Introduction

Acute myeloid leukemia (AML) is the most common hematologic malignancy in adults with a high incidence rate and low survival probability [1,2,3]. AML progresses rapidly due to the rapid growth of abnormal white blood cells that accumulate in the bone marrow and interfere with the production of red blood cells, platelets, and normal white blood cells. If left untreated, AML is usually fatal within weeks or months after diagnosis. FLT3 (FMS-like tyrosine kinase 3), a cell surface receptor belonging to the class III receptor tyrosine kinase family, plays a pivotal role in the differentiation and survival of the hematopoietic stem cells in bone marrow [4,5]. FLT3 is one of the most commonly mutated genes in AML [6,7]. Activating FLT3 mutations, FLT3-ITD (an internal tandem duplication mutation in the juxtamembrane domain) and FLT3-TKD (a missense mutation within the kinase domain), are frequently observed in approximately 30% of adult AML patients [8,9,10,11]. FLT3-activating mutations critically regulate leukemic transformation by accelerating proliferation and suppressing apoptosis and are significantly associated with poor prognosis [12,13]. These findings highlight FLT3-ITD and FLT3-TKD as highly attractive therapeutic targets for drug development in human AML.

There are now several classes of small molecule FLT3 inhibitors that have entered clinical trials. However, effective drugs have not yet been identified in clinics [14,15,16]. Although these inhibitors have demonstrated promising anti-cancer activity in in vitro and in vivo preclinical models, clinically positive responses in AML patients receiving single-agent FLT3 inhibitors are limited due to the transient reduction of peripheral blasts but not bone marrow blasts or the occurrence of inhibitor-resistant FLT3 mutations in patients [17,18,19,20]. Therefore, combinatorial strategies of FLT3 inhibitors and other chemotherapeutic agents may be beneficial approaches to improve FLT3 inhibitor therapy and to overcome treatment failures [21,22]. The FLT3 inhibitor CEP-701 (lestaurtinib) combined with standard AML chemotherapeutic agents has the potential to improve clinical outcomes in AML patients [23]. In addition, histone deacetylase inhibitors (HDACi), a class of compounds that can induce cancer cell growth arrest and cell death by altering the acetylation status of both histone and non-histone proteins, can enhance the activity of FLT3 inhibitors on AML cell apoptosis [24,25,26]. The HDACi vorinostat (SAHA) exhibits clinical activity in AML; however, its efficacy as a single
agent is only moderate [27,28]. In this study, we report data characterizing the pharmacological profile of a new FLT3 kinase inhibitor, BPR1J-340, and elucidate the possible molecular mechanism of the strongly synergistic effects in combination with SAHA in FLT3-ITD+ cells.

The BPR1J-340 compound exhibits potent FLT3 inhibitory activity, with a 50% inhibitory concentration (IC50) of 25±5 nM and growth inhibitory effects on FLT3-ITD+ leukemia MOLM-13 and MV4;11 cells with a GC50 value of 3.4±1.5 and 2.8±1.2 nM, respectively. The IC50 values were approximately 1 nM against FLT3-ITD and 1 nM against STAT5 phosphorylation in MV4;11 cells. In addition, BPR1J-340 exhibits favorable pharmacokinetic properties and significant anti-tumor activity in FLT3-ITD murine xenograft models. The combination of the HDAC inhibitor SAHA with BPR1J-340 exhibits strongly synergistic anti-leukemia effect in FLT3-ITD+ cells. These results highlight the therapeutic potential of BPR1J-340 and SAHA in AML and support its preclinical or clinical development.

Materials and Methods

Chemicals and reagents

The FLT3 inhibitors, BPR1J-340 and AC220, were synthesized by our laboratory. The histone deacetylase inhibitor vorinostat (SAHA) was purchased from SelleckBio (Houston, TX, USA). All inhibitors were dissolved in dimethylsulfoxide (DMSO) at a stock concentration of 10 mM. The anti-FLT3 (sc-480, Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-p-FLT3-Tyr919 (#9363, Cell Signaling Technology, Beverly, MA, USA) anti-STAT5 (9363, Cell Signaling Technology), anti-pSTAT5-Tyr694 (9351, Cell Signaling Technology), anti-cleaved poly ADP-ribose polymerase (PARP) (9452, Cell Signaling Technology), anti-Mcl-1 (#4572, Cell Signaling Technology), anti-caspase 3 (#9662, Cell Signaling Technology) and anti-β-actin (GTX110546, GeneTex, Irvine, CA, USA) antibodies were purchased for Western blotting analysis. The preparation of recombinant proteins, FLT3 (residues 567–899), VEGFR1 (residues R781–11338) and VEGFR2 (residues V789–V1356), for biochemical kinase assay was described previously [29]. The VEGFR3 (residues M500–Y1363) proteins were purchased from Upstate (Billerica, MA, USA).

Cell culture

RS4;11, MV4;11, U937 and K562 cells were obtained from American Type Culture Collection (ATCC, Manassas, VA, USA). MOLM-13 cells were purchased from the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Braunschweig, Germany). All leukemic cell lines were grown in RPMI 1640 (Invitrogen, USA) with 10% fetal bovine serum (FBS) (Fisher Scientific, Pittsburgh, PA, USA). The HEK293T and FLT3-transfected HEK293T cells were cultured in DMEM (Invitrogen, USA) medium with 10% fetal bovine serum (FBS). The other 14 non-leukemic cell lines, HCC2927, H1975, H1650, H2228, NCH1460, A431, HCT-116, MKN45, MiaPaCa-2, RT4, MCF-7, HuH7, Hep3B, and Detroit 551, were cultured in medium according to the ATCC recommendations.

In vitro kinase activity assay

The FLT3 and VEGFR1/2 Kinase-Glo kinase assays were performed as reported by our earlier study [30]. The VEGFR3 activity analysis was conducted using the same protocol described in the VEGFR1/2 assay. The kinase inhibition profiling and inhibitory activity of FLT3-D835Y, CSF1R, and TRKA were determined by Invitrogen SelectScreen® kinase profiling service (Carlsbad, California, USA).

Cell viability and cell growth assays

Cell viability was assessed with an MTS assay as carried out by the method described previously [31]. Cells were seeded in 96-well plates at a density of 10,000 cells per well for 16 hours and then treated with vehicle or various concentrations of compound in medium. The change in viable cells was quantitated using the MTS method (Promega, Madison, WI, USA) according to the manufacturer’s recommended protocol or by cell counting under a microscope. The GC50 value was defined as the amount of compound that caused a 50% reduction in cell viability in comparison with DMSO-treated (vehicle) control and was calculated using Prism version 4 software (Graph-Pad Software, Inc., San Diego, CA, USA). The data are presented as the mean ± S.D. from three independent experiments. Cell growth were measured by trypan blue dye exclusion method. Cells (1×106 mL−1) were cultured in 12-well plates for 24 hours and then treated with different amounts of test compound and the same volumes of vehicle for 72 hours. The number of viable cells was counted by trypan blue dye staining using a hemocytometer at the indicated time point after drug treatment. The result was expressed as the mean ± S.D. from triplicate determinations.

Western blotting

Cells were lysed in lysis buffer (50 mM Tris, pH 8.0, 150 mM NaCl, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, 1 mM sodium orthovanadate, 1 mM PMSF, and 1 mM DTT). Protein lysates were resolved by SDS-PAGE and transferred onto a polyvinylidene difluoride (PVDF) membrane (Millipore, Bedford, MA, USA). The membranes were immunoblotted with appropriate antibodies and detected using the SuperSignal reagent (Pierce, Rockford, IL, USA) followed by exposure to X-ray film.

Apoptosis assay

The number of apoptotic cells was determined by fluorescence-activated cell sorting (FACS) analyses using Annexin V-FITC staining. In brief, cells were centrifuged after drug treatment and resuspended in 1× binding buffer containing 2.5 mM of calcium (Ca2+). Cells were incubated with Annexin V-FITC (Biolegend, San Diego, CA) and propidium iodide (PI) (Sigma–Aldrich, St. Louis, MO) in the dark at room temperature for 20 minutes. Next, samples were reconstituted with 1× binding buffer and subjected to flow cytometry FACS Calibur (BD Bioscience, Franklin Lakes, NJ) to analyze the Annexin V-positive population using CellQuest Pro (BD Bioscience) and FlowJo (Treestar, Inc., San Carlos, CA).

Pharmacokinetic analysis of BPR1J-340

The uses of animals were approved by The Institutional Care and Use Committee of the National Health Research Institutes. Briefly, male Sprague–Dawley rats weighing 300–400 g each were obtained from BioLASCO, Taiwan Co., Ltd., Ilan, Taiwan. One day prior dosing, rats were surgically prepared with a jugular-vein cannula and fasted overnight (~18–20 hr). Water was available ad libitum. Food was provided at 4 hours after dosing. Single 1.5 mg/kg intravenous dose of BPR1J-340, as a PEG400/water (80/20, v/v) solution, was separately administered to groups of 3 rats each via the jugular-vein cannula. At 0 (prior to dosing), 2, 5, 15 and 30 min. and at 1, 2, 4, 6, 8 and 24 hr after dosing, a blood sample was collected. Plasma was separated from the blood by centrifugation (14,000 g for 15 min at 4°C in a Beckman Model AllegraTM 6R centrifuge) and stored in a freezer (~20°C) until
Subcutaneously tumor xenograft experiments

Male nude mice (Nu-Fox1nu) of eight weeks old were purchased from BioLasco (Ilan, Taiwan). The nude mice (n = 5–7 per group) were inoculated subcutaneously with MOLM-13 (1 x 10^6 cells/flank). All human cancer cells were determined to be free of Mycoplasma spp before the inoculation. When tumors size reached 100–200 mm³ and a large size of >500 mm³, the animals were grouped and treated with the BPR1J-340 (5 and 20 mg/kg, i.v.) or vehicle as control once daily for 5 days per week for two or three weeks. Tumor sizes were measured and calculated with the formula of length x width²/2 after the initiation of the treatments. Tumor size and animal body weight were measured twice a week after the tumor cell inoculation. At the end of the study, animals were euthanized by carbon dioxide inhalation followed by cervical dislocation. The significant difference between the treated and vehicle control were analyzed by using one-way ANOVA and Student-Newman-Keuls test. The level of a statistical significance was set at p<0.05.

Results

In vitro kinase profiling of BPR1J-340

BPR1J-340, a urea-substituted 3-phenyl-1H-5-pyrazolylamine-based compound, was designed through chemical modification of sulfonamide series of derivatives (BPR1J-097 series) by a structure-activity relationship (SAR) study (Fig. 1) [31,32]. For kinase inhibition specificity screening, BPR1J-340 was tested against 59 protein kinases covering the major oncogenic kinases of human protein kinome. This kinase inhibition profile revealed that BPR1J-340 is a highly selective kinase inhibitor and most of the tested kinases were not significantly inhibited at the concentration of 100 nM (Table S1). Subsequently, the most potent kinases from the 59 kinases screening were further evaluated. As shown in Table 1, BPR1J-340 potently inhibited wild-type FLT3 (IC50 = 29.5 nM, as compared to ABT869 with an IC50 of 38±5 nM), mutant FLT3-D835Y (IC50 = 19 nM), CSF1R (FMS) (IC50 = 78 nM), KDR (VEGFR2) (IC50 = 28±9 nM), and FLT4 (VEGFR3) (IC50 = 29±7 nM). In addition, BPR1J-340 inhibited TRKA, which is associated with the growth and metastasis of breast cancers, with an IC50 value of 8 nM. Given the high similarity of the ATP binding pockets of FLT3 and the Aurora kinases, the inhibitory activity toward Aurora A and Aurora B were measured. The IC50s of BPR1J-340 were determined to be 1,040 nM and 1,344 nM for Aurora A kinase and Aurora B kinase, respectively. Taken together, BPR1J-340 is a selective kinase inhibitor with in vitro activity against FLT3, VEGFR2, VEGFR3 and TRKA receptor tyrosine kinases.

BPR1J-340 inhibits the proliferation of FLT3-ITD⁺ cells

After the selective and potent growth inhibitory effects of BPR1J-340 were demonstrated in FLT3-ITD-bearing leukemic cells, the cell proliferation inhibitory activity was tested in a panel of leukemic and non-leukemic cells. BPR1J-340 inhibited the cellular proliferation of FLT3-ITD⁺ cells (MOLM-13 and MV4;11 FLT3-dependent cell lines) with an IC50 of approximately 5 nM. Cells that were not dependent on FLT3 signaling for growth, including RS4;11, U937, and K562 leukemic cells [33,34,35], and the non-leukemic cell lines tested were either weakly inhibited or were not inhibited by BPR1J-340 (Table 2). The growth of the FLT3-independent RS4;11 cell line which contains wild-type FLT3 was only weakly inhibited by BPR1J-340 with an IC50 of 770±360 nM. The sensitivity toward BPR1J-340 varies between the FLT3-dependent cells MOLM-13/MV4;11 and FLT3-independent cells RS4;11, suggesting that the FLT3 signaling pathway in FLT3-ITD+ cells is almost completely disturbed by BPR1J-340. Furthermore, our results suggest that BPR1J-340 could inhibit FLT-ITD activity by in vitro biochemical and cellular assays. Overall, BPR1J-340 is a potent and highly selective inhibitor for the proliferation for FLT3-driven cells.

BPR1J-340 inhibits FLT3-STAT5 signaling

To address whether BPR1J-340 was able to inhibit the FLT3 signaling pathway in FLT3-driven cells, MV4;11 cells were treated...
signaling pathway, particularly the FLT3-ITD pathway. inhibition of cellular FLT3 phosphorylation and modulates the FLT3 phosphorylation by Western analysis. As shown in Figure 2B, 1 hr, and FLT3 ligand (50 ng/mL) was added for 5 min to cells were treated with BPR1J-340 at various concentrations for 1 hour, and the phosphorylation of all the FLT3-WT, FLT3-ITD and FLT3-D835Y to BPR1J-340, HEK293T mutants FLT3-ITD and FLT3-D835Y was inhibited by BPR1J-340, with IC50 values of 10 to 100 nM. Taken together these data demonstrate that BPR1J-340 induces apoptosis in FLT3-ITD expressing cells with different concentrations of BPR1J-340 for 1 hour, and the phosphorylation status of FLT3 and STAT5 (Signal Transducer and Activator of Transcription 5), a critical down-stream modulator that plays a major role in FLT3-ITD signal transduction for cell expansion and survival, was examined by Western blot analysis. As shown in Figure 2A, BPR1J-340 suppressed the phosphorylation of FLT3 and STAT5 in a dose-dependent manner with an IC50 value of approximately 1 nM. To investigate the difference in sensitivity between FLT3-WT and the activating mutants FLT3-ITD and FLT3-D835Y to BPR1J-340, HEK293T cells engineered to express FLT3-WT or mutants FLT3 (FLT3-ITD, FLT3-D835Y) were analyzed [31]. The HEK293T-FLT3 cells were treated with BPR1J-340 at various concentrations for 1 hr, and FLT3 ligand (50 ng/mL) was added for 5 min to prepare the cell lysate for detection of changes in FLT3 phosphorylation by Western analysis. As shown in Figure 2B, the phosphorylation of all the FLT3-WT, FLT3-ITD and FLT3-D835Y was inhibited by BPR1J-340, with IC50 values of 10 to 100 nM. Taken together these data demonstrate that BPR1J-340 inhibits cellular FLT3 phosphorylation and modulates the FLT3 signaling pathway, particularly the FLT3-ITD pathway.

BPR1J-340 induces apoptosis in FLT3-ITD expressing cells

As the inhibition of FLT3 signaling results in a loss of growth potential and an induction of apoptosis in FLT3-ITD expressing cells, the effects of BPR1J-340 on cell apoptotic response in FLT3-ITD+ cells was investigated. The MOLM-13 and MV4;11 cells were treated with BPR1J-340 at different concentrations for 24 hours and analyzed for the induction of apoptosis using active caspase-3 and cleaved PARP (poly-ADP-ribose polymerase) analysis. The ability of BPR1J-340 to induce apoptosis is evident as shown in Figure 3 where significant cleavage of caspase-3 (active caspase-3) and PARP (active PARP) was observed in MOLM-13 and MV4;11 cells treated with BPR1J-340 at 10 nM.

SAHA in combination with BPR1J-340 enhances cytotoxicity against FLT3-ITD+ expressing cells

Some histone deacetylase inhibitors (HDACi) will enhance the cytotoxic activity of FLT3 inhibitor on FLT3-ITD+ cell via degradation of FLT3-ITD and STAT5 [24,26,36,37]. To determine whether SAHA, an HDACi, can synergize with BPR1J-340 to enhance cytotoxicity in FLT3-ITD expressing cells by interfering with the FLT3-ITD and STAT5 axis, the combined effects of SAHA and BPR1J-340 on cytotoxicity were examined. The cell growth rate of MOLM-13 and MV4;11 with SAHA and BPR1J-340 combinational treatment was decreased compared with the single-drug treatment (Fig. 4A). Next, we examined the SAHA treatment effect on BPR1J-340-induced apoptosis in FLT3-ITD-expressing cells. Sole treatment with SAHA (at 300 nM) for 48 hours did not increase the rate of apoptotic cells in control population 10%, whereas SAHA/BPR1J-340 co-treatment resulted in a greater induction of apoptosis (increased 20% in MOLM-13 and 12% in MV4;11 cells, respectively) as compared with BPR1J-340 drug treatment alone (Fig. 4B and Fig. 4C). These data suggested that SAHA enhance BPR1J-340-induced apoptosis in FLT3-ITD+ cells. We further elucidated the protein levels of FLT3-ITD and STAT5 in this SAHA/BPR1J-340 co-treatment enhanced cytotoxicity. We treated MOLM-13 cells with SAHA, BPR1J-340, or their combination for 20 hours. Compared with single-drug treatment, the combination of SAHA and BPR1J-340 remarkably decreased the protein levels of FLT3-ITD and STAT5 (Fig. 4D). Furthermore, BPR1J-340/SAHA treatment profoundly reduced Mcl-1 (myeloid cell leukemia-1, an anti-apoptotic member of the BCL-2 family) protein levels in FLT3-ITD+ cells (Fig. 4D). This results of Mcl-1 reduction may explain the enhanced cytotoxicity even the STAT5 phosphorylation at Tyr694 was completely block by BPR1J-340 (Fig. 4D) Mcl-1 plays an essential role in resistance to chemotherapy in AML cells [38]. Thus, targeting MCL-1 using SAHA/BPR1J-340 may be a promising strategy to overcome drug resistance in FLT3-ITD-positive AML. Our results suggest that the reduction in the total protein levels of FLT3-ITD, STAT5 and Mcl-1 by SAHA addition provides a possible mechanism to explain the enhanced cytotoxicity with SAHA/BPR1J-340 coadministration.

Pharmacokinetic parameters of BPR1J-340

To evaluate the pharmacokinetic properties of BPR1J-340, we measured the plasma concentration of BPR1J-340 over a 24-hr period after a single intravenous administration (Fig. 5 and Table 3). After a 1.5 mg/kg intravenous dose administered to Sprague-Dawley rats, BPR1J-340 achieved a maximum plasma concentration of 7.9 μM (4,296 ng/mL) at 2 min after dosing. The plasma concentration exceeded 80 ng/mL for 6 hours and remained steady at 9.9 ng/mL (C24 hour, 18 nM) after 24 hour dosing. Therefore, the plasma concentration of BPR1J-340 reaches sufficient levels to inhibit proliferation and induce apoptosis of FLT3-ITD+ cells based on cellular experiments. The total body clearance was 20.4±5.2 mL/min/kg and the volume of distribution at the steady state (Vss) was 10.3±2.5 L/kg for BPR1J-340 in rats. The apparent plasma half-life was approximately 8.8 hr.

Table 2. Anti-proliferative activity of BPR1J-340 against a panel of tumor cell lines.

| Cell lines   | Proliferation GI50, nM |
|-------------|------------------------|
| Leukemias   |                        |
| MOLM-13     | 3.4 ± 1.5              |
| MV4;11      | 2.8 ± 1.2              |
| RS4;11      | 770 ± 360              |
| U937        | 2,400 ± 780            |
| K562        | 6,370 ± 2,070          |
| Non-Leukemias |                    |
| HCC827      | 2,470 ± 380            |
| H1975       | 2,610 ± 270            |
| H1650       | 3,725 ± 977            |
| H2228       | 2,520 ± 420            |
| NCIH460     | 13,160 ± 2,840         |
| A431        | 6,570 ± 680            |
| HCT-116     | 9,330 ± 2,320          |
| MKN45       | 3,580 ± 1,210          |
| MiaPaCa-2   | 9,960 ± 570            |
| RT4         | 7,030 ± 1,510          |
| MCF-7       | 11,300 ± 1,110         |
| Huh7        | 9,740 ± 1,680          |
| Hep3B       | 8,310 ± 490            |
| Detroit551  | 8,340 ± 680            |

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with substantial volumes. The plasma concentration of BPR1J-340 remained steady at 9.9 ng/mL (C24 hour, 18 nM) after 24 hour dosing. Therefore, the plasma concentration of BPR1J-340 reaches sufficient levels to inhibit proliferation and induce apoptosis of FLT3-ITD+ cells based on cellular experiments. The total body clearance was 20.4±5.2 mL/min/kg and the volume of distribution at the steady state (Vss) was 10.3±2.5 L/kg for BPR1J-340 in rats. The apparent plasma half-life was approximately 8.8 hr.
Tumor growth suppressive activity of BPR1J-340

FLT3-ITD\(^+\) MV4;11 and MOLM-13 xenografts have been widely used to evaluate the \textit{in vivo} efficacy of FLT3 inhibitors against AML [39,40]. We and other groups have found the MV4;11 xenograft model is noticeably more sensitive to FLT3 inhibitors [30,31,39,41]. To assess the ability of BPR1J-340 to inhibit tumor growth in an AML xenograft model, we therefore selected the MOLM-13 tumor xenograft model. BPR1J-340 was administered intravenously (5 mg/kg qd) on days 1–5 of each week for 3 weeks; by the end of this period the MOLM-13 tumors had reached an average volume of 200 mm\(^3\). Treatment via this regimen resulted in complete regression of the tumor in all animals by day 22; complete regression (CR) occurred in 4 of 6 treated animals within 31 days of post-treatment observation (Fig. 6A). In this efficacy study, mice with larger tumors (average tumor volume 630 mm\(^3\), \(n = 3\)) were given a single intravenous dose of 20 mg/kg BPR1J-340 on days 1–5 of each week for 2 weeks, which resulted in complete and durable tumor regression with no palpable tumors detected during the 48-day follow up (Fig. 6B). No loss of body weight or mortality was observed in animals treated with any dose.

Figure 2. BPR1J-340 inhibits FLT3-dependent signaling. (A) MV4;11 cells were treated with BPR1J-340 at the indicated concentrations for 1 hour. The phosphorylation status of FLT3 and STAT5 were evaluated by Western blot analysis. (B) HEK 293T cells were transfected with FLT3-WT, FLT3-ITD, or FLT3-D835Y-expressing plasmids for 24 hours and then incubated with various concentrations of BPR1J-340 for 1 hour. The FLT3 phosphorylation status in the transfected cells was evaluated by Western blot analysis.
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Figure 3. BPR1J-340 induces apoptosis in MOLM-13 and MV4;11 cells. Western blotting revealed that BPR1J-340 was able to induce apoptosis in FLT3-ITD-driven FLT3-ITD\(^+\) AML cells. MOLM-13 (A) and MV4;11 (B) cells were treated with BPR1J-340 at the indicated concentrations for 24 hours, and the cell lysates were then subjected to Western blot analysis using antibodies against caspase-3 and PARP (poly-ADP-ribose polymerase). (full length caspase-3 (FL-caspase-3), cleavage of caspase-3 (CL-caspase-3), full length poly(ADP-ribose) polymerase (FL-PARP), cleavage of PARP (CL-PARP)).
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of BPR1J-340 (data not shown). These results demonstrate that BPR1J-340 effectively reduces tumor size at intravenous doses of 5 and 20 mg/kg/day; BPR1J-340 is well-tolerated in mice at these effective doses.

Discussion

Given that the abnormal expression of FLT3 kinase, including amplified or aberrantly activated FLT3, is frequently observed in

Table 3. Pharmacokinetic parameters of BPR1J-340 in rats.

| Compound  | T1/2 (hr) | CL (mL/min/kg) | Vss (L/kg) | AUC(0–24) (ng/mL-hr) |
|-----------|----------|----------------|------------|----------------------|
| BPR1J-340 | 8.8±0.6 | 20.4±5.2        | 10.3±2.5   | 1130±260             |

Figure 4. The effect of BPR1J-340 and SAHA versus drug combination on AML FLT3-ITD+ cells. MOLM-13 or MV4;11 cells were seeded at an initial concentration of $1 \times 10^5$ cell mL$^{-1}$ for 24 hours and then treated with BPR-1J340 either alone or in combination with SAHA in the indicated concentrations. (A) Viable cells were counted after staining with trypan blue dye at indicated time point (B) At 48-hours after treatment, cells were stained with FITC-labeled annexin V and propidium iodide and percentages of apoptotic cells were analyzed by flow cytometry. (C) The representative flow cytometry histograms of assays of annexin positive apoptotic cells at 48 hours drug treatment (D) MOLM-13 cells were treated with drugs for 20 hours, and then followed by Western blot analysis to assess changes in protein levels with the indicated antibodies. Statistical analysis was performed using Student’s t-test. *Indicates significant difference compared to vehicle control (* p<0.05, ** p<0.01). Data shown are representative of multiple independent experiments.

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Figure 5. Pharmacokinetic profile of BPR1J-340 in rats. A single intravenous bolus dose (1.5 mg/kg) of BPR1J-340 was administered to adult male Sprague-Dawley rats (n = 3). Data illustrate mean values (n = 3) ± S.D. of plasma concentrations of BPR1J-340 at each timepoint.

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inhibitors that not only increased the potency of FLT3 inhibitors, we intended to search other series of such as BPR1J-097 [31]. In continuing to our efforts to develop pyrazolylamine-based compounds as potent inhibitors of FLT3.

Previously, we identified a sulfonamide series of 3-phenyl-1H-5-pyrazolylamine-based compounds as potent inhibitors of FLT3 such as BPR1J-097 [31]. In continuing to our efforts to develop potent FLT3 inhibitors, we intended to search other series of inhibitors that not only increased the in vitro growth-inhibitory effect on AML cells but also prolonged the duration of action in vivo. Through rational design, we discovered BPR1J-340, which is a urea series of 3-phenyl-1H-5-pyrazolylamine-based FLT3 inhibitor, with effectively inhibits FLT3-WT or FLT3-ITD activity in vitro and in vivo. Because multiple signaling pathways influence the growth and metastatic potential of tumor cells, many of the inhibitors in clinical development are designed as multi-targeted inhibitors that block a limited number of oncogenic kinases. Thus, the kinase selectivity profiling of BPR1J-340 was performed to identify additional targets in a panel of 59 tested oncogenic kinases. In further biochemical assay, BPR1J-340 demonstrated potent inhibition (20–190 nM) against the angiogenic kinases VEGFR1, VEGFR2, and VEGFR3, which all play an important role in the tumor microenvironment (Table 1). In addition, BPR1J-340 potently inhibited TRKA activity with an IC_{50} value of 8 nM. Taken together, BPR1J-340 is characterized as a selective multi-targeted inhibitor with potent inhibition activity against FLT3-WT, FLT3-ITD, VEGFR2, VEGFR3, and TRKA. This inhibition profile may allow BPR1J-340 to inhibit tumor growth directly by blocking the aberrant FLT3 signaling pathway and indirectly by targeting tumor angiogenesis. BPR1J-340 may also have clinical potential in tumor driven by abnormally expressed TRKA receptors, which can occur in brain, prostate, pancreatic, and breast cancer [42,43,44,45].

BPR1J-340 inhibited cellular FLT3 phosphorylation and modulated the FLT3 signaling pathway, which resulted in inhibition of proliferation and induction of apoptosis. BPR1J-340 demonstrated potent growth inhibition, predominantly in FLT3-dependent cells (MOLM-13 and MV-4;11) but not in FLT3-independent cells. BPR1J-340 inhibited the proliferation of mutant FLT3-ITD cells with an GC_{50} of 2–6 nM and inhibited FLT3-ITD phosphorylation with an IC_{50} of 10 nM; even the cells transfected with the FLT3-D835V mutant were also inhibited by BPR1J-340 with an IC_{50} of approximately 100 nM. Consistent with these results, BPR1J-340 effectively induced apoptosis in FLT3-ITD cells.

HDAC inhibitors may exhibit growth inhibition activity against AML cells and significantly improve the therapeutic efficacy of FLT3 inhibitors [24,46]. A recent study reported that HDACi LBH589 plus an FLT3 inhibitor (PPKC412, AC220) combination treatment could synergistically induce apoptosis through FLT3-ITD and STAT5 degradation [26]. It also demonstrated that activated caspase-3 (CL-caspase 3) contributes to the degredation of FLT3-ITD and STAT5 [26]. FLT3-ITD degradation was also reported secondary to HDACi-induced up-regulation of UCHL8 and down-regulation of HSP90 [36,47]. McI-1, an anti-apoptotic protein that promotes survival of FLT3-ITD+ cells via STAT5 activation, is down-regulated by FLT3 inhibition [25]. The HDACi SAHA also induced down-regulation of McI-1 [48]. Furthermore, McI-1 protein is a direct cleavage substrate of activated caspase-3 [49]. We noted that the amount of McI-1 correlated with induction of activated caspase-3 (Figure 4D). Our results demonstrate that SAHA enhances BPR1J-340 inhibition activity in FLT3-ITD+ cells and suggests that the enhancement is...
These data demonstrated that BPR1J-340 is superior to the inhibition of FLT3 activity in rats or mice. BPR1J-340 dosing once a day is sufficient for continuous inhibition of FLT3-ITD. In addition, the high Vss indicated that the distribution of BPR1J-340 were close to the IC50 value for inhibition of FLT3-ITD. Even at 24 hour after the single dosing, the plasma levels of BPR1J-340 into deep tissue compartments, including tumor tissue, is expected. These pharmacokinetic properties suggest that BPR1J-340 into deep tissue compartments, including tumor tissue, is expected. These pharmacokinetic properties suggest that BPR1J-340 dosing once a day is sufficient for continuous inhibition of FLT3 activity in rats or mice.

To examine whether BPR1J-340 exhibits antitumor activity in vivo, MOLM-13 cells were subcutaneously implanted into nude mice. Our results demonstrated that BPR1J-340 administration resulted in significant tumor regression and tumor shrinkage in this MOLM-13 tumor model. In comparison with sulfonamide BPR1J-97 [I/8 CR, 25 mg/kg] in the same model [31], BPR1J-340 results in a higher CR ratio (67%) at a lower dose (5 mg/kg). These data demonstrated that BPR1J-340 is superior to the sulfonamide compound BPR1J-097 in an in vivo efficacy study. In conclusion, results from this study demonstrate that BPR1J-340 exhibits high potency and excellent selectivity against FLT3 kinase, strong suppression of the FLT3-ITD survival signaling pathway, favorable pharmacokinetic properties, and complete tumor regression in a FLT3-ITD+ xenograft model.

Supporting Information

Table S1 Kinase inhibition profiles of BPR1J-340. Percentage inhibition was determined by Invitrogen SelectScreen kinase profiling service.

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Author Contributions

Conceived and designed the experiments: WHL WTJ TKY CTC JT-AH. Performed the experiments: WHL KJY Chun-Hwa Chen CTH SYH LHC CPG Chun-Hsien Chiu, LCK. Analyzed the data: WHL CPG TKY CTC JT-AH. Contributed reagents/materials/analysis tools: WTJ YSC CTC JT-AH. Wrote the paper: WHL WTJ TKY CTC JT-AH.

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