In vitro Antioxidant and Enzymatic Approaches to Evaluate Neuroprotector Potential of Blechnum Extracts without Cytotoxicity to Human Stem Cells

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ABSTRACT

BACKGROUND: Investigation of selected plant extracts on multi-targets related to neurodegeneration, such as monoamine oxidases (MAO), cholinesterase enzymes, and antioxidant activities (AOA) is a useful tool for identification of new scaffolds. OBJECTIVE: This work investigated biological effects of three Blechnum methanol extracts from Brazil and chemical profile of the most active sample. MATERIALS AND METHODS: AOA included scavenging of hydroxyl and nitric oxide radicals, also lipid peroxidation inhibition. Enzymatic modulation of Blechnum binervatum, Blechnum brasiliense, and Blechnum occidentale extracts on MAO and cholinesterases was conducted. Moreover, total phenol content was performed with all samples, and high-performance liquid chromatography-diode array detection mass spectrometry analysis was carried out with B. brasiliense. Possible toxic effects were evaluated on Wistar rats polymophonuclear cells (PMN) and human stem cells. RESULTS: B. brasiliense extract presented the highest phenolic amount (9.25 g gallic acid equivalent/100 g extract) and lowest IC50 values (112.3 ± 2.61 and 176.1 ± 1.19 µg/mL) against hydroxyl radicals and on lipid peroxidation, respectively, showing strong AO effects. On nitric oxide assay and cholinesterase inhibition, all extracts were considered inactive. MAO-A selective action was evidenced, being B. brasiliense powerful against this enzyme (IC50: 72.7 µg/mL), followed by B. occidentale and B. binervatum (IC50: 130.85 and 165.2 µg/mL). No cytotoxic effects were observed on PMN and human stem cells treated with Blechnum extracts. HPLC-DAD-MS analysis of B. brasiliense allowed the identification of chlorogenic and rosmarinic acids. CONCLUSION: Our results especially highlight B. brasiliense, with pronounced phenols content and strong effects on selected targets related to neurodegeneration, being characterized as a natural safe source of bioactive hydroxycinnamic acids.

Key words: Antioxidants, Blechnum, high-performance liquid chromatography-diode array detection-mass spectrometry, monoamine oxidases inhibitors, phenolic content, stem cells

SUMMARY

• Blechnum crude extracts showed high phenolic amounts and valuable IC50 values on targets related with neurodegenerative disorders
• Blechnum brasiliense was the most active sample, with strong radical scavenging and lipid peroxidation inhibition, also with monoamine oxidases:

A selective modulation
• No cytotoxic effects were observed on polymophonuclear cells rat cells and human stem cells treated with Blechnum extracts
• High-performance liquid chromatography-diode array detection-mass spectrometry analysis of Blechnum brasiliense allowed the identification of hydroxycinnamic derivatives: Chlorogenic and rosmarinic acids.

INTRODUCTION

Ferns are vascular plants, globally distributed, covering approximately 13,600 species, found in several biomes. Additionally, ferns present a great chemical diversity and several biological activities, including anti-inflammatory, antioxidant and are capable to modulate enzymes related to neurodegeneration.1,2 The genus Blechnum is chemically characterized by the presence of phenolic compounds such as flavonols, hydroxycinnamic acids, and lignans; compounds with recognized potential central nervous system activities. Among species studied in this work, Blechnum occidentale is used in folk medicine for treatment of inflammatory and pulmonary diseases, urinary tract infections, and others.3,4 Nonato et al.3,4 showed that methanol extracts of B. occidentale leaves, when administered intraperitoneally

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and orally, produced anti-inflammatory and antinociceptive effects in animals. Our research group have investigated the multifunctional feature of extracts from selected plants and isolated compounds able to inhibit targets related to neurodegeneration, such as monoamine oxidases (MAO), cholinesterase enzymes and antioxidative activities, according to the proposal scheme by Novaroli et al.[4] The imbalance between reactive species formation and repairing overproduction thereof, results in oxidative damage to membranes, proteins, lipids, and nucleic acids. Evidence have supported its involvement in the etiology of several diseases, including neurodegeneration.[3]

Pharmacological activity associated with preclinical evaluation of toxicity from natural product extracts should be performed by appropriate bioassays. Determination of action mechanisms is important, also selection of the most active crude mixture devoid of toxicity, aiming to search novel potential natural sources of compounds, as by bioassay-guided fractionation.[6]

Hence, the aim of this study was to perform in vitro antioxidant, enzymatic and toxicological studies of methanolic extracts of Blechnum binervatum, Blechnum brasiliense, and B. occidentale fronds, in addition to quantify the total phenol contents. Different biological systems were employed, including stabilization of hydroxyl and nitric oxide radicals, inhibition of lipid peroxidation, effects against MAO and cholinesterases, and possible toxic responses of rat cells and human stem cells after incubation with Blechnum extracts. In addition, high-performance liquid chromatography-diode array detection-mass spectrometry (HPLC-DAD-MS) analysis of most active sample was performed in order to search substances responsible for powerful properties.

MATERIALS AND METHODS

Materials

Human MAO-A and MAO-B supersomes were acquired from BD Gentest (Woburn, MA). Acetylcholinesterase from Electrophorus electricus, butyrylcholinesterase from equine serum, acetylthiocholine iodide (ACHe), S-butyrylthiocholine iodide (BChE), 5,5′-dithiobis (2-nitrobenzoic acid) (Ellman’s reagent), phystostigmine, kynurenine, pargyline, clorgyline, 2-deoxy-D-ribose, sodium nitroprusside dihydrate, sulfanilamide, 2-thiobarbituric acid (TBA, 1%) were performed. The plates were incubated for 30 min and absorbance measured at 532 nm in SpectraMax® apparatus (Molecular Devices, CA, USA). As positive control, chlorogenic and caffeic acids were employed.[9] Analyzes were performed in triplicate, and half maximal inhibitory concentration values (IC_{50}) were estimated.

Stabilization of hydroxyl radical

Extracts (1-500 µg/mL) were diluted in phosphate buffer and added to the reaction system in 96-well plate, containing 2-deoxyribose (50 mM), ferrous sulfate (60 µM), and hydrogen peroxide (29%) and phosphate buffer (20 mM, pH 7.2). After, the addition of phosphoric acid (4%) and thiobarbituric acid (TBA, 1%) were performed. The plates were incubated for 30 min and absorbance measured at 532 nm in SpectraMax® apparatus (Molecular Devices, CA, USA). As positive control, chlorogenic and caffeic acids were employed.[9] Analyzes were performed in triplicate, and half maximal inhibitory concentration values (IC_{50}) were estimated.

Nitric oxide radical scavenging activity

Samples (1-500 µg/mL) were prepared in phosphate saline buffer (20 mM, pH 7.2) and added to sodium nitroprusside solution (20 mM) in 96-well plates. Incubation was performed for 60 min, at room temperature. After, Griess reagent (sulphanilamide [2%] and naphthylethylenediamine dihydrochloride [0.1%]) was included and incubation of reactions was performed in the dark, for 7 min. Finally, formed nitrite levels were quantified at 546 nm using SpectraMax® apparatus.[10] The experiment was carried out in triplicate and IC_{50} values from extracts and standards (chlorogenic and caffeic acids) were calculated.

Thiobarbituric acid reactive substances assay

To perform thiobarbituric acid reactive substances (TBARS) experiment, approval by Ethics Committee on Animal Use from UFRGS was obtained (Protocol 23374). Cortex and hippocampus of adult male Wistar rats were employed, weighing between 180 and 220 g. Animals were sacrificed by decapitation and brain structures were immediately obtained and incubated with tris (hydroxymethyl)aminomethane, methanol, acetonitrile and formic acid were purchased from Sigma-Aldrich Chemical (St. Louis, MO, USA). Phosphate salts, potassium chloride, and sodium hydroxide came from Fluka (Buchs, Switzerland). Trichloroacetic acid, hydrogen peroxide, and ferrous sulfate were obtained from Synth (Diadema, SP, Brazil). Phosphoric and 2-thiobarbituric acids were obtained from Merck (Darmstadt, Germany). Lactate dehydrogenase (LDH) commercial kit was purchased from Doles (Goiás, Brazil). Phosphoric and 2-thiobarbituric acids were obtained from Merck (Darmstadt, Germany). Lactate dehydrogenase (LDH) commercial kit was purchased from Doles (Goiás, Brazil).

Plant material

Blechnum species were harvested in Rio Grande do Sul state, with previous approval from Conselho de Gestão do Patrimônio Genético and identified by botanical Maria Angelica Kieling-Rubio. Exsiccates were deposited in the Herbarium of Instituto de Biociências (ICN) from Federal University of Rio Grande do Sul (UFRGS). B. binervatum (ICN 171553) and B. brasiliense (ICN 177668) were harvested in Morro Reuter (29°32′17″S, 51°04′51″W) city while B. occidentale (ICN 177667) was harvested in Campo Bom (29°40′39″S, 51°01′97″W).

Preparation of methanolic extracts

Fronds of Blechnum species were dried at room temperature in the shadow. Reduction was performed using mill of knives and plants were extracted exhaustively with methanol by maceration (3 × 5 days), using 1:20 proportion (drug: solvent). Extracts were combined and evaporated under reduced pressure at temperature below 40°C. Yields obtained were 13.6%, 11.6%, and 15.8% for B. binervatum, B. brasiliense, and B. occidentale, respectively.

Determination of total phenol content

Total phenol content was determined according to Folin-Ciocalteu method, using a spectrophotometer at 760 nm. Results were expressed as grams of gallic acid equivalent (GAE)/100 g of crude extract.[12] Antioxidant assays

Stabilization of hydroxyl radical

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Thiobarbituric acid reactive substances assay

To perform thiobarbituric acid reactive substances (TBARS) experiment, approval by Ethics Committee on Animal Use from UFRGS was obtained (Protocol 23374). Cortex and hippocampus of adult male Wistar rats were employed, weighing between 180 and 220 g. Animals were sacrificed by decapitation and brain structures were immediately removed and washed with tris (hydroxymethyl)aminomethane (TRIS) buffer (20 mM, pH 7.4). Tissues were placed in Potter-Elvehjem homogenizer to obtain a cell homogenate which was centrifuged at 7500 rpm, for 5 min, removing the supernatant. Samples were diluted in purified water (1–500 µg/mL) and incubated with homogenate brain tissue supernatant, FeSO_4 (10 mM) and ascorbic acid (0.1 mM), for 60 min at 37°C. After, addition of trichloroacetic acid (28%) and TBA (2%) was performed. The plates were incubated for 30 min and absorbance measured at 532 nm in spectrophotometer (SpectraMax®).[11] Analyzes were performed in triplicate, and IC_{50} values were obtained.
Inhibition of monoamine oxidase enzyme

In black microplates were added in this order: Phosphate saline buffer (20 mM, pH 7.4), kynuramine (5 mM), and samples (1-500 µg/mL) in DMSO (final concentration of 1%). Mixtures were incubated at 37°C, for 20 min. Following, enzyme isoforms were added separately, MAO-A (0.09 mg/mL) and MAO-B (0.15 mg/mL), and microplates were incubated at 37°C, for 30 min. At the end of incubation, sodium hydroxide (2 M) was added to stop reactions. Fluorescence readings (excitation λ = 320 nm and emission λ = 400 nm) were measured in SpectraMax® plate reader.[12] Clorgyline and pargyline were used as control inhibitors of MAO-A and MAO-B, respectively. IC₅₀ values of samples were calculated after triplicate analysis.

Inhibition of acetyl and butyrylcholinesterase enzymes

To evaluate inhibition of acetyl and butyrylcholinesterase, Ellman reagent (10 mM), substrate solution 14 mM (AChE and BChE), samples diluted in DMSO (1%) and respective enzyme (1 U/mL) were added in 96-well plates. After enzyme addition, reactions were immediately measured in kinetic mode at 412 nm (SpectraMax®), during 6 min at intervals of 30 s.[13,14] Physostigmine was used as positive control and Blechnum extracts (100-500 µg/mL) were evaluated in triplicate to estimate IC₅₀ values.

Evaluation of cytotoxicity and cell viability

Polymorphonuclear animal cells assay

Polymorphonuclear cells (PMN) obtained from plasma of Wistar rats (final concentration of 1.5 × 10⁵ cells/mL) were employed to evaluate cytotoxicity of Blechnum extracts. Animals were previously treated to obtain a PMN cells pellet, as described by Andrade et al.[15] PMN cells were preincubated with samples at 37°C, for 30 min. After centrifugation, enzyme substrate and ferric alum were added to supernatant and mixture was incubated for 3 min. Nicotinamide adenine dinucleotide was added, and incubation for 5 min was performed. At the end, stabilization solution was added to reaction medium and analysis was performed at 492 nm using a commercial kit of LDH enzyme (Doles Reagents, Goiás, Brazil). Extracts were assessed in triplicate (1 mg/mL), and Triton X-100 (1%) was used as positive control.

Human mesenchymal stem cells assay

Stem cells from human exfoliated deciduous teeth (SHED) were isolated as previously described by Bernardi et al.[16] in partnership with Graduate Program in Dentistry at UFRGS, approved by the Ethics Committee of UFRGS (Protocol 36403514.6.0000.5347). Cell cultures were seeded and maintained in DMEM supplemented medium (pH 7.4) in an atmosphere containing 5% CO₂ at 37°C. After culture, SHED characterization was performed by flow cytometry (FACSARia III BD®, CA, USA) incubated with specific monoclonal antibodies (CD14, CD34, CD45, CD73, CD90, CD105, and human leukocyte antigen - antigen D related [HLA]-DR). Additionally, cells were analyzed for their ability to differentiate into osteoblasts, chondrocytes, and adipocytes through induction media.[17] SHED were plated (7 × 10⁴ cells/well) and after reached confluence, were treated with crude extracts at concentrations of 100, 250 and 500 µg/mL, dissolved in DMSO (2%). As negative control, DMSO 2% was used and positive control consisted of Triton X-100 (1%). After 24 h incubation, cell viability was determined by reduction of MTT.[18] In this assay, cells were incubated with MTT (0.25 µg/mL) during 4 h; after, supernatant was removed, and DMSO (200 µL) was added. The measure of absorbance at 570 and 630 nm was performed in SpectraMax® and results were obtained by two independent experiments, each one in triplicate.

Extension of cell membrane integrity was determined in supernatant of cultures, using LDH assay, through commercial test kit (Labtest Diagnostica SA, Minas Gerais, Brazil) as standardized by Pranke et al.[19] SHED culture was treated with extracts, as described above. Readings were performed in LabMax 560 equipment (Labtest Diagnostica SA), and cytotoxicity was calculated according to the damage of cell membrane, expressed as a percentage of intracellular LDH release compared to negative control (100%). Independent experiments were performed in triplicate.

Chromatographic and mass spectrometric analysis

Based on biological results, we performed liquid chromatography coupled with photodiode array (PDA) detector (HPLC-PDA) analysis of the most active sample, in order to verify chemical composition and detect active compounds. Separation Module Waters 2695 and ultraviolet (UV) visible Waters 996 PDA detector were employed, data acquisition and integration were managed with Waters Empower software (Waters, Milford, MA, USA). A C18 reversed-phase column (Kromasil, 150 mm × 4.6 mm × 5 µm) was used, operating at a temperature of 24 ± 2°C. Dried methanol crude extract from B. brasiliense was dissolved in methanol (LC grade) to obtain final concentration of 10 µg/mL. Sample was filtered through a 0.45 µm pore size membrane (Millipore, Bedford, USA) before LC system analysis. A linear gradient system was used with mobile phases consisting of a ultrapure water: formic acid (100:2; v/v) mixture (A) and acetonitrile (100; v) (B). Gradient profile was: 0–45 min from 5 to 35% of B, 45–46 min from 35 to 50% of B, 46–47 min from 50 to 100% of B, 47–50 min 100% of B. The flow rate was 0.8 mL/min, and injection volume was 10 µL. Compound detection was performed at 320 nm, and identity was confirmed by comparison with reference substances, injected under the same conditions.

Additional chemical structure information was obtained by mass spectrometry with electrospray ionization (ESI-MS). A mass spectrometer Waters microTOF-Q Micromass (Waters Corp., Milford, MA, USA) was employed, high purity nitrogen was used as nebulizer and auxiliary gas argon as the collision gas. MS/MS spectra was obtained in negative ion mode, with 25 eV collision energy, by direct injection. Data acquisition was performed using Waters MassLynx software (Waters, Milford, MA, USA).[18]

Statistical analysis

Statistical analysis from determination of total phenols, assessment of cell viability and cytotoxicity were evaluated by analysis of variance followed by Bonferroni’s test, using Prism 5.0 software (GraphPad Software, Inc., CA, USA). Significant differences were considered when P < 0.05. Results obtained in antioxidant assays, and enzyme inhibition tests were interpreted using the same program, and IC₅₀ values were obtained after adjustment of experimental data (% inhibition vs. inhibitor concentration) for nonlinear regression curves, through equation described below (Equation 1):

\[ Y = Bottom + \left(\frac{Top - Bottom}{1 + 10^{\left(-\log IC_{50} - X\right)\cdot HillSlope}}\right) \]

Equation 1. Calculation of IC₅₀ value from samples, using nonlinear regression curves obtained by Prism 5.0 software (GraphPad Software, Inc., CA, USA).

RESULTS

Total polyphenols content

Results of total polyphenols content in methanolic extracts of Blechnum species demonstrated B. brasiliense with higher content (9.25 ± 0.35 g GAE/100 g extract), followed by B. occidentale (7.22 ± 0.20 g GAE/100 g extract) and B. binervatum (5.17 ± 0.32 g GAE/100 g extract), with values significantly different between samples (P < 0.05).
Antioxidant assays

Hydroxyl radicals are reactive oxygen species mainly responsible for biological damage and lipid oxidation.\(^{19}\) Blechnum extracts were effective in scavenger hydroxyl radicals, formed by deoxyribose degradation. \(B.\) \(brasiliense\) showed the highest activity in hydroxyl radical stabilization, with \(IC_{50}\) of 112.3 ± 2.61 µg/mL [Table 1]. Against nitric oxide, Blechnum extracts showed low activity, with \(IC_{50}\) values above 500 µg/mL. Hydroxycinnamic acids, used as standards, were less active compared with results for other used methods. In addition, antioxidant activity of extracts was assessed by TBARS assay, which is based on the formation of malondialdehyde, a byproduct of lipid peroxidation. Potential activity was observed for \(B.\) \(brasiliense\) extract with \(IC_{50}\) value of 176.1 ± 1.19 µg/mL [Table 1]. Chlorogenic and caffeic acids presented \(IC_{50}\) values equal to 37.5 ± 1.25 and 55.9 ± 1.22 µM, respectively.

Enzyme inhibition

According to results presented in Table 2, all samples were more active against isoform A of MAO, with pronounced activity for \(B.\) \(brasiliense\) extract (\(IC_{50}\) 72.7 ± 1.09 µg/mL). In addition, \(B.\) \(brasiliense\) showed a good selectivity index, since its MAO-A inhibition was 3.92 times higher than MAO-B modulation (\(IC_{50}\) 285.2 ± 1.03 µg/mL). The irreversible MAO-A inhibitor, clorgyline, showed \(IC_{50}\) value of 0.009 ± 0.0002 µM and selective MAO-B inhibitor, pargyline, presented \(IC_{50}\) of 0.207 ± 0.0112 µM.

All Blechnum extracts presented \(IC_{50}\) values above of 500 µg/mL, in both AChE and BChE inhibition, being considered inactive against cholinesterase. Physostigmine, a positive control, exhibited strong inhibitory activity with \(IC_{50}\) values of 0.0165 ± 0.0019 µM (AChE) and 0.0718 ± 0.0018 µM (BChE).

Toxicity evaluation in rodents cells

Cell toxicity was not observed for Blechnum samples, tested by measuring the release of LDH. At 1 mg/mL, results were similar to negative control (cells nontreated), indicating the feasibility of membrane from PMN [Figure 1]. On the other hand, Triton X-100 (1%) showed strong cell membrane damage.

Assessment of viability and cytotoxicity on stem cells

This study also evaluated effects of Blechnum extracts on stem cells viability, through MTT assay [Figure 2a]. Characterization of mesenchymal stem cells was performed according to established criteria by the International Society for Cellular Therapy, showing typical fibroblasts morphology. Cells presented plastic adherence and expressed positivity (≥95%) for CD73, CD105, and CD90. In addition, cells did not express (≤2%) surface markers: CD34, CD45, CD14, CD11b and HLA-DR and were differentiated into osteoblasts, adipocytes, and condroblastos in vitro.\(^{19}\)

Added of extracts, (100-500 µg/mL), cells maintained viability, with values comparable to nontreated group \((P < 0.05)\). Triton X-100 (1%) showed extensive cell death [Figure 2a].

In agreement with MTT results, cells treated with Blechnum extracts by 24 h showed no cytotoxic effect by LDH release method [Figure 2b]. Triton X-100 (1%) showed significant increase in LDH leakage of treated cells. Samples demonstrated similar profiles and were comparable to negative control \((P < 0.05)\). In summary, the results showed Blechnum extracts are capable of maintaining stem cells viability, and no cytotoxic effects were observed at concentrations of 100–500 µg/mL.

Table 1: Half maximal inhibitory concentration values of Blechnum extracts, chlorogenic and caffeic acid standards, in antioxidant experiments

| Assay               | \(IC_{50}^a\) (µg/mL) | \(IC_{50}^b\) (µM) | \(IC_{50}^c\) (µg/mL) | \(IC_{50}^d\) (µM) |
|---------------------|-----------------------|-------------------|-----------------------|-------------------|
| Hydroxyl radical    | 311.1±1.66            | 112.3±2.61        | 198.2±1.86            | 45.6±1.52         |
| Nitric oxide radical| >500                  | >500              | >500                  | 159.8±1.44        |
| TBARS               | 489.5±2.98            | 176.1±1.19        | 455.7±2.56            | 37.5±1.25         |

\(^a\)Thiobarbituric acid reactive substances assay; \(^b\)Chlorogenic acid; \(^c\)Caffeic acid; \(^d\)Values expressed as mean±standard deviation. 

Table 2: Enzyme inhibition of Blechnum extracts against monoamine oxidases and cholinesterase enzymes

| Sample            | MAO-A   | MAO-B   | SI\(^a\) | AChE  | BChE  |
|-------------------|---------|---------|----------|-------|-------|
| \(IC_{50}^a\) (µg/mL) |         |         |          |       |       |
| \(B.\) \(binnervatum\) | 165.2±1.10 | 499.7±2.24 | 3.02     | >500   | >500   |
| \(B.\) \(brasiliense\) | 72.7±1.09  | 285.2±1.03 | 3.92     | >500   | >500   |
| \(B.\) \(occidentale\) | 130.8±1.05 | 421.5±1.01 | 3.22     | >500   | >500   |
| \(IC_{50}^a\) (µM) |         |         |          |       |       |
| Clorgyline        | 0.009±0.0002 | NT**    | -        | NT**   | NT**   |
| Pargyline         | NT**    | 0.207±0.0112 | -        | NT**   | NT**   |
| Physostigmine     | NT**    | 0.0165±0.0019 | 0.0718±0.0018 |       |       |

\(^a\)Selectivity index \((IC_{50}^b/IC_{50}^a)\); \(^b\)Values expressed as mean±standard deviation; \(^c\)NT: Not-tested; \(IC_{50}^a\): Half maximal inhibitory concentration; MAO: Monoamine oxidases; AChE: Acetylcholinesterase; BChE: Butryrylcholinesterase
Chromatographic and mass spectrometric analysis

Analysis by HPLC-PDA was conducted with B. brasiliense extract, most active sample in antioxidant assays and inhibition of MAO. This sample presented four detectable peaks, being two of them, the majorities, with retention times of 13.73 and 29.76 min [Figure 3]. Compounds showed maxima UV absorption characteristic of hydroxycinnamic acids.[20] Comparing with standard reference, compound in 13.73 min was characterized as chlorogenic acid.

In addition, compound with retention time of 29.76 min was analyzed by mass spectroscopy in negative mode. The substance presents C_{18}H_{16}O_{8} molecular formula and molecular mass of 360 g/mol. Pseudomolecular ion (m/z 357 [359-H_2^+]_-, was fragmented by MS/MS [Figure 3d], generating characteristic product-ions: m/z 197, m/z 179 and base peak m/z 161.[21-23] Compiled data conducted to the identification of rosmarinic acid in B. brasiliense extract.

DISCUSSION

For some species of the Blechnum genus, Bohm[24] showed the presence of rosmarinic and chlorogenic acids in ethanol extracts, using thin layer chromatography analysis. Blechnic acid was also observed in B. spicant extract. According to Farias et al.,[25] the crude extract of B. brasiliense presented 2.352 g GAE/100 g extract. Comparing with our results, we observed the higher content of total polyphenols in this species.

Antioxidant action mechanism of polyphenols, evaluated by hydroxyl scavenging assay, is usually attributed to the scavenging ability of these compounds. However, many studies have been shown that these phenols do not only block the degradation of deoxyribose by stabilization of hydroxyl radicals but also act as antioxidants via formation of complexes with iron. Thus, iron cannot participate in Fenton’s reaction, which is responsible to triggering generation of free radicals through oxidation of organic substrates.[26,27] With regards to the NO• radical, literature reports as one of reactive nitrogen species that contributes to the development of several diseases, including inflammation and neurodegenerative disorders.[28] TBARS was performed since therapeutic strategies capable to modulate lipid peroxidation may be promising in the prevention of several disorders.[29] Thus, substances with free radical scavenging properties, also lipid peroxidation inhibition are valuable for new drugs development. Farias et al.[25] previously demonstrated inhibition of DPPH formation by B. brasiliense, which IC_{50} values of crude extract and ethyl acetate fraction were 4.14 and 1.43 mg/mL, respectively. Significant differences observed
can be explained by chemical nature of samples and also by employed assay. Distinct methodologies present limitations and applicability, mainly due to complex nature of phytochemicals and their interactions. Thus, multiple antioxidant methodologies become an indispensible approach to evaluate complex phenol mixtures, which are involved in scavenging free radicals and inhibiting lipid peroxidation in tissues.[30]

MAO activity was determined, in this study, by fluorometric assay proposed by Novaroli et al.[31] employing kynurenine as substrate. This substance, after oxidation, produces 3-hydroxykynoline, a fluorescent compound, easily detected and quantified.[1,12] MAO inhibitors are considered key prototypes for drug development, applied for neurodegenerative diseases and also depression and anxiety treatment.[32] Mazzio et al.[33] ranked plant crude extracts regarding their MAO-B inhibition, according to levels: Very strong (IC50 <70 µg/mL), strong (IC50 <200 µg/mL), moderately strong (IC50 >200 <400 µg/mL), moderate (IC50 >400 <700 µg/mL) and weak (IC50 = 700 µg/mL). Considering this classification, B. brasilienne extract showed moderately strong inhibition (IC50: 285.2 µg/mL), B. binervatum and B. occidentale extracts presented moderate inhibition of MAO-B (IC50: 499.7 and 421.5 µg/mL, respectively).

Covering the proposal of multiple targets search, the inhibitory activity of AChE was assessed, which is characterized as the main strategy employed in Alzheimer’s disease treatment.[34] This activity promotes cognition improvement in patients. Besides AChE, BChE is also responsible for inactivation of acetylcholine in brain tissue.[35] Our results showed no effects of Blechnum extracts against cholinesterases, highlighting the modulation of targets related to Parkinson disease.[4]

Some of the available MAO modulator drugs in therapeutics of depression and Parkinson diseases present many side effects and their physiological activity can persist for up to 2-3 weeks.[36] Thus, the search for novel inhibitors with few adverse effects is strongly encouraged. In this sense, natural compounds from different classes are being studied as MAO inhibitors, as well as, multifunctional substances on targets related to neurodegeneration. Flavonoids and coumarins have already demonstrated strong MAO inhibition, being agenin the most active compound, among the tested (IC50: 1 µmol).[37]

Generally, bicyclic compounds, including coumarin and conjugated hydroxycinnamic acids, are known to present considerable inhibitory activity against both MAO-A and MAO-B enzymes.[19] Gallic acid and derivatives, isolated from Liquidambar formosana fruits, showed strong MAO inhibition, with IC50 values lower than 2 µg/mL.[38] Recent studies showed that infusions from green tea and citrus peels presented MAO inhibitory activity,[40] being linked to their phenolic content, due to its structural similarity to synthetic inhibitors.[41] Phenolic compounds are well known antioxidant and anti-inflammatory substances and these properties, added to their MAO inhibition, makes them interesting neuroprotective compounds designed to drug discovery.

Our previous research studies assessed 11 species of ferns from Brazil, regarding its antioxidant and anti-inflammatory potential (10 µg/mL) and also inhibitory capacity against MAO-A and MAO-B enzymes (100 µ/mL). Asplenium serra, Lastroepis amplissima, and Cyathea dichromatolepis showed potential antioxidant activity by total reactive antioxidant potential method (TRAP). MAO-A inhibition was predominantly observed for Dydymochlaena truncatula, Alsophila setosa, Cyathea phalerata, and Cyathea delgadii, whose inhibition values ranged from 70.32% to 82.61% at the concentration tested. No toxic effects were observed to PMN cells, when incubated with fern species, at 100 µg/mL.[41]

In addition to the toxic assay using animal cells, our studies have evaluated effects of Blechnum extracts on human stem cells, considering their high rate of proliferation and capacity of differentiation in several cell types, including neurons. Thus, this cells can predict toxicity of plant extracts and isolated molecules in specific differentiated cells.[42-44] Furthermore, human exfoliated deciduous teeth allows obtaining stem cells with minimal invasive damage to donor since teeth are discarded after natural replacement process.[45] Human cells usage enables more sensitivity design to predict toxicity and contribute to animals replacement.[46]

According with HPLC-DAD-MS analysis, the most active sample presented two hydroxycinnamic acids in extract composition, characterized as chlorogenic and rosmarinic acid. Studies have reported effects of chlorogenic acid on H2O2 induced damages[47] as well as its protective ability against neurotoxicity promoted by oxidative stress and reduction of lipid peroxidation in brain structures.[48] For rosmarinic acid, significant antioxidant activity has been reported,[49] being responsible for cytoprotective effects of this molecule.[50] Moreover, this substance presented protective ability on human dopaminergic cell line (SH-SYSY).[51] Thus, our findings point to new safe sources of neuroactive compounds, capable of modulate targets related to neurodegenerative disorders, stimulating the development of new scaffolds to compose therapeutic strategies to combat these pathologies.

CONCLUSION

Our results highlight the importance of fern species studies, especially from Blechnum genus, which few reports are described in the literature. This is the first report assessing Blechnum species for antioxidant activity and enzymatic selected methods demonstrating safety of these plants in animal and human stem cells. Taken together, data indicate B. brasilienne extract with most relevant results, which can be related to higher levels of total phenols and also to the presence of chlorogenic and rosmarinic acids. These compounds contributed to the pronounced capacity of stabilize hydroxyl radical and inhibit lipid peroxidation. In addition, B. brasilienne extract displayed strong and selective inhibition of MAO-A enzyme, showing no toxic effects and maintaining viability of rodents and human stem cells. Compiled data point to these fern species as important sources of natural and safe compounds, capable to prevent damage caused by oxidants and lipid peroxidation, which is associated with numerous pathologies. The lack of AChE and BChE enzyme activities also shows selectivity, indicating these plant species as inhibitors of specific biochemical targets related degenerative disorders, such as Parkinson’s disease.

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Conflicts of interest

There are no conflicts of interest.

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