Cotylenin A and tyrosine kinase inhibitors synergistically inhibit the growth of chronic myeloid leukemia cells

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Abstract. The treatment of chronic myeloid leukemia (CML) with tyrosine kinase inhibitors (TKIs) has substantially extended patient survival. However, TKIs do not effectively eliminate CML stem cells. In fact, CML stem cells persist and cause relapse in the majority of patients upon discontinuation of the drug treatment. Transcriptomic and proteomic analyses have revealed that p53 and c-Myc play defining roles in CML stem cell survival, suggesting that the dual targeting of p53 and c-Myc may selectively eliminate stem cells in patients with CML. Since the downregulation of c-Myc and then upregulation of p21 (a target gene of p53) are commonly observed during the differentiation of acute myeloid leukemia cells induced by differentiation inducers, we hypothesized that differentiation-inducing agents may be useful in regulating c-Myc and p53 expression in CML cells. In the present study, we demonstrate that some differentiation-inducing agents effectively suppress the self-renewal ability of CML cells, and that the combination of these inducers with TKIs results in significantly greater inhibitory effects on CML cell growth compared to the use of TKIs or the inducer alone. The KU812 cells were treated with various concentrations of the inducers in the presence or absence of 30 nM imatinib for 4 days. Among the differentiation inducers we tested, cotylenin A (CN-A) was the most potent at inhibiting the self-renewal ability of the CML cells. CN-A induced the robust expression of CD38, a marker of committed progenitor and more differentiated myelomonocytic cells, and rapidly suppressed c-Myc expression and upregulated p21 expression in CML cells. Thus, these results suggest that CN-A may have potential to promote the elimination of stem cells in CML.

Introduction

Chronic myeloid leukemia (CML), a clonal disease affecting hematopoietic stem cells, is driven by the BCR/ABL oncoprotein, a constitutively active tyrosine kinase. Patients with CML in the chronic phase are treated with imatinib (IM) or other tyrosine kinase inhibitors (TKIs), which are highly effective at inducing remission and prolonging survival. However, TKIs do not completely eliminate leukemia stem cells (LSCs), even in patients who achieve deep molecular responses (1-3). Thus, the identification of drugs that can target these LSCs is of primary importance in order to achieve the eradication of CML. Transcriptomic and proteomic analyses have revealed that p53 and c-Myc play defining roles in CML-LSC survival (4). These results suggest that the dual targeting of p53 and c-Myc may selectively eliminate LSCs in patients with CML.

Human and murine myeloid leukemia cells can be induced to differentiate into mature granulocytes and macrophages by various differentiation-inducing agents (5,6). There are a number of methods with which to induce granulocytic or monocytic differentiation in leukemia cells, since differentiation-inducing agents act by different mechanisms to induce the production of more or less identical end-stage cells (7-9). However, the downregulation of c-Myc and the upregulation of p21 (a target gene of p53) are commonly observed during the differentiation of myeloid leukemia cells induced by various inducers of differentiation (10-12). These results suggest that the dual targeting of p53 and c-Myc may selectively eliminate LSCs in patients with CML.

In this study, we examined the effects of agents that induce the differentiation of acute myeloid leukemia (AML) cells on the proliferation of CML cells in the presence of TKIs. Among the former, cotylenin A (CN-A) was found to be the most effective at inhibiting the clonogenic potential and proliferation of CML cells. CN-A, which is a novel fusicoccane-diterpene glycoside with a complex sugar moiety, has been shown to affect the differentiation of leukemia cells that have been freshly isolated from patients with AML in primary culture (6,13). The administration of CN-A has been shown to significantly prolonged the survival of mice inoculated with retinoid-resistant human promyelocytic leukemia NB4 cells, and no appreciable adverse effects were observed (14). These findings thus suggest that CN-A may be useful in CML therapy when combined with TKIs.

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Materials and methods

Materials. RPMI-1640 medium, all-trans retinoic acid (ATRA), doxorubicin, rapamycin and cytosine arabinoside (AraC) were purchased from Sigma-Aldrich Japan (Tokyo, Japan). Dimethyl sulfoxide (DMSO), 1α,25-dihydroxy vitamin D3 (VD3) and sodium butyrate were purchased from Wako Chemicals (Osaka, Japan). CN-A was a gift from Professor Takeshi Sassa, phycocerythrin (PE)-labeled anti-CD38 antibody (cat. no. 555460) was obtained from BD Immunocytometry Systems (San Jose, CA, USA). IM, dasatinib (DAS) and nilotinib (NIL) were obtained from Selleck Chemicals (Houston, TX, USA).

Patients with leukemia. Leukemic bone marrow specimens were collected at diagnosis, after the patients provided written informed consent for sample collection in accordance with institutional policy. The Shimane University Institutional Committee on Ethics (Shimane, Japan) approved the present study. The samples were obtained from the following patients: Case 1 was a 71-year-old Japanese male who was admitted to the Department of Oncology/Hematology, Shimane University Hospital (Shimane, Japan), presenting with hyperleukocytosis. Laboratory data upon admission were WBC, 60,420/µl; myelocytes, 12.7%; metamyelocytes, 15.5%; and lactate dehydrogenase (LDH), 861 U/l. Samples from case 1 were obtained in November, 2017. Case 2 was a 64-year-old Japanese female who was admitted to the Department of Oncology/Hematology, Shimane University Hospital, presenting with hyperleukocytosis. Laboratory data upon admission were WBC, 60,420/µl; myelocytes, 12.7%; metamyelocytes, 4.2%; and LDH, 674 U/l. Samples from case 2 were collected in December, 2017. In both cases, CML upon admission were WBC, 169,040/µl; myelocytes, 13.5%; metamyelocytes, 15.5%; and lactate dehydrogenase (LDH), 861 U/l. Samples from case 1 were obtained in November, 2017. Case 2 was diagnosed on the basis of bone marrow morphology and cytogenetic translocation, t(9;22), in 98% of the cells by fluorescence in situ hybridization (FISH) analysis.

Cells and cell culture. K562 (15) and KU812 (16) cells were obtained from the respective founders and were maintained in RPMI-1640 medium supplemented with 10% fetal bovine serum (Biowest, Nuaillé, France) and 80 µg/ml gentamicin (MSD Co., Ltd., Tokyo, Japan) at 37˚C in a humidified atmosphere of 5% CO2 in air.

Assay of cell growth and properties of differentiated cells. Suspensions of cells (2x10^4 cells/ml) in 1 ml of culture medium were incubated with or without the test compounds in multidishes. Cell numbers were counted using a Model Z1 Particle Counter (Beckman Coulter, Tokyo, Japan). The surface expression of CD38 was determined by FACScan flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA). The cell number to one-half of that in untreated cells (IC50) was calculated from the cell counts and the dilution used when feeding the culture. Cell numbers were counted using a Beckman Coulter Z1 Particle Counter.

Colonial formation assay. The K562 or KU812 cells (3x10^5 per dish) were plated into 2 ml of a semi-solid medium with 0.8% methylcellulose and 20% fetal bovine serum in triplicate multiwell plates (12 wells, 3.5 cm^2 growth area/well) for 7-14 days. A solution of 0.1 ml of PBS containing various concentrations of the drugs was added to the semi-solid medium. To determine the colony-forming ability of the leukemia cells from patients with CML, heparinized bone marrow aspirations were diluted with RPMI-1640 medium supplemented with 10% fetal bovine serum, overlaid on 15 ml of Ficoll-Paque Plus (GE Healthcare Biosciences, Uppsala, Sweden) and centrifuged at 500 x g for 30 min. The mononuclear cells were washed twice and suspended in RPMI-1640 medium supplemented with 10% fetal bovine serum, plated in semi-solid culture medium with 20% serum at 10^5 cells/dish for colony formation, and incubated at 37˚C in a humidified atmosphere of 5% CO2 in air. Colonies were photographed under an inverted microscope (Model CKX41; Olympus, Tokyo, Japan). In serial colony formation assays, the cells were serially replated after 7 days of culture.

Western blot analysis. The cells were packed after being washed with cold PBS, and then lysed at 1.5x10^7 cells/ml in sample buffer [63 mM Tris-HCl (pH 6.8), 15% glycerol, 2% sodium dodecyl sulfate (SDS), 5% 2-mercaptoethanol and 0.005% bromophenol blue]. The resultant lysates were resolved on 10% SDS-polyacrylamide gels. Protein concentration was quantified using the Protein Quantification kit-Rapid (Wako Pure Chemical Industries, Ltd.). Equal amounts of protein (10 µg) were separated by SDS/PAGE (10% gels) prior to transfer to polyvinylidene fluoride membranes (Bio-Rad Laboratories, Hercules, CA, USA), and then blocked with Block Ace (DS Pharma Biomedical Co., Ltd., Osaka, Japan) for 60 min at room temperature. The membranes were then immunoblotted with anti-p21 (#2947), anti-cMyc (#5606) and anti-β-actin (#4970) antibodies (1:500 dilution) antibodies which were purchased from Cell Signaling Technology Japan (Tokyo, Japan). Horseradish peroxidase (HRP)-conjugated antibody (#7074S and 7076S; Cell Signaling Technology, Danvers, MA, USA) was used as a secondary antibody (1:2000 dilution). The bands were developed by treatment with the Immun-Star HRP Chemiluminescent kit (Bio-Rad Laboratories) for 5 min at room temperature, and detected using a Fuji Lumino Image Analyzer LAS-4000 system (Fuji Film, Tokyo, Japan).

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR). Total RNA was extracted from the cells using TRIzol reagent (Sigma-Aldrich). Total RNA was converted to first-strand cDNA primed with random hexamer in a reaction volume of 20 µl using an RNA PCR kit (qPCR RT Master Mix; Toyobo Co., Ltd., Osaka, Japan), and 2 µl of this reaction was used as a template in real-time PCR. The primers
were used as previously described (17). The quantitative PCR reaction was performed using a Takara TP860 Real-Time PCR system (Takara Bio, Tokyo, Japan) according to the manufacturer's instructions (40 cycles of duration at 95°C for 15 sec and annealing and extension at 60°C for 60 sec). The threshold cycle values were normalized to the threshold value of glyceraldehyde-3-phosphate dehydrogenase. Data analysis was performed using the 2^ΔΔCq method (18).

Statistical analysis. The results are expressed as the means ± standard deviation (SD). Statistical analysis was conducted with SPSS 19.0 software (IBM Japan, Tokyo, Japan). Statistical significance between multiple groups was determined by ANOVA with a Bonferroni post-hoc test, and between 2 groups using a Student's t-test. Significant differences were considered to exist for probabilities below 5% (P<0.05).

Results

Combined effects of IM and differentiation-inducing agents on the proliferation of CML cells. IM inhibited the growth of the K562 and KU812 cells in a concentration dependent manner; the IC50 values at 7 days were 25 and 18 nM, respectively (data not shown). The KU812 cells were treated with 30, 60 and 100 nM IM for 14 days, washed with PBS, resuspended in drug-free culture medium, and cultured for an additional 7 days. The cells were first treated with 100 nM IM for 14 days. After washing, the cells were further treated without or with IM. While continuous IM treatment effectively suppressed proliferation, the depletion of IM from the medium allowed the cells to re-grow, indicating that IM did not completely block the ability of the cells to repopulate (Fig. 1A). Similar results were obtained when the K562 cells were treated with 100 nM IM (data not shown). We then examined the effects of several agents that are known to induce the differentiation of AML cells on the proliferation of CML cells (Table I). KU812 cells were treated with various concentrations of the inducers in the presence or absence of 30 nM IM. Most of the compounds we used effectively synergized with IM to inhibit the growth of KU812 cells. Low concentrations of ATRA and IM synergistically inhibited the proliferation of KU812 cells, but ATRA did not affect the proliferation of K562 cells even in the presence of IM. Combination of butyrate and IM was not evident, suggesting that butyrate is not a good partner of IM. On the other hand, CN-A alone greatly inhibited the long-term proliferation of KU812 cells. In this condition, therefore, the combined effects with IM were modest. These results suggest that CN-A at concentrations less than IC50 are enough to inhibit the long-term proliferation of CML cells in the presence of IM. The results of treatment with IC50 concentrations for 4 days are shown in Table I. CN-A was less cytotoxic than the other compounds tested, but was the most potent at inhibiting cell proliferation for 14 days (Table I), suggesting that CN-A effectively suppressed the self-renewal ability of these cells. CN-A and IM cooperatively inhibited cell proliferation (Fig. 1B). These compounds also exerted combined effects on the K562 cells. While the K562 cells that were treated with 2 µg/ml of CN-A or 30 nM IM for 14 days still grew in the drug-free culture, those treated with CN-A plus IM lost the ability to proliferate (Fig. 1C). Although CN-A alone hardly affected the apoptosis of the CML cells, it markedly enhanced the IM-induced apoptosis of the K562 cells (data not shown). These compounds also exerted combined effects on the clonogenic activity of the cells (Fig. 1D). The colony-forming ability of the K562 cells was markedly inhibited by treatment with CN-A alone at a concentration of >4 µg/ml, and IM cooperatively inhibited the colony-forming ability of the K562 cells treated with CN-A. Treatment with CN-A at 1 µg/ml completely suppressed the colony-forming ability of the KU812 cells and IM did not alter this effect of CN-A on colony formation (data not shown).

The effectiveness of combined treatment with CN-A and IM on primary CML cells was examined and compared with that of treatment with either IM or CN-A alone. CN-A significantly inhibited colony formation by leukemia cells from a patient with CML (case 1) (Fig. 2A). Although IM alone did not significantly suppress colony formation, it significantly enhanced the inhibitory effects of CN-A on colony formation (Fig. 2A and B). The clonogenic activity of the leukemia cells was completely blocked by treatment with CN-A alone or in combination with IM (Fig. 2B).

Table I. Effects of differentiation inducers on the growth of KU812 cells in the presence or absence of IM.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Concentration</th>
<th>None</th>
<th>+30 nM IM</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATRA 3 nM</td>
<td>412,000±26,000</td>
<td>84,000±7,000</td>
<td></td>
</tr>
<tr>
<td>AraC 1.7 nM</td>
<td>216,400±18,200</td>
<td>76,000±6,200</td>
<td></td>
</tr>
<tr>
<td>CN-A 1.2 µg/ml</td>
<td>54,000±4,200</td>
<td>42,000±3,400</td>
<td></td>
</tr>
<tr>
<td>VD3 300 nM</td>
<td>590,000±62,000</td>
<td>186,400±12,000</td>
<td></td>
</tr>
<tr>
<td>Butyrate 730 µM</td>
<td>238,000±19,200</td>
<td>178,000±12,600</td>
<td></td>
</tr>
<tr>
<td>DMSO 105 mM</td>
<td>265,000±22,000</td>
<td>138,600±11,600</td>
<td></td>
</tr>
<tr>
<td>Rapamycin 0.13 nM</td>
<td>1,269,000±104,000</td>
<td>138,800±14,200</td>
<td></td>
</tr>
<tr>
<td>Doxorubicin 12.7 nM</td>
<td>233,600±20,200</td>
<td>167,200±13,600</td>
<td></td>
</tr>
<tr>
<td>AraC 1.7 nM</td>
<td>216,400±18,200</td>
<td>76,000±6,200</td>
<td></td>
</tr>
</tbody>
</table>

aIC50 concentrations for 4 days. bCells were cultured with various differentiation-inducing agents in the presence or absence of 30 nM IM for 14 days. Results are presented as the means ± SD of 3 separate experiments.
The proliferation of KU812 cells was also inhibited by treatment with IM plus CN-A, although the combined effects on the cells from case 2 were less prominent than those on the cells from case 1. The inhibitory effects of CN-A on the stem cell potential of primary cells from case 2 were confirmed using serial colony formation assays (Fig. 2C). CN-A, alone or in combination with IM, completely reduced the replating efficiency of the cells from case 2. By contrast, treatment with 30 nM IM did not reduce colony formation in secondary cultures from case 2.

Two second-generation TKIs have been developed and represent viable alternatives to IM. Thus, we examined the combined effects of CN-A and these TKIs. Similar results were observed when the cells were treated with CN-A plus other TKIs, such as DAS and NIL (Fig. 2D and E).

**Modulation of the phenotypes of KU812 cells by CN-A.** Since CN-A is a potent inducer of the differentiation of AML cells (6,13), in this study, we examined the effects of CN-A on the differentiation-associated phenotypes of the K562 and KU812 cells. CN-A induced morphological changes in these cells, whereas ATRA, AraC and IM alone did not (Fig. 3A). The enlargement of the cytoplasm, decreased cytoplasmic basophilia and compact nuclei were observed in the CN-A-treated cells. In the KU812 cells, morphological changes induced by CN-A were not affected by IM. CN-A did not induce CD11b expression, α-naphthyl acetate esterase activity or the reduction of nitroblue tetrazolium (markers of myelomonocytic differentiation) in either cell line even in the presence of IM (data not shown), although CN-A has been shown to effectively induce these differentiation-associated phenotypes in AML cell lines and AML cells in primary culture (13). Similar results were obtained in the CN-A-treated K562 cells (data not shown).

CD38 is an important marker of human hematopoietic stem cells. CD34+CD38− bone marrow cells are highly enriched for long-term repopulating hematopoietic stem cells, while CD34+CD38+ cells are more committed progenitor...
cells. These same markers also apply to CML cells (19). Therefore, in this study, we examined the expression of CD38 in the CN-A-treated cells. CN-A at concentrations as low as 0.5 µg/ml efficiently converted the KU812 cells from CD38- to CD38+ (Fig. 3B), consistent with its role in promoting myeloid differentiation to an intermediate stage, but not to mature stages. IM did not affect CD38 expression even in the presence of CN-A.

Effect of CN-A on the expression of c-Myc and p21 in KU812 cells. A previous study indicated that the dual targeting of p53 and c-Myc selectively eliminated CML stem cells (4). Since p21 is a target gene of p53, we examined p21 expression as a marker of the p53 signal transduction pathway. CN-A effectively decreased the c-Myc protein level and increased the p21 protein level in the KU812 cells, while ATRA did not affect the c-Myc protein level and decreased the p21 protein level (Fig. 4A). IM did not affect the protein levels of p21 or c-Myc in the KU812 cells, even in the presence of CN-A (Fig. 4B). We then examined the effect of CN-A on c-Myc mRNA expression in the KU812 cells. The downregulation of c-Myc mRNA by CN-A was observed within 6 h (Fig. 4C), suggesting that it is an early event in the action of CN-A. Neither IM nor ATRA affected mRNA expression (Fig. 4D).

Discussion

In the present study, the combination of TKIs and CN-A was found to be more effective than either drug used alone at reducing the CML cell bulk, and TKIs did not interfere
with the detrimental effect of CN-A on stem cell potential or clonogenic activity.

Differentiation-inducing agents can alter the phenotype of cancer cells. A previous study indicated that ATRA converted KCL-22 CML cells from CD38− to CD38+ cells (20), and the results of the present study demonstrated that ATRA effectively inhibited the growth of KU812 cells (Table I). However, ATRA did not affect the growth of the K562 cells and did not effectively convert the KU812 cells from CD38− to CD38+ cells. CN-A altered the phenotype of CML cells more effectively than ATRA. In a previous study, CN-A affected the differentiation of leukemic cells that had been freshly isolated from patients with AML in primary culture. It significantly stimulated both the functional and morphological differentiation of leukemia cells in 9 of 12 cases. Its differentiation-inducing activity was shown to be more potent than those of ATRA and VD3 (13). However, CN-A did not induce CD11b or NBT reduction in CML cells, although it induced CD38 expression. These results indicated that CN-A induces the differentiation of CML cells to an intermediate stage. Previous studies have indicated that CN-A and related compounds inhibit clonogenic activity, cell-surface expression of cancer stem cell markers and stemness-associated gene expression in some malignant cells (21-23). CN-A and related compounds have been shown to significantly inhibit the growth of malignant cells as xenografts without apparent adverse effects (22,24). Combined treatment with CN-A and TKIs had a selective cytotoxic effect on more primitive LSCs at clinically tolerable doses.

A receptor of fusicoccin, which is closely related to CN-A, has been reported to be a member of a family of 14-3-3 proteins that are commonly found in a wide variety of signaling and regulatory pathways (25). The 14-3-3 proteins bind to discrete phosphoserine-containing motifs present in many signaling molecules. The 14-3-3 proteins are associated with dynamic nucleo-cytoplasmic shuttling. A number of nuclear proteins can become phosphorylated, bind to 14-3-3 proteins and accumulate in the cytoplasm. The 14-3-3 proteins negatively regulate histone deacetylase 4 (26) and cyclin-dependent kinase inhibitor p27 (27) by preventing their
nuclear localization. A special subfamily of 14-3-3 proteins may bind CN-A and affect its interaction with some signaling molecules. This modification may lead to downregulation of the c-Myc gene (Fig. 4), which is an early gene that is suppressed by CN-A in CML cells. c-Myc upregulates BCR/ABL expression and is necessary for BCR/ABL-induced transformation and LSC maintenance (28). However, further studies are warranted in order to elucidate the mechanism through which the downregulation of c-Myc is related to the effects of CN-A on the 14-3-3 signaling pathway.

Discontinuation trials with second generation TKIs have shown that 40-50% of patients with CML maintain the treatment response without continued therapy. However, cardiopulmonary adverse effects are now impacting treatment choice (29,30). Therefore, more effective TKI therapy combined with targeted therapy against pathways involved in CML-LSC survival should be explored. CML-LSCs have been extensively characterized, with the aim of developing novel curative approaches based on the eradication of LSCs. A number of potential molecular targets have been identified to eradicate LSCs in CML. Several studies have presented other possible targets to eliminate LSCs in CML therapy in combination with TKIs. These include interleukin-1 signaling, hypoxia-inducible factor and mitochondrial oxidative phosphorylation (31-34). At present, however, no drug is available for clinical use against these targets. A number of pathways and mechanisms may promote the survival of LSCs in CML. Therefore, the dual or triple inhibition of these targets may be more effective in the treatment of CML with TKIs.

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

Authors’ contributions
YH, TO and JS designed the study. FI and YH performed the experiments. YH, TU and JS wrote, edited and revised the manuscript critically for important intellectual content. FI and YH analyzed the results. All authors have read and approved the final manuscript.

Ethics approval and consent to participate
Leukemic bone marrow specimens were collected at diagnosis, after the patients provided written informed consent for sample collection in accordance with institutional policy. The Shimane University Institutional Committee on Ethics (Shimane, Japan) approved the present study.
References


The authors declare that they have no competing interests.

Not applicable.

Competing interests

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