-terminal conserved domain of HSP90 or interfering with...
the interaction between HSP90 and other accessory chaperone proteins, thus reducing the stability of chaperone protein (38).

Previously, the synergistic effect of traditional Chinese medicine extract and chemotherapy drugs has been reported (39). Similarly, a previous study has also demonstrated that both HDE and BIIB021 downregulated telomerase activity, and combined treatment further enhanced the tumor apoptosis-inducing effects of the compounds (40). BIIB021 outperforms other drugs as it displays independent anticancer effects with expression of multidrug resistance (MDR) proteins, such as P-gp and/or MDR-related protein 1, which typically confer resistance to a broad range of chemotherapeutic agents and molecularly targeted drugs (41-44). The prolonged inhibitory effects of BIIB021 on proliferation could increase its clinical applicability. In vitro experiments indicated that BIIB021 can competitively bind to the N-terminal ATP/ADP domain of HSP90, downregulate the expression of chaperones HER2, Akt, and Raf-1, and upregulate HSP70 and HSP27, with a broad spectrum of antitumor activities (20,45).

HDE may share a similar anticancer mechanism with BIIB021, but further investigation is required. In addition, HDE is extracted from the natural herb *Hedyotis diffusa*, which has been used for cancer treatment in China for centuries (46). Previous studies have demonstrated that TCHD could induce apoptosis in leukemia cell lines Kasumi-1, KG-1, THP-1, U937 and K562 (47,48). The present study in SKM-1 cells further demonstrated the broad-spectrum antileukemia benefits of TCHD. In the future, a preclinical study should be conducted to identify other purified coumarin components with fewer side effects and improved efficacy.

The present study indicated that downregulation of hTERT by siRNA might serve as a more direct strategy to inhibit cancer cell proliferation. Following hTERT knockdown, the flow cytometry results suggested that the apoptotic rate was increased in HDE- or BIIB021-treated SKM-1 cells. Antitumor effects were observed with a combination of the two drugs following hTERT knockdown. The western blotting results further indicated that cleaved caspase 3, cleaved

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Figure 5. Effects of HDE and BIIB021 on apoptosis-related protein expression levels in SKM-1 cells. (A) Apoptosis-related protein expression levels in SKM-1 cells were analyzed by western blot analysis. (B) Apoptosis-related protein expression levels were semi-quantified using gray scanning. BMSC, bone marrow-derived stem cell; hTERT, human telomerase reverse transcriptase; si, small interfering RNA; NC, negative control; PARP, peroxisome proliferator-activated receptor. *P<0.05, **P<0.01.
caspase 8 and downstream target cleaved PARP, which are involved in apoptosis pathways, were upregulated following hTERT knockdown compared with the hTERT-siNC group. Collectively, the results indicated an additive effect between the two drugs. However, the lack of cell proliferation analysis with trypan blue was a key limitation of the present study. Therefore, further mechanisms underlying HDE require investigation in future studies.

In conclusion, the present study identified a new active component HDE in *Oldenlandia diffusa* Willd, which displayed apoptosis-inducing effects on the MDS-derived leukemia cell line SKM-1, and the underlying mechanism, which involved downregulation of hTERT expression. The present study indicated that HDE, due to its potent anti-MDS activity, may be valuable for the development of new therapies for patients with MDS.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
SL, YunZ and BW supervised the project. BW, JJ and YS designed the study. YunZ, JJ, ST, SL, YS and LW performed the experiments. BW, ST and SL wrote the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Not applicable.

Patient consent for publication
Not applicable.

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

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