Bothrops moojeni L-amino acid oxidase induces apoptosis and epigenetic modulation on Bcr-Abl+ cells

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Abstract

Background: Resistance to apoptosis in chronic myeloid leukemia (CML) is associated with constitutive tyrosine kinase activity of the Bcr-Abl oncoprotein. The deregulated expression of apoptosis-related genes and alteration in epigenetic machinery may also contribute to apoptosis resistance in CML. Tyrosine kinase inhibitors target the Bcr-Abl oncoprotein and are used in CML treatment. The resistance of CML patients to tyrosine kinase inhibitors has guided the search for new compounds that may induce apoptosis in Bcr-Abl+ leukemic cells and improve the disease treatment.

Methods: In the present study, we investigated whether the L-amino acid oxidase isolated from Bothrops moojeni snake venom (BmooLAAO-I) (i) was cytotoxic to Bcr-Abl+ cell lines (HL-60.Bcr-Abl, K562-S, and K562-R), HL-60 (acute promyelocytic leukemia) cells, the non-tumor cell line HEK-293, and peripheral blood mononuclear cells (PBMC); and (ii) affected epigenetic mechanisms, including DNA methylation and microRNAs expression in vitro.

Results: BmooLAAO-I induced ROS production, apoptosis, and differential DNA methylation pattern of regulatory apoptosis genes. The toxin upregulated expression of the pro-apoptotic genes BID and FADD and downregulated DFFA expression in leukemic cell lines, as well as increased miR-16 expression – whose major predicted target is the anti-apoptotic gene BCL2 – in Bcr-Abl+ cells.

Conclusion: BmooLAAO-I exerts selective antitumor action mediated by H₂O₂ release and induces apoptosis, and alterations in epigenetic mechanisms. These results support future investigations on the effect of BmooLAAO-I on in vivo models to determine its potential in CML therapy.

Keywords:
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Background

Chronic myeloid leukemia (CML) is a myeloproliferative neoplasm characterized by the Bcr-Abl oncprotein expression, which promotes the uncontrolled cell proliferation and resistance to apoptosis [1,2]. The CML patients are currently treated with tyrosine kinase inhibitors (TKI) that target the Bcr-Abl oncprotein, including imatinib mesylate, dasatinib, and nilotinib. Despite the high rates of molecular response to TKI therapy, many patients may acquire resistance to this therapy [3,4] due to mutations in the kinase domain of the Bcr-Abl oncprotein, and to Bcr-Abl-independent mechanisms that mediate activation of alternative cell survival signaling pathways, which are associated with epigenetic and apoptotic deregulation [5].

Epigenetic deregulation plays a significant role on the development, maintenance and progression of different neoplasms, including the hematological neoplasms [6–8]. Epigenetic deregulation involves gene expression changes that promote heritable phenotype alterations with preserved DNA sequence [9]. DNA methylation and microRNA (miRNA) expression are key epigenetic regulation mechanisms for silencing gene expression [10].

DNA methylation regulates gene transcription and maintain genome stability [11]. The presence of genespecific hypermethylation in the promoter region results in transcriptional repression of genes in different types of neoplastic cells [2,6,12].

miRNA is an endogenous small non-coding RNA molecule capable of inhibiting the gene expression [13] mostly by miRNA-mRNA binding, which could result in cleavage or degradation of the expressed mRNA [14,15]. miRNA can control a variety of biological processes, such as cell differentiation, proliferation, and apoptosis, but abnormal miRNA expression is associated with pathogenesis of solid tumors and hematological neoplasms such as lymphoma, acute and chronic lymphocytic leukemia, acute promyelocytic leukemia, and CML [16,17].

The miRNA molecules that mediate regulation of apoptosis-related genes expression, known as apoptomiR, are deregulated in CML. Our research team has reported the differential apoptomiR expression in CML patients in chronic and accelerated phase, as well as in patients resistant to the TKI imatinib mesylate [17]. Patients resistant to TKI therapy present an apoptomiR expression profile linked to overexpression of anti-apoptotic genes and downregulation of pro-apoptotic genes [17]. The apoptomiR miR-15a and miR-16, which are involved in the control of the Bcl-2 protein expression, are also low expressed in Bcr-Abl+ leukemic cells [18,19]. Thus, it is relevant to seek new compounds that induce apoptosis and modulate the CML epigenetic machinery to increase sensitivity of leukemic cells to TKI therapy.

In this context, L-amino acid oxidases (LAAO) isolated from snake venom (SV-LAAO) have attracted great interest of the scientific and medical fields due to their antitumor potential [20–23]. The mechanisms by which SV-LAAO exerts their biological actions such as cytotoxicity and apoptosis induction remain unclear, but there are evidences that they are mediated by H\textsubscript{2}O\textsubscript{2} production. H\textsubscript{2}O\textsubscript{2} is a reactive oxygen species (ROS) that destabilizes mitochondrial membranes and induces cell death [22,24–27].

We have recently reported the antitumor potential of LAAO from Bothrops pirajai (BpirLAAO-I) and Calloselasma rhodostoma (CR-LAAO) venom, which induce apoptosis in Bcr-Abl+ cell lines [28–30]. The LAAO isolated from Bothrops moojeni (BmooLAAO-I) exerts antitumor action on Ehrlich ascites carcinoma cells and HL-60 acute promyelocytic leukemia cells [31]. The long-term enzymatic stability of BmooLAAO-I makes it possible to assess its pharmacological effects [32]. The present study examined whether BmooLAAO-I affected the apoptotic and epigenetic machineries of Bcr-Abl+ cell lines resistant and responsive to imatinib mesylate.

Methods

BmooLAAO-I isolation

BmooLAAO-I was isolated from a B. moojeni snake venom sample that was kindly donated by the Center for the Study of Venoms and Venomous Animals (CEVAP) of São Paulo State University (UNESP – Botucatu, SP, Brazil), and stored at \( -20 \) °C.

Crude venom (200 mg) was purified according to the protocol reported by Stábeli and colleagues [31], with minor modifications. Initially, unpurified venom sample was concentrated by ultrafiltration using an AMICON\textsuperscript{®} apparatus equipped with a 10,000 Da cutoff membrane. The concentrated fraction was purified by hydrophobic chromatography on CM-Sepharose and Phenyl-Sepharose CL-4B columns (1.0×26 cm) previously equilibrated with 0.02 M Tris-HCl buffer, pH 7.4. Elution was carried out using a reverse linear NaCl gradient (4–0 M) at a flow rate of 72 mL/h, at 25 °C, and fractions of 3.0 mL were collected. The fractions with LAAO activity were pooled, concentrated by ultrafiltration using a 30,000 Da cutoff membrane, and submitted to a third purification step by affinity chromatography on a Benzamidine Sepharose column (1.8×10 cm) previously equilibrated with 20 mM Tris-HCl, pH 7.4. The sample was eluted using a step gradient of 20 mM Tris-HCl supplemented with 1.0 M NaCl, pH 7.4, at a flow rate of 1 mL/h. Fractions of 3 mL were collected and followed by recording absorbance at 280 nm. The LAAO-active fraction was collected, concentrated by ultrafiltration using a 30,000 Da cutoff membrane, and stored at 4 °C for subsequent analysis. The purification parameters of BmooLAAO-I were summarized in the Additional file 1, it yielded 2.0% protein, specific activity of 2,806 U/mg with 12.3-purification fold.

The purity of the isolated BmooLAAO-I sample was analyzed by 12% (w/v) SDS-PAGE and high-performance liquid chromatography using a C18 reverse phase column (0.46×25 cm) equilibrated with 0.1% (v/v) trifluoroacetic acid. Elution was performed at a flow rate of 1 mL/min, for 90 min, using a concentration gradient (0–100%, v/v) of 70% acetonitrile in
0.1% trifluoroacetic acid (v/v). The fold purity of BmoLAAO-I was 12.3-fold.

**Determination of enzymatic activity of BmooLAAO-I**

Total protein concentration was determined using the BCA Protein Assay Kit (Thermo Fischer Scientific, Rockford, IL, USA), according to the manufacturer’s instructions. The BmooLAAO-I specific enzymatic activity was determined, prior to assays, by spectrophotometry using L-leucine as substrate [33].

**Cell lines**

We used the cell lines HEK-293 (embryonic epithelial cells of human kidney), HL-60 (human promyelocytic leukemia cells), HL-60.Bcr-Abl (HL-60 infected with retrovirus carrying the BCR-ABL1 gene), K562-S (imatinib mesylate-sensitive Bcr-Abl cells), and K562-R (imatinib mesylate-resistant Bcr-Abl cells). The K562-R and K562-S cells were obtained from a human CML patient in blastic phase. The tumor cell lines were kindly provided by Dr. Gustavo P. Amarante-Mendes (Institute of Biomedical Sciences, University of São Paulo, São Paulo, SP, Brazil), while HEK-293 cells were kindly provided by Dr. Andreia Machado Leopoldino (School of Pharmaceutical Sciences of Ribeirão Preto, University of São Paulo, Ribeirão Preto, SP, Brazil). All the leukemic cell lines were cultured in complete RPMI (Roswell Park Memorial Institute) 1640 medium, while HEK-293 cells were cultured in complete DMEM (Dulbecco’s Modified Eagle Medium); both media were supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin, and all the cell lines were cultured at 37 °C, under 5% CO2.

**Isolation of peripheral blood mononuclear cells**

Peripheral blood samples from three human healthy donors aged between 20 and 35 years old were collected into vacuum tubes containing EDTA (BD Vacutainer®). Peripheral blood mononuclear cells (PBMC) were isolated using the Ficoll-Hypaque density gradient method (Ficoll® Paque Plus, GE Healthcare), according to the manufacturer’s instructions. Cell viability was determined using the trypan blue exclusion assay. The Human Research Ethics Committee of the School of Pharmaceutical Sciences of Ribeirão Preto, University of São Paulo, Brazil, approved the study protocol (CAAE number 55672816.6.000.5403). All healthy volunteers signed the informed consent form and agreed to participate in the study.

**Cytotoxicity assay**

Cell viability was assessed using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) method described by Mosmann [34]. The five cell lines and PBMC cells (2×105) were cultured in 180 µL of RPMI 1640 complete medium in 96-well plates and treated with BmooLAAO-I (0.000765-0.392 µg/mL) for 24 h. The cell lines were cultured with BmooLAAO-I in the presence or not of catalase (100-400 μg/mL) (Sigma-Aldrich, St. Louis, MO, USA) [22,35,36]. Untreated cells were used as the negative control. Next, 20 µL of MTT solution (5 mg/mL) were added to each well, and the plates were incubated (4 h, 37 °C) and centrifuged (660 × g, 3 min, room temperature). The supernatants were discarded and the formazan crystals were dissolved with 200 µL of dimethyl sulfoxide. After 30 min of incubation at room temperature, absorbance was recorded at 570 nm, and the results were expressed as percentage of viable cells relative to the negative control. The cell viability percentage was used to calculate the toxin concentration that reduced cell viability by 50% (IC50), with the aid of the Calcsusyn 2.1 software. The experiments were performed in independent triplicates.

**Quantification of intracellular ROS**

Intracellular ROS generation was measured using 2’,7’-dichlorodihydrofluorescein diacetate (H2DCFDA), as reported by Wang and Joseph [37]. HL-60, HL-60.Bcr-Abl, and HEK-293 cells (2×104/well) were seeded in 96-well sterile black plates for 24 h, and treated with phosphate-buffered saline (PBS; negative control) or BmooLAAO-I at different concentrations, from 0.003075 to 0.038 µg/mL (HL-60 cells) or from 0.01225 to 0.192 µg/mL (HL-60.Bcr-Abl and HEK-293 cells). HL-60 and HL-60.Bcr-Abl cells were also treated with BmooLAAO-I in the presence or absence of 200 µg/mL catalase (Sigma-Aldrich, St. Louis, MO, USA). The supernatant was discarded, and the cells were washed with PBS and incubated with 100 µL of 10 µM H2DCFDA for 30 min, at 37 °C. The positive control cells were treated with 20 µL of 20 µM H2O2 for 20 min and washed with PBS. Finally, 100 µL of PBS were added to each well, and the fluorescence intensity was recorded in the fluorescent microplate reader Synergy H1, at the excitation and emission wavelengths of 485 and 528 nm, respectively. The percentage of intracellular ROS levels was normalized considering untreated cells (negative control) as 100%, and was calculated from the ratio between fluorescence intensity of each treated sample and fluorescence intensity of the negative control, multiplied by 100.

**Cell death quantification**

HL-60, HL-60.Bcr-Abl, K562-S, and K562-R cells (1×105) were seeded in 6-well plates and treated for 24 h with BmooLAAO-I at different concentrations: 0.003075-0.038, 0.01225-0.192, 0.01225-0.148, and 0.01225-0.166 μg/mL, respectively. Etoposide (VP-16) at 25 μM was used to induce cell death, as the positive control. Untreated cells were used as the negative control. The apoptosis levels were quantified using the annexin V-FITC/PI [38] and hypotonic fluorescent solution (HFS) methods [39].

**Annexin V assay**

After BmooLAAO-I treatment, cells were collected, centrifuged (240 × g, 10 min, 4 °C), washed with 200 µL of annexin-binding buffer, and suspended in 100 µL of annexin V-FITC previously diluted 1:2,000 in annexin-binding buffer. After a 20-min incubation at room temperature, 1 µL of 250 µg/mL propidium
iodide (PI) solution was added to each sample. Data from 10,000 events were analyzed in the FACS Canto flow cytometer (Becton-Dickinson, San Jose, CA, EUA) and the percentage of cells stained only with annexin V (annexin V/PI−) or with both annexin V and PI (annexin V+/PI+) were quantified.

HFS assay
After BmooLAAO-I treatment, cells were collected, centrifuged (240 xg, 10 min, 4 °C), and suspended in 400 μL of HFS (50 μg/mL PI in 0.1% sodiumcitrate and 0.1% Triton X-100). After incubation (20 min, 4 °C), data from 5,000 events were acquired and hypodiploid nuclei were quantified in the FACS Canto flow cytometer (Becton-Dickinson).

Analysis of protein expression
HL-60, HL-60.Bcr-Abl, K562-S, and K562-R cells (1×10⁶) were seeded in 6-well plates and treated for 24 h with BmooLAAO-I at 0.003075-0.038, 0.01225-0.192, 0.01225-0.148, and 0.01225-0.166 μg/mL, respectively, or 25 μM VP-16 (positive control). Untreated cells were used as the negative control. Next, cells were collected and suspended in Western blotting lysis buffer (20 mM Tris-HCl pH 7.4, 150 mM NaCl, 1 mM EDTA, and phosphatase and protease inhibitors).

Total protein concentration was determined using the BCA Protein Assay Kit (ThermoFisher Scientific), according to the manufacturer’s instructions. Equal protein amounts (20 μg) were submitted to SDS-PAGE analysis and transferred to polivinilidene fluoride membranes (Amersham ECL Plus, GE Healthcare Life Sciences, Pittsburgh, PA, USA). The membranes were treated with blocking solution (5% non-fat dry milk, 0.01% sodium azide) for 2 h and incubated overnight at 18 °C with primary antibodies against caspase 3 (code #96625), caspase 8 (code #9746), caspase 9 (code #9502), PARP (code #9541, anti), and Bcl-2 (code #2870) (Cell Signaling Technology, Danvers, MA, USA), previously diluted in blocking solution to appropriate concentrations. After removing unbound primary antibodies, the membranes were washed with TBS-T buffer (100 mM Tris-HCl, 300 mM NaCl, 1 % Tween20) and incubated (1 h, room temperature) with the respective peroxidase-conjugated antimouse or anti-rabbit secondary antibody (Sigma-Aldrich) diluted 1:2,000. The membranes were revealed by chemiluminescence according to the manufacturer’s instructions (Amersham ECL Plus), GE Healthcare Life Sciences). The proteins β-tubulin (code #2146; Cell Signaling Technology), β-actin (code #A1978; Sigma-Aldrich), and γ-tubulin (code #T3320; Sigma-Aldrich) were used for sample loading normalization.

DNA extraction and methylation pattern analysis
The methylation pattern was analyzed in K562-S and K562-R cells (5×10⁶ cells/well) treated for 24 h with BmooLAAO-I at 0.01225 and 0.0245 μg/mL in 6-well plates. Total DNA was extracted with QIAamp DNA Mini Kit and purified with minElute Reaction Cleanup kit (Qiagen Company; Hilden, Frankfurt, Germany). Total DNA concentration was determined using the NanoVue spectrophotometer (GE Healthcare Life Sciences). One μg of each sample was submitted to DNA restriction digestion using the EpiTect Methyl II DNA Restriction Kit (Qiagen Company). Next, the EpiTect Methyl PCR Arrays (Qiagen Company) was used to analyze methylation of the promoter region of the following apoptosis-related genes: APAF1, BAK, BAG1, BAX, BCL2, BCL2L11, BCLAF1, BID, BIK, BIRC2, BNP3L, CASP3, CASP9, CIDEB, CRADD, DAPK1, DFFA, and FADD. Samples were analyzed by real-time PCR (StepOnePlus®, Applied Biosystems) and the percentage of methylated and non-methylated genes were determined. The results were expressed as percentage of total input DNA (% of methylated DNA) associated with the corresponding heatmap created using the Hierarchical clustering method, with the aid of the Software MeV v4.8.1. All the assay kits were used according to the respective manufacturer’s instructions.

RNA extraction, cDNA synthesis, and real-time PCR analysis
HL-60 cells (5×10⁶ cells/well) were treated with BmooLAAO-I at 0.003075 and 0.00615 μg/mL, while the Bcr-Abl+ cells HL-60. Bcr-Abl, K562-S, and K562-R cells (5×10⁶ cells/well) were treated with BmooLAAO-I at 0.01225 and 0.0245 μg/mL, for 24 h in 6-well plates. Untreated cells were used as the negative control. Total RNA was extracted using the Trizol® method, following the manufacturer’s instructions (Invitrogen Life Technologies, Carlsbad, CA, USA). RNA concentrations were calculated from ratio of absorbance recorded at 260 nm and 280 nm (A260/A280), using the NanoVue spectrophotometer (GE Healthcare Life Sciences). Complementary DNA (cDNA) was reverse transcribed from 1 μg of the total RNA extracted, using the High Capacity cDNA reverse transcription kit (Applied Biosystems, Foster City, EUA), according to the manufacturer’s recommendations. cDNA (diluted 1 to 4) samples were used to quantify gene expression (hypermethylated genes) by real-time PCR (StepOnePlus™ equipment, Applied Biosystems), using the assay kit SYBR Green PCR Master Mix (Applied Biosystems, Carlsbad, CA, USA) for the genes BID (BH3 interacting domain death agonist) and FADD (Fas associated via death domain) and the reference genes B-ACTIN and B2M, and the assay kit TaqMan® Gene Expression assays (Applied Biosystems, Foster City, EUA) for the gene DFFA (DNA fragmentation factor subunit alpha) and the reference genes B-ACTIN and GAPDH. The Ct (cycle threshold) was used to calculate values of gene expression using the equation 2−ΔΔCt (fold change). Three independent experiments were performed. The oligonucleotide sequences used for quantification of target gene expression are listed in Additional file 2.

miRNA expression levels in leukemic cell lines
To analyze the level of apoptomiR expression in HL-60, HL-60.Bcr-Abl, K562-S, and K562-R cells, 2.5 ng of the total RNA extracted were used to synthesize cDNA, using the High Capacity
cDNA reverse transcription assay kit (Applied Biosystems’), with specific loop-primers for each miRNA (Applied Biosystems’). cDNA (diluted 1:4) was used to detect expression of the apoptomiR miR-15a (predicted target gene BCL2), miR-16 (predicted target gene BCL2), and hsa-let-7d (predicted target gene BCL2L1). The expression levels of each miRNA were normalized using the reference miRNA RNU24 and RNU44. miRNA expression was quantified by real-time PCR (StepOnePlus™ equipment, Applied Biosystems), using the ‘TaqMan’ microRNA assay kit for human samples (Applied Biosystems’, Foster City, USA). The Ct (cycle threshold) was used to calculate values of miRNA expression using the equation $2^{-\Delta\Delta C_T}$ (fold change).

**Data analysis**

One-way Analysis of Variance (ANOVA) followed by the Tukey’s post-hoc test was used to compare the results from experimental groups (cell lines treated with BmooLAAO-I at different concentrations) and their respective negative control groups (cell lines not treated with the toxin), in the assays to determine ROS production, cytotoxicity, apoptosis, formation of hypodiploid nuclei, gene expression, and miRNA expression. The first four assays were carried out in three biological and technical replicates, while the last two assays were carried out in two biological and technical replicates. All the experimental data were analyzed using the GraphPad Prism version 5.0 software (GraphPad Software, San Diego, California, USA), with values of p<0.05 considered as significant. The power of the statistical test applied for each cell line and assay was determined by one-way ANOVA with 5% of significance. All the tests had power greater than 0.962.

**Results**

**Isolation and purification of BmooLAAO-I**

BmooLAAO-I was successfully purified through three chromatographic steps. After the first purification step on a CM-Sepharose column, the LAAO-active fraction was identified using enzymatic assays (Additional file 3A). The resulting fraction was concentrated and subsequently applied onto a Phenyl-Sepharose CL-4B column (Additional file 3B). The resulting LAAO-active fraction was concentrated and applied onto a Benzamidine Sepharose affinity column (Additional file 3C). An LAAO-active fraction was obtained with high purity, as analyzed by C18 reversed-phase HPLC (Additional file 3D) and SDS-PAGE (Additional file 3D, inset).

The N-terminal amino acid sequence of the isolated protein obtained by automatic Edman degradation resulted in a sequence of 35 amino acid residues (ADDRNLPEECFRETDYEEFLETAKNGLSTTSNKKL) with high identity (95%) to the N-terminal sequence of the BmooLAAO-I isolated from *B. moojeni* venom by Stábeli et al. [31].

**BmooLAAO-I selective cytotoxicity to leukemic cell lines is mediated by H$_2$O$_2$ release**

We assessed the BmooLAAO-I cytotoxicity to the tumor cell lines HL-60, HL-60.Bcr-Abl, K562-S, and K562-R, as well as to the non-tumor cell line HEK-293 (Fig. 1) and PBMC (Additional file 4). The toxin decreased cell viability of HL-60 cells at concentrations greater than 0.00615 μg/mL, in a concentration-dependent manner, reducing cell viability to 9% at the highest concentration tested (0.392 μg/mL, Fig. 1A) and yielding IC$_{50}$ = 0.038 μg/mL. BmooLAAO-I at the highest concentration tested (0.392 μg/mL) strongly reduced cell viability to around 30% in HL-60.Bcr-Abl, K562-S, and K562-R cells, and afforded IC$_{50}$ values of 0.192, 0.148, and 0.166 μg/mL, respectively (Fig. 1B-D). The toxin significantly reduced cell viability of HL-60, Bcr-Abl and K562-S cells at concentrations greater than 0.049 μg/mL, and of K562-R cells at concentrations greater than 0.098 μg/mL. Compared with untreated cells, treatment with BmooLAAO-I did not significantly alter the cell viability (90-100%) of HEK-293 cells (Fig. 1E) and PBMC (Additional file 4), under the conditions assessed.

To address whether BmooLAAO-I cytotoxicity was associated with H$_2$O$_2$ release, we analyzed cell viability in the presence of catalase, a H$_2$O$_2$-degrading enzyme. Catalase at 100 μg/mL increased cell viability of all leukemic cell lines, and exerted stronger effects in HL-60.Bcr-Abl, K562-S, and K562-R treated with the highest toxin concentration (0.392 μg/mL), in which catalase augmented cell viability by 30 to 50% (Fig. 1B-D).

Each cell line was then treated with three BmooLAAO-I concentrations – the two lowest concentrations that reduced cell viability and the IC$_{50}$ value – in the presence or not of catalase at 200-400 μg/mL (Additional file 5). The three catalase concentrations tested increased cell viability to the same extent as the previously tested concentration of 100 μg/mL, in all tumor cell lines. Catalase significantly augmented cell viability of HL-60 and HL-60.Bcr-Abl cells treated with the two highest toxin concentrations (Additional file 5A-B), and of K562-S and K562-R cells treated with the three toxin concentrations tested (Additional file 5C-D).

**BmooLAAO-I increases ROS generation in leukemic cell lines**

To confirm that BmooLAAO-I cytotoxicity resulted from ROS production, we quantified intracellular ROS generation in the leukemic cell lines HL-60 and HL-60.Bcr-Abl, and in the non-tumor cell line HEK-293 (Fig. 2). BmooLAAO-I at intermediate concentrations (0.01225 and 0.0245 μg/mL) enhanced ROS levels in HL-60 cells (Fig. 2A) and, at concentrations greater than 0.049 μg/mL, it enhanced ROS levels in HL-60.Bcr-Abl cells (Fig. 2D). The toxin did not alter ROS levels in HEK-293 cells (Fig. 2C). Catalase mitigated the toxin-elicited increased ROS levels in HL-60 and HL-60.Bcr-Abl cells (Fig. 2B, 2E). These results indicate the association between ROS production and BmooLAAO-I cytotoxicity in these leukemic cell lines.
Figure 1. Cytotoxicity of BmooLAAO-I towards four tumor cell lines. (A) HL-60, (B) HL-60.Bcr-Abl, (C) K562-S, and (D) K562-R cells were treated with the toxin for 24 h, in the presence or absence of 100 μg/mL of catalase. (E) HEK-293 cells were treated with the toxin for 24 h only without catalase due to the lack of cytotoxicity of BmooLAAO-I towards this cell line. Results are expressed as mean ± standard deviation of the percentage of cell viability from three independent experiments assayed in triplicate. NC: negative control (untreated cells); CC: catalase control (cells treated only with catalase). *p < 0.05 vs. NC, *p < 0.05 vs. catalase (−) (one-way ANOVA combined with the Tukey’s post-hoc test).

Figure 2. Intracellular ROS generation in cells treated with BmooLAAO-I for 24 h. (A, B) Fluorescence intensity of H₂DCFDA was measured in HL-60, (C) HEK-293 and (D, E) HL-60.Bcr-Abl cells. HL-60 and HL-60.Bcr-Abl cells were treated with the toxin in (A, D) the absence or (B, E) presence of 200 μg/mL of catalase. Results are expressed as mean ± standard deviation of three independent experiments. NC: negative control (untreated cells); CC: catalase control (cells treated only with catalase). Values not sharing the same letter are significantly different from each other (p < 0.05; one-way ANOVA combined with the Tukey’s post-hoc test).
BmooLAAO-I induces apoptosis in leukemic cell lines

Considering the cytotoxic effect of BmooLAAO-I in leukemic cell lines, we investigated whether BmooLAAO-I induced apoptosis in these tumor cells (Fig. 3). BmooLAAO-I at the concentration of 0.0245 and 0.038 µg/mL increased the percentage of annexin V⁺/PI⁻ cells (from 23 to 32%), annexin V⁺/PI⁺ cells (from 16 to 33%), and hypodiploid nuclei (from 40 to 76%) in HL-60 cells (Fig. 3A-B). The toxin at the concentration range of 0.049-0.192 µg/mL augmented the percentage of annexin V⁺/PI⁺ cells (38-75%) and hypodiploid nuclei (34-70%), but not of annexin V⁺/PI⁻ cells in HL-60.Bcr-Abl cells (Fig. 3C-D).

![Figure 3](image-url)

**Figure 3.** Quantification of apoptosis levels induced by BmooLAAO-I. The percentage of annexin V⁺/PI⁻ (white bars) and annexin V⁺/PI⁺ (black bars) was determined in (A) HL-60, (C) HL-60.Bcr-Abl, (E) K562-S, and (G) K562-R cells. The percentage of hypodiploid nuclei was determined in (B) HL-60, (D) HL-60.Bcr-Abl, (F) K562-S, and (H) K562-R cells, using the hypotonic fluorescent solution method. Results are expressed as mean ± standard deviation of three independent experiments. NC: negative control (untreated cells); VP-16: etoposide. *p < 0.05 vs. NC (annexin V⁺/PI⁻ and hypodiploid nuclei); *p < 0.05 vs. NC (annexin V⁺/PI⁺) (one-way ANOVA combined with the Tukey’s post-hoc test).
In K562-S cells, the toxin increased the percentage of annexin V+/PI- cells (35-61%) and hypodiploid nuclei (40-87%) at 0.049-0.148 µg/mL, and the percentage of annexin V+/PI- cells (26-28%) at 0.098 and 0.148 µg/mL (Fig. 3E-F). In K562-R cells, the toxin augmented the percentage of annexin V+/PI- cells (18-31%) at 0.0245-0.166 µg/mL, and the percentage of annexin V+/PI- cells (33-52%) and hypodiploid nuclei (35-52%) at 0.049-0.166 µg/mL (Fig. 3G-H). The toxin at 0.098 and 0.166 µg/mL induced weaker formation of hypodiploid nuclei in K562-R cells than in K562-S cells (Additional file 6).

VP16 induced cell apoptosis in HL-60 and K562-R cells, but not in HL-60.Bcr-Abl and K562-S cells (Fig. 3). This compound induced the formation of hypodiploid nuclei with strong intensity in HL-60 cells (87%, Fig. 3B), moderate intensity in K562-R cells (40%, Fig. 3H), and weak intensity in K562-S (24%, Fig. 3F) and HL-60.Bcr-Abl (18%, Fig. 3D) cells.

**BmooLAAO-I activates caspases 3, 8 and 9 in leukemic cells**

To examine whether BmooLAAO-I induced apoptosis via the intrinsic or extrinsic pathway activation, we analyzed the expression levels of caspases 3, 8, and 9. The toxin lowered expression of pro-caspases 3, 8, and 9 in all tumor cell lines (Fig. 4) and increased the levels of cleaved caspase 8 in K562-R cells (Fig. 4D) and HL-60.Bcr-Abl cells (Fig. 4B).

**BmooLAAO-I modulates the methylation pattern of apoptosis-related genes in K562-S cells**

Considering the mechanisms by which BmooLAAO-I induced apoptosis in leukemic cells, we examined whether it modulated the DNA methylation pattern in the promoter region of apoptosis-related genes in K562-S (Fig. 5) and K562-R (Additional file 7).

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**Figure 4.** Western blotting analysis of BmooLAAO-I-induced expression of cleaved caspases in tumor cell lines. (A) The levels of caspases 3, 8 and 9 were determined in HL-60, (B) HL-60.Bcr-Abl, (C) K562-S, and (D) K562-R cells after 24 h of treatment with BmooLAAO-I. Decreased pro-caspase expression and increased expression of active (cleaved) forms indicates caspase activation. NC: negative control (untreated cells), VP-16: etoposide (positive control).
cells. The cells were treated with BmooLAAO-I at the sublethal concentrations of 0.01225 and 0.0245 μg/mL. Compared with the negative control (Fig. 5A), the pro-apoptotic genes BID, FADD, and DFFA were hypermethylated (52.42%, 60.45%, and 68.51%, respectively) in K562-S cells treated with the toxin at 0.0245 μg/mL (Fig. 5C). Heatmap analysis showed that the three genes were hypermethylated, but the other apoptosis-related genes exhibited methylation levels similar to those detected in the negative control (Fig. 5D). The toxin did not alter the methylation pattern of apoptosis-related genes in K562-R cells (Additional file 7).

**BmooLAAO-I modulates expression of BID, FADD, and DFFA genes**

Since BmooLAAO-I promoted hypermethylation of the genes BID, FADD, and DFFA, we analyzed their expression levels in K562-S cells (Fig. 5E-G). The toxin at 0.01225 μg/mL upregulated BID expression [fold change (fc) = 1.78, Fig. 5E] and downregulated DFFA expression (fc = 0.26, Fig. 5G). At the concentration of 0.0245 μg/mL, the toxin did not alter the BID and DFFA expression levels, but it upregulated FADD expression (fc = 2.42, Fig. 5F).

![Figure 5](image-url)

**Figure 5.** BmooLAAO-I modulates the apoptosis-related gene methylation pattern in K562-S cells. The percentage of methylation of the promoter region of apoptosis-related genes was quantified by real-time PCR in cells treated with BmooLAAO-I for 24 h. (A) Untreated cells (negative control). (B) Cells treated with the toxin at 0.01225 μg/mL. (C) Cells treated with the toxin at 0.0245 μg/mL. (D) Heatmap of sample clustering according to the percentage of methylated DNA. The horizontal bar in the top of the heatmap represents the color scale of percentage of methylation ranging from 0-100%. Expression levels of hypermethylated genes in K562-S cells treated with BmooLAAO-I. Expression of the genes (E) BID, (F) FADD, and (G) DFFA was quantified by real-time PCR after a 24 h treatment with BmooLAAO-I at sublethal concentrations (0.01225 and 0.0245 μg/mL). Results were expressed as mean fold change ± standard deviation of three independent experiments. NC: negative control (untreated cells). *p < 0.05 vs. NC (one-way ANOVA followed by the Tukey’s post-hoc test).
BmooLAAO-I modulates miR-16 and Bcl-2 protein expression in K562-R cells

Considering that regulation by miRNA is another important epigenetic mechanism involved with tumorigenesis and apoptosis resistance, we detected the expression levels of the apoptomiR miR-15a, miR-16, and hsa-let-7d in leukemic cell lines treated with BmooLAAO-I at sublethal concentrations. BmooLAAO-I did not alter the expression levels of miR-15a (Additional file 8A) and hsa-let-7d (Additional file 8B). Notably, the toxin downregulated miR-16 expression in K562-S cells at the concentration of 0.01225 µg/mL (fc = 0.75), and upregulated miR-16 expression in K562-R cells at the concentrations of 0.01225 and 0.0245 µg/mL (fc = 1.26 and 2.12, respectively) (Fig. 6A). The expression level of the Bcl-2 protein, one of the miR-15a/miR-16 predicted target gene, decreased in K562-S and K562-R cells treated with both toxin concentrations (Fig. 6B).

Discussion

Natural compounds, including SV-LAAO, have exhibited strong antitumor activity [22,28–30,32,40]. In the present study, BmooLAAO-I decreased cell viability of all the leukemic cell lines tested: HL-60, HL-60.Bcr-Abl, K562-S, and K562-R. To address whether H₂O₂ released during the LAAO enzymatic activity of BmooLAAO-I mediated cell death, we carried out the cytotoxicity assay in the presence of 100 µg/mL catalase. Addition of such H₂O₂-degrading enzyme increased cell viability of the four leukemic cell lines treated with different toxin concentrations. This finding is in line with our previous reports that 100 µg/mL catalase mitigates cytotoxicity of CR-LAAO towards HL-60, HL-60.Bcr-Abl, K562, and KCL22 cells [30], as well as with other literature reports that catalase mitigates cytotoxicity of LAAO from B. pirajai, Bothrops atrax, Bothrops pauloensis, Lachesis muta, and Ophiophagus hannah to the tumor cell lines SBKR-3, Jurkat, EAT, MCF-7, A549, Jurkat, and B16/F10 [22,35,36,41,42].

One essential mechanism for BmooLAAO-I cytotoxicity is ROS overproduction. We confirmed that BmooLAAO-I induced ROS production in leukemic cell lines in a concentration-dependent manner, but it did not alter ROS levels in the non-tumor HEK cells. Catalase not only mitigated the toxin-mediated ROS overproduction due to degradation of H₂O₂ but also increased cell viability, indicating that H₂O₂ was a key player on the BmooLAAO-I cytotoxicity to leukemic cell lines, especially to Bcr-Abl⁺ cells. In agreement with our findings, LAAO isolated from Bothrops alternatus and Bothrops jararacussu snake venoms induce ROS overproduction in many tumor cell types, including hepatocarcinoma, leukemia T, breast cancer, adenocarcinoma, and melanoma; however, the enhanced ROS levels are associated with anti-tumor effects due their contribution to cell death and DNA damage [43,44].

Considering the strong pharmacological effects and the risk of toxicity of snake venom toxins to humans, their cytotoxicity must be evaluated in non-tumor cells. This study demonstrated that BmooLAAO-I was not cytotoxic to the non-tumor cell line HEK-293 and to PBMC cells, corroborating previous reports that SV-LAAO is more cytotoxic to tumor cells than to non-tumor cells [28,29,35,45].

After analyzing the BmooLAAO-I-induced cytotoxicity and ROS production, we examined the toxin ability to sensitize and/or induce apoptosis in leukemic and non-leukemic cells. Apoptosis is an important cell process to be analyzed during antitumor drug development, because it is one of the main routes of clearance of neoplastic cells [46–48]. It is well-known that Bcr-Abl⁺ cells are more resistant to apoptosis induced by chemotherapeutic agents and classical apoptogenic stimuli.
Hence, our finding that BmooLAAO-I was capable of inducing apoptosis in both Bcr-Abl+ and Bcr-Abl leukemia cells is relevant. The fact that K562-S cells are more sensitive to the TKI imatinib mesylate than K562-R cells [51,52] may explain, at least in part, why this cell was more sensitive to BmooLAAO-I cytotoxicity.

The glycans moiety of SV-LAAO favors their anchoring in tumor cells and increases the local concentration of H2O2, which thereby causes oxidative damage and induces apoptosis [53]. Many studies have demonstrated the apoptosis-inducing potential of SV-LAAO in a variety of tumor cell lines, such as HL-60, A2780, K562, Jurkat, B16/F10, and A549 [20,22,23,25,54–56]. Our research team has reported that BpirLAAO and CR-LAAO stimulate apoptosis in primary cells from chronic myeloid leukemia (CML) patients and in Bcr-Abl+ cell lines [28,29].

Apoptosis stimulation by BmooLAAO-I is accompanied by activation of caspasases 3, 8, and 9 in the tumor cell lines, indicating that this toxin induces apoptosis via the intrinsic and extrinsic pathways. This finding is in line with previous reports from our research team that BpirLAAO-I, CR-LAAO, and BjuussLAAO-II (a LAAO isolated from B. jararacussu venom) promote activation of caspasases 3, 8 and 9 in tumor cell lines [23,28,30,33]. Many authors have reported the potential of SV-LAAO to activate caspases in tumor cell lines, including HeLa, A549, and MCF-7 cells [36,57,58].

The deregulated expression of a number of apoptosis-related genes in CML patients may be explained, at least in part, by epigenetic deregulation such as DNA methylation and altered miRNA expression [6,17]. To better understand the molecular mechanisms by which the toxin induced apoptosis, we examined whether BmooLAAO-I modulated the DNA methylation pattern of apoptosis-related genes, as well as the expression levels of apoptomiR.

In K562-S cells, the toxin at 0.0245 µg/mL induced hypermethylation of the promoter region of the genes BID, FADD, and DFFA, but it did not supress their gene expression levels. Paradoxically, the toxin increased the expression of the pro-apoptotic genes BID and FADD at 0.01225 and 0.0245 µg/mL, respectively, and decreased DFFA expression at 0.01225 µg/mL; hence, the hypermethylation level was not sufficient to suppress BID and FADD gene expression in cells treated with 0.0245 µg/mL BmooLAAO-I. Essentially, DNA methylation does not imply gene silencing because suppression of gene expression requires recruitment of regulatory proteins and chromatin remodeling factors to the methylated region. However, DNA methylation is the key step of these gene regulation [59]. We should also consider that other epigenetic mechanisms can regulate gene expression, such as miRNA expression and histone acetylation [60]. Additional studies are required to better elucidate the role that DNA methylation plays on the BmooLAAO-I-induced expression of apoptosis genes.

The BmooLAAO-I-induced upregulation of BID and FADD gene expression suggests that the toxin sensitizes leukemic cells to apoptosis, which is an advantage for the treatment of neoplasias, including CML [61]. During apoptosis activation, FADD is recruited with pro-caspase 8 and activates caspase 8, which in turn activates caspase 3. The protein BID represents a connection point between the extrinsic and intrinsic apoptosis pathways, and it is activated by caspase 8-mediated cleavage [62,63]. DFFA plays a controversial role on apoptosis induction because it can act as an anti- or pro-apoptotic gene [64].

Analysis of miRNA expression pattern in some type of tumors can explain or elucidate the mechanisms involved in tumor resistance or sensitivity to different treatments, and help to select therapeutic approaches [65,66]. We analyzed the apoptomiR expression profile in K562-S and K562-R cells. The major finding was the elevated expression of the apoptomiR miR-16 in K562-R cells treated with BmooLAAO-I. The increased expression of miR-16 is associated with lowered levels of the anti-apoptotic protein Bcl-2 [17–19]. BmooLAAO-I reduced Bcl-2 levels in K562-R cells, indicating that the expression levels of miR-16 and Bcl-2 were inversely correlated. Upregulation of miR-16 expression by BmooLAAO-I is of great importance because it lowers the levels of Bcl-2 and increases the sensitivity of Bcr-Abl+ leukemic cells to apoptosis. The Bcrl-tyrosine kinase activity is associated with apoptosis inhibition mediated by increased levels of anti-apoptotic proteins [49,67].

Conclusion
Taken together, the findings of the present study demonstrate the antitumor potential of BmooLAAO-I, which acts through ROS production, apoptosis induction, and upregulation of expression of the apoptomiR miR-16. In addition, the selective toxicity towards tumor cells and the low toxicity to non-tumor cells and PBMC cells open new perspectives for application of BmooLAAO-I in antitumor therapy.

Abbreviations
ANOVA: analysis of variance; apoptomiR: microRNA that regulate apoptosis-related genes; BmooLAAO: L-amino acid oxidase from Bothrops moojeni; BpirLAAO: L-amino acid oxidase from Bothrops pirajai; cDNA: complementary DNA; CML: chronic myeloid leukemia; CR-LAAO: L-amino acid oxidase from Calloselasma rhodostoma; H2DCFDA: 2’,7’-dichlorodihydrofluorescein diacetate; HFS: hypotonic fluorescent solution; LAAO: L-amino acid oxidase; miRNA: microRNA; MTT: 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; PBMC: peripheral blood mononuclear cells; ROS: reactive oxygen species; SV-LAAO: L-amino acid oxidase from snake venom; TKI: tyrosine kinase inhibitor; VP-16: etoposide.

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**Availability of data and materials**

Not applicable.

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**Competing interests**

The authors report no conflicts of interest.

**Author’s contributions**

SMB, MCC and JGC contributed equally to the conceiving and writing of this manuscript. SMB designed and performed the experiments, analyzed the data, and drafted the manuscript. MCC and JGC performed the experiments, analyzed the data, discussed the results and drafted the manuscript. RADG, LMGA and ARTM performed the intracellular ROS assay. FSD performed the heatmaps. LA performed the western-blotting assays. ACOC purified the toxin. SVS provided the toxin, searched funding resources for this study and discussed the results. FAC searched funding resources for this study, devised the project, designed the study, discussed the results, and drafted the paper. All authors carefully revised the paper and approved the final version.

**Ethics approval and consent to participate**

We declare that the Human Research Ethics Committee of the School of Pharmaceutical Sciences of Ribeirão Preto, University of São Paulo, Brazil, approved the study protocol, which was registered under the CAAE number 55672816.6.00005403. We declare that all participants signed the informed consent form and agreed to participate in the study.

**Consent for publication**

Not applicable.

**Supplementary material**

The following online material is available for this article:

**Additional file 1.** Parameters of L-amino acid oxidase from Bothrops moojeni snake venom purification procedure.

**Additional file 2.** Oligonucleotide sequences used for quantification of target gene expression.

**Additional file 3.** Purification of the L-amino acid oxidase BmooLAAO-I from Bothrops moojeni snake venom. (A) Chromatographic profile of 200 mg of desiccated venom applied onto a CM-sepharose column previously equilibrated with 0.05 M NH₄HCO₃ buffer, pH 8.0, and eluted with a concentration gradient of up to 1.0 M of the same buffer (buffer B) at room temperature. The LAAO-active fraction eluted in the first peak. (B) Chromatographic profile of the pooled and concentrated LAAO-active fraction obtained in (A), applied onto a phenyl-sepharose column previously equilibrated with 0.02 M Tris–HCl buffer, pH 7.6 (buffer B) containing 4 M NaCl, and eluted with a discontinuous gradient of 4–0 M NaCl in the same buffer. (C) Chromatographic profile of the pooled and concentrated LAAO-active fraction obtained in (B), applied onto a benzamidine sepharose column pre-equilibrated with 20 mM Tris–HCl, pH 7.4 (buffer A), and eluted with a step gradient of 20 mM Tris–HCl containing 1.0 M NaCl, pH 7.4 (buffer B). (D) Purity of the active enzyme BmooLAAO-I assessed by SDS-PAGE (inset) and reversed-phase HPLC.

**Additional file 4.** Cytotoxicity of BmooLAAO-I towards PBMC, at 24 h post-treatment. Results are expressed as mean ± standard deviation of the percentage of cell viability from three samples assayed in triplicate. NC: negative control (untreated cells). p > 0.05 vs. NC (one-way ANOVA combined with the Tukey’s post-hoc test).

**Additional file 5.** Cytotoxicity of BmooLAAO-I towards tumor cell lines in the presence of 200–400 µg/mL of catalase. (A) HL-60 cells, (B) HL-60.Bcr-Abl cells, (C) K562-S cells, and (D) K562-R cells. Results are expressed as mean ± standard deviation of the percentage of cell viability from three independent experiments assayed in triplicate. Cells were treated with the toxin for 24 h. NC: negative control (untreated cells); CC: catalase control (cells treated with catalase only). *p < 0.05 vs. (−) catalase (one-way ANOVA combined with the Tukey’s post-hoc test).

**Additional file 6.** Comparison between BmooLAAO-I-induced apoptosis levels in K562-S and K562-R cells. (A) Total percentage of annexin-V-stained cells. (B) Percentage of hypodiploid nuclei. Results are expressed as mean ± standard deviation of three independent experiments. NC: negative control (untreated cells). *p < 0.05 vs. NC (one-way ANOVA combined with the Tukey’s post-hoc test).

**Additional file 7.** BmooLAAO-I did not alter the methylation pattern of apoptosis-related genes in K562-R cells. The percentage of methylation of the promoter region of apoptosis-related genes was quantified by real-time PCR in cells treated with BmooLAAO-I for 24 h. (A) Untreated cells (negative control). (B) Cells treated with the toxin at 0.01225 µg/mL. (C) Cells treated with the toxin at 0.0245 µg/mL. (D) Heatmap of sample clustering according to the percentage of methylated DNA. The horizontal bar in the top of the heatmap represents the color scale of percentage of methylation ranging from 0-100%.
Additional file 8. ApoptomiRs expression in tumor cell lines treated with BmooLAAO-I. Real-time PCR quantification of the apoptomiRs (A) miR-15a and (B) has-let-7d in HL-60, HL-60.Bcr-Abl, K562-S, and K562-R cells treated for 24 h with BmooLAAO-I at sublethal concentrations. Results are expressed as mean fold change ± standard deviation of three independent experiments. NC, negative control (untreated cells). *p < 0.05 vs. NC (one-way ANOVA followed by the Tukey’s post-hoc test).

References


