Molecular evaluation of anti-inflammatory activity of phenolic lipid extracted from cashew nut shell liquid (CNSL)

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Abstract

Background: Anacardium occidentale L phenolic lipid (LDT11) is used in traditional medicine as anti-inflammatory, astringent, antidiarrheal, anti-asthmatic and depurative. Phenolic derivatives, such as anacardic acid, extracted from cashew nut shell liquid (CNSL) have demonstrated biological and pharmacological properties, and its profile makes it a candidate for the development of new anti-inflammatory agents. The objective of the present study was to evaluate the anti-inflammatory profile of a derivative, synthesized from LDT11, on an in vitro cellular model.

Methods: Organic synthesis of the phenolic derivative of CNSL that results in the hemi-synthetic compound LDT11. The cytotoxicity of the planned compound, LDT11, was analyzed in murine macrophages cell line, RAW264.7. The cells were previously treated with LDT11, and then, the inflammation was stimulated with lipopolysaccharide (LPS), in intervals of 6 h and 24 h. The analysis of the gene expression of inflammatory markers (TNFα, iNOS, COX-2, NF-κB, IL-1β and IL-6), nitric oxide (NO) dosage, and cytokine IL-6 were realized.

Results: The results showed that the phenolic derivative, LDT11, influenced the modulatory gene expression. The relative gene transcripts quantification demonstrated that the LDT11 disclosed an immunoprotective effect against inflammation by decreasing genes expression when compared with cells stimulated with LPS in the control group. The NO and IL-6 dosages confirmed the results found in gene expression.

Discussion: The present study evaluated the immunoprotective effect of LDT11. In addition to a significant reduction in the expression of inflammatory genes, LDT11 also had a faster and superior anti-inflammatory action than the commercial products, and its response was already evident in the test carried out six hours after the treatment of the cells.

Conclusion: This study demonstrated LDT11 is potentially valuable as a rapid immunoprotective anti-inflammatory agent. Treatment with LDT11 decreased the gene expression of inflammatory markers, and the NO, and IL-6 production. When compared to commercial drugs, LDT11 showed a superior anti-inflammatory action.

Keywords: Fenolic lipid, Anacardic acid, Gene expression, Cashew nut shell liquid, CNSL, Anti-inflammatory activity
Background

Cashew nut shells are considered a residue of the cashew nut processing by the agribusiness. However, for those in search of useful substances from renewable sources it has proven to be valuable bio-based material. The cashew nut shell contains a liquid (CNSL) that is a caustic, viscous oil comprising 25% of the fruit weight in natura. CNSL extracted by the cashew processing industry, which separates the almond and the oil, is one of the most abundant sources of non-isoprenoid phenolic lipid, such as anacardic acid, cardol, cardanol and methylcardol (Fig. 1) [1]. The CNSL components, in addition to an aromatic nucleus and several distinct functional groups, has an acyclic side chain containing multiple instabilities in the aliphatic chain, which results in an amphipathic behavior. From a synthetic point of view, CNSL properties characterize it as an extremely versatile material [2].

Nations in South America, Africa, and Asia have for decades use *Anacardium occidentale* L. phenolic lipid, extract from CNSL, in traditional medicine [3–5]. In folk medicine, CNSL is used as anti-inflammatory, astringent, antidiarrheal, anti-asthmatic, depurative and tonic medication. It is also used as diabetes medication [4, 6, 7] and wounds and wart treatment [8–10].

Past research has confirmed that phenolic and semi-synthetic derivatives of CNSL have biological properties [11], such as antibacterial, anti-inflammatory [12–14], and antioxidant activity [15]. Additionally, pharmacological properties included enzymatic inhibition [16, 17] and antiproliferative activity [16, 18]. A recent review by Hemshekhar et al. [19] reinforced anacardic acid multi-target pharmacological profile and its potentiality for the development of new anti-inflammatory drugs.

Inflammation is part of the complex biological response by body tissue to harmful stimuli, caused by infections, injuries or trauma. It is a complicated process regulated by several pro-inflammatory mediators, such as TNF-α, COX-2, iNOS, NF-κB, IL-1β, and IL-6 [20]. The rapid release of pro-inflammatory cytokines by activated macrophages plays a crucial role in triggering local immune response [21]. However, excessive production of inflammatory mediators may be more damaging than the event that triggered the immune response and may be associated with autoimmune diseases, diabetes, sepsis, diffuse intravascular coagulation, tissue injury, hypotension, and death [22]. The inhibition of these inflammatory mediators employing pharmacological modulators has been used as an effective therapeutic strategy to reduce inflammatory reactions [23].

Considering that biosynthetic molecules derived from CNSL have been tested in cellular models in vitro [24, 25], the present work proposes to evaluate the anti-inflammatory profile of *Anacardium occidentale* L. phenolic lipid (LDT11, Fig. 2), in the cellular model. Results of this analysis may offer alternative therapeutic strategies for the treatment of inflammation.

Methods

The production of inflammatory mediators was analyzed on RAW 264.7- TIB-71 murine macrophages cell culture (American Type Culture Collection - ATCC), previously treated with LDT11. Cells were purchased from the cell bank of the Adolf Lutz Institute (São Paulo, Brazil), and cultured according to the ATCC criteria.

Synthesis and characterization of LDT11 as a potential anti-inflammatory agent

LDT11 is a derivative designed from cashew nut shell liquid (CNSL) phenolic lipid. Compounds from library of the Laboratory of Development of Therapeutic Innovations (LDT), part of the University of Brasilia (Brazil) were used in this research. LDT11 synthesis was performed as follows: to a solution of the mixture of anacardic acids (5 g, 14.5 mmol for average molecular wt 344) in ethanol (50.0 mL) was added 10% palladium-carbon (0.2 g) and shaken in a Parr apparatus (Parr Instrument Company©, Moline, IL, USA), under hydrogen atmosphere (4 atm, 60 psi) at room temperature. After six hours, the mixture was filtered and the solvent was evaporated under reduced pressure. The residue was recrystallized from hexane to afford a saturated anacardic acid (LDT11) as a white solid (4,55 g, 90%, mp 81 °C–83 °C, Rf 0.48 – Hex:AcOEt 4:1). IR (KBr) ν_{max} cm^{-1}: 3326 (ν_{OH}); 2954 (ν_{asCH3}); 2920

![Fig. 1 Non-isoprenoid phenolic lipid constituent of the CNSL](image-url)
steroid dexamethasone (DEX) (Decadron, Aché Laboratórios Farmacêuticos S.A, Brazil). LDT11 has a similar chemical structure to LDT11, and corticosteroid salicylic acid (ASA) (Sedalive, Vitamedic, Brazil), which activity was compared to two commercial drugs: acetylsalicylic acid (ASA) (Sedalive, Vitamedic, Brazil), which protective tests for inflammation in vitro. LDT11 was used for immunoprotective tests for inflammation in vitro. LDT11’s biological activity was compared to two commercial drugs: acetylsalicylic acid (ASA) (Sedalive, Vitamedic, Brazil), which has a similar chemical structure to LDT11, and corticosteroid dexamethasone (DEX) (Decadron, Aché Laboratórios Farmacêuticos S.A, Brazil).

Quantitation of viable cell number - WST-8 assay
The cytotoxicity of the macrophages treated with synthetic phenolic derivatives was determined using the WST-8 assay (Cell Counting kit-8, Sigma-Aldrich, St. Louis, MO, USA). Cells were grown in complete Dulbecco’s Modified Eagle’s Medium (DMEM) at a concentration of 1 × 10^5 in 96-well plates and incubated at 37 °C in an atmosphere of 5% CO₂ for 48 h. The culture medium was subsequently exchanged for 100 μL of LDT11, the medium was discarded, the cells were washed twice with phosphate buffered saline solution (1X PBS, pH 7.4), and 100 μL of DMEM supplemented, and 50 μg/mL of neutral red was added to the wells. The plate was incubated under the conditions described in the item 2.2. The medium was then discarded, and the cells were washed five times with 1X PBS to remove the excess of dye; which was followed by the addition of 100 μL of alcohol-acid solution (50% ethanol, 1% acetic acid and 49% distilled water) to each well to fix the neutral red to the cells. The plate was shaken for 10 min, and the absorbance of the samples was read in a TP-Reader microplate spectrophotometer (Thermoplate, Palm City, FL, USA) using a 450 nm wavelength filter.

Neutral red uptake assay
The Neutral red uptake assay was performed following the protocol described by Tanner et al. [26], with some modifications. The assay was carried out under the same conditions as the WST-8 assay. After 48 h of incubation with the LDT11, the medium was discarded, the cells were washed twice with phosphate buffered saline solution (1X PBS, pH 7.4), and 100 μL of DMEM supplemented, and 50 μg/mL of neutral red was added to the wells. The plate was incubated under the conditions described in the item 2.2. The medium was then discarded, and the cells were washed five times with 1X PBS to remove the excess of dye; which was followed by the addition of 100 μL of alcohol-acid solution (50% ethanol, 1% acetic acid and 49% distilled water) to each well to fix the neutral red to the cells. The plate was shaken for 10 min, and the absorbance of the samples was read in a TP-Reader microplate spectrophotometer (Thermoplate, Palm City, FL, USA) using a 492 nm filter. The results were expressed as a percentage, the value obtained for the positive control (untreated cells), being considered as 100% viability. The equation used was: viability (%) = (number of viable cells / total number of untreated cells) × 100.

Analysis of gene expression
To evaluate the influence of LDT11 on RAW 264.7 gene expression, after stimulation with LPS, quantitative real-time polymerase chain reaction (qPCR) was performed. The RNA from cells was extracted, purified and quantified, and the resulting complementary DNA was prepared for the qPCR reaction, as follow: RAW 264.7 cells were cultured in six well plates at a concentration of 5 × 10^5 cells/well, each containing 3 mL 10% complete DMEM medium. The plates were incubated (approximately 24 h) until an estimated confluence of 90% was obtained. The medium was discarded, and 3 mL of colorless DMEM medium without FBS supplementation

Quantitation of viable cell number - WST-8 assay
The cytotoxicity of the macrophages treated with synthetic phenolic derivatives was determined using the WST-8 assay (Cell Counting kit-8, Sigma-Aldrich, St. Louis, MO, USA). Cells were grown in complete Dulbecco’s Modified Eagle’s Medium (DMEM) at a concentration of 1 × 10^5 in 96-well plates and incubated at 37 °C in an atmosphere of 5% CO₂ for 48 h. The culture medium was subsequently exchanged for 100 μL of DMEM supplemented with 5% colorless fetal bovine serum (FBS), and 100 μL of LDT11 was added to the wells at concentrations of 25 μM, 50 μM, 75 μM, 100 μM, 125 μM, 150 μM, totaling a volume of 200 μL per well. After that, the plate was once more incubated under the previously described conditions.

The assay was performed using samples and controls in triplicate. After 48 h, the medium was discarded, and 100 μL of colorless DMEM, supplemented with 5% FBS was added, followed by the addition of 10 μL of WST-8 to each well. Non-stimulated cells were used as positive controls. Cell death control was performed using cells treated with 10 μL of 1% Triton-X. After an incubation of four hours with WST-8, the absorbance of the samples was measured in a TP-Reader microplate spectrophotometer (Thermoplate, Palm City, FL, USA) using a 450 nm wavelength filter.
was added in order to restrain the cells growth rate. The immune-protective properties of LDT11 were analyzed in: (a) untreated cells (NT); (b) cells stimulated exclusively with LPS; (c) cells treated with LDT11, and subsequently exposed to LPS; (d) cells treated with ASA, and then exposed to LPS; (e) cells treated with DEX, and exposed to LPS.

**Extraction, purification, quantification of RNA and cDNA synthesis**

For the extraction and purification of RNA, Direct-zol ™ RNA Miniprep Kit (R2051, Zymo Research, Orange, CA, USA), was used following the manufacturer’s recommendations. The quantification of extracted RNA was determined by using the NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA). The cDNA synthesis was performed using High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Waltham, MA, USA), according to manufacturer’s instructions.

**Gene analysis and characterization**

Genes involved in the inflammatory response were selected to test the biological activity of LDT11. Seven primers synthesized by Integrated DNA Technologies (Skokie, IL, USA) were used, as described in Table 1. The design of the primer pairs was performed using the Primer Express Program (Applied Biosystems, Waltham, MA, USA) based on sequences obtained from the Mouse Transcriptome Database (http://www.informatics.jax.org).

**Quantitative real-time polymerase chain reaction**

The Quantitative Real-Time Polymerase Chain Reaction (qPCR) was performed in triplicates, using SYBR Green system (Absolute qPCR SYBR Green Rox Mix - Thermo Fisher Scientific Inc., Vilnius, Lithuania, USA) in a solution containing 50 ng of cDNA, 5 pmol of each primer forward and reverse and QSP of ultrapure water, totaling 20 μL. Amplification assays were performed on a StepOnePlus Real-Time PCR System (Applied Biosystems, Carlsbad, CA, USA) under the following conditions: initial denaturation temperature at 95 °C for 10 min, 40 cycles of denaturation at 95 °C for 15 s, annealing and extension at 60 °C for 1 min and 72 °C for 30 s. All reactions were performed three times for each gene, and GAPDH gene was used both as endogenous control and normalizing gene. Water was used as negative control substituting cDNA.

The result, expressed in CT value, refers to the number of cycles in qPCR required for the fluorescent signal to reach the threshold detection. The analysis of the gene expression of the inflammatory markers was obtained by the relative quantification of their transcripts by the Delta-Delta Ct (ΔΔCt) method, which allows a relative comparison with the group that did not receive treatment (NT) and cells stimulated with LPS. The melting curve was used as quality control of amplification products.

For the data analysis, the values found for the control group, LPS-stimulated cells, were considered 100% inflamed. The other tests, with their respective treatments (LDT11, ASA and DEX) were analyzed comparing with the inflamed group.

**Nitric oxide quantification**

For the analysis of Nitric Oxide (NO), was used the Griess method, by adding 100 μL of Griess reagent (1% [w/v] sulfanilamide, in a 5% phosphoric acid, and 0.1% [w / v] N-1- naphthyl-ethylenediamide dihydrochloride (NEED) in water). Samples of culture supernatants were analyzed by microplate reader (TP-Reader microplate spectrophotometer, Thermoplate, Palm City, FL, USA), using the spectrum 450 nm, and the results expressed in

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**Table 1** RAW 264.7 macrophages genes analyzed in the study. Primers Description Sequences Mt.*

| Primers | Description | Sequences | Mt* |
|---------|-------------|-----------|-----|
| GAPDH   | Glyceraldehyde-3-phosphate dehydrogenase | 5’-CCCGTGCTGATATGCG-3’ | 85,22 |
| COX-2   | Cyclooxygenase 2 | 5’-TCAGTGACGAGGCTTTC-3’ | 83,05 |
| IL-1β   | Interleukin 1 beta | 5’-TAATGGCAGCGATATGC-3’ | 54,50 |
| IL-6    | Interleukin 6 | 5’-CAGTACGGCGATATGC-3’ | 73,48 |
| iNOS    | Induced nitric oxide synthase | 5’-GCCGAGCTGACGAGCTT-3’ | 77,81 |
| NF-κB   | Nuclear Factor kappa b | 5’-ACAGGCGTGATATGC-3’ | 77,81 |
| TNF-α   | Tumor necrosis factor alpha | 5’-TCCTCTCATTCGCTTG-3’ | 81,70 |

*Mt Melting temperature (°C)
μmol/L of NO - comparing the optical density (OD) obtained with a standard curve of NO - ranging from 1.56 μM to 100.0 μM.

IL-6 quantification
The quantification of Interleukin-6 (IL-6) was performed by applying the competitive immunoenzymatic assay, as established in the manufacturer’s kit (#27768 Mouse IL-6 Kit – Immuno-Biological Laboratories Co., Ltda, Hamburg, Germany). The same treatment used in the gene expression assessment was applied in this assay, using the supernatants of RAW 264.7 cells treated with the synthetic phenolic derivative LDT11 at 50 μM concentration.

Statistical analysis
Statistical analysis was performed using the Analysis of Variance (ANOVA) test, followed by Post-Hoc tests (Bonferroni, Dunnnett and t-test) when applicable, according to GraphPad Prism version 4.0 (GraphPad Software, San Diego, USA). Statistically significant differences were considered when $p < 0.05$.

Results
Cytotoxicity analysis and quantitation of viable cell number
The determination of the cytotoxic effect of LDT11 by the WST-8 and Neutral Red methods was performed using the following concentrations: 25 μM, 50 μM, 75 μM, 100 μM, 125 μM, and 150 μM.

WST-8 assay
LDT11 showed approximately 100% cell viability in a concentration of 25 μM. The cell viability declined to 90% with a LDT11 concentration of 50 μM. Concentrations equal or over 75 μM ensued a cell viability equal or lower than 60% and were considered cytotoxic (Fig. 3).

Neutral red uptake assay
Neutral Red Uptake Assay was performed to confirm the cell viability obtained by the WST-8 assay on the RAW264.7 cell line, using similar LDT11 concentrations. The assay showed results similar to those found by the WST-8 method (Fig. 4).

From the results of the cell viability assays, toxic concentrations of LDT11 were excluded from the study. The standardized concentration for the other tests was 50 μM. For further comparison, the following assays with ASA and DEX were also made using the same concentration established for LDT11, 50 μM.

Modulation and relative quantification of transcripts of inflammatory genes
Gene modulation of the inflammatory process was analyzed after interaction with the CNSL derivative, LDT11. Gene expression was evaluated by the relative quantification of TNF-α, COX-2, iNOS, NF-kB, IL-1β and IL-6 genes in RAW264.7 cells. Inflammation was induced with 1 μg/mL LPS, both in control cells and in LDT11 treated cells. The analysis of the results was carried out 6 and 24 h post- treatment.

As seen in Fig. 5, the gene expression of TNF-α, six hours after interaction with LDT11, showed an eight-fold decrease (88%) in comparison with the gene expression disclosed by the control cells (LPS). The results obtained from the interaction with ASA and dexamethasone (DEX) were respectively one and a half times lower (33%) for ASA and two times lower (50%) for DEX (Fig. 5a). The results obtained 24 h after treatment were similar with those obtained after six hours. Both had a decrease in gene expression in all treatments analyzed. Cells treated with LDT11 disclosed 10 times (91%) less gene expression than the positive control, while cells treated with ASA and DEX respectively showed a gene expression two and three times lower (55 and 64%) than control cells, Fig. 5b.
COX-2 was the second gene analyzed (Fig. 6). After six hours of treatment with LTD11, COX-2 gene expression decreased more than five-fold (81%) when compared to control cells (Fig. 6a). Cells treated with ASA did not show a significant difference, and those treated with DEX showed a four-fold decrease (74%) in gene expression when compared to control cells. After 24 h of interaction with LTD11 (Fig. 6b), the gene expression disclosed a six-fold decrease (84%) in treated cells than in control cells. The reduction in gene expression caused by both ASA and DEX was approximately 2-fold higher (30%) than that observed in control cells.

After six hours of LTD11 treatment, the expression of the iNOS gene, (Fig. 7) showed a 200 times decrease (100%) comparing with control cells. Treatment with ASA resulted 20-fold (99%) decreased expression, while cells treated with DEX showed a 60-fold (98%) gene suppression (Fig. 7a). After 24 h (Fig. 7b), both LTD11 and DEX treated cells showed approximately a 2-fold decreased gene expression (50%) while a threefold decrease (68%) was observed in ASA-treated cells.

The action of LTD11 on the NF-κB gene expression can be seen in Fig. 8. In comparison with control group, cells treated with LTD11 resulted an eight-fold (88%) decrease in NF-κB gene expression. The treatment of the cells with ASA and DEX produced respectively a decrease in gene expression equivalent to 1.5-fold (33%) for the first and two-fold (50%) for the second drug (Fig. 8a). In the treatment performed after 24 h (Fig. 8b), both LTD11 and ASA treated cells showed a decrease of about two-fold (55%) in the NF-κB gene expression. On the other hand, cells treated with DEX failed to show any significant difference when compared to the control cells.

The IL-1β gene also showed decreased expression in most of the assays (Fig. 9). After a six-hour interaction with LTD11, IL-1β gene expression decreased more than 14-fold (93%). Treatment with ASA did not cause a significant decrease in its expression whereas DEX led to a three-fold reduction (69%) when compared to control cells (Fig. 9a). After 24 h (Fig. 9b), when compared with control cells, those treated with LTD11 showed an approximately six-fold decrease (83%) in gene expression, whereas ASA and DEX caused respectively a two-fold (39%) and five-fold (75%) decline.

After six hours of treatment with LTD11 there was a 65-fold (98%) decrease in the gene expression of IL-6, in
comparison with the control cells. Gene expression of IL-6 decreased respectively 8-fold (88%) in ASA-treated cells and 27-fold (96%) in DEX-treated cells (Fig. 10a). After 24 h (Fig. 10b) the gene expression decreased 5-fold (79%) in the cells treated with LDT11 while the decrease was respectively two-fold (45%) in the cells treated with ASA and three-fold (65%) in the cells treated with DEX.

Assessment of nitric oxide and cytosine IL-6

Nitric oxide, resulting from the presence of oxygen and nitrogen reactive metabolites produced during the inflammatory process, causes an increase in the oxidative stress of the cells. The presence of NO in the supernatant of RAW 264.7 cells was measured to confirm the influence of LDT11 on anti-inflammatory activity. The following assays were performed with the same timing (six and 24 h) of the previous analysis. The LDT11 was able to protect RAW264.7 cells against oxidative stress by reducing NO production by about eight times (95%) after six hours, and about 15 times (100%) after 24 h (Fig. 11).

Additionally, to confirm the results obtained in the analysis of gene expression, the cytokine IL-6 was also assayed. The pre-treatment of the cells with LTD11 resulted in a significant protective effect against inflammation, reducing IL-6 production by more than 1700-fold (76%) after six hours (Fig. 12a) and more than 1400-fold (60%) after 24 h (Fig. 12b). Cells treated with ASA reduced IL-6 production respectively by more than 120-fold (52%), after six hours and by more than 400 (17%) after 24 h. Treatment with DEX reduced the IL-6 production at about 1000-fold (42%) after six hours and at about 1400-fold (60%) after 24 h.

Discussion

The present study evaluated the immunoprotective effect of LDT11; a compound synthesized from the anacardic acid extracted from the CNSL. This evaluation was performed through the quantification of gene transcripts involved in the inflammatory response, and the measurement of NO and IL-6 production in RAW 264.7 murine macrophages. The murine macrophages culture was firstly treated with LTD11 and posteriorly stimulated with LPS that mimicked the inflammatory response and induced high production cytokine and oxidative stress [27–29]. Overproduction of pro-inflammatory mediators such as TNF-α, COX-2, iNOS, NF-κB, IL-1β and NO has been implicated in several
inflammatory diseases [27, 29, 30]. Therefore, an agent that prevents the release of these mediators or downregulates the expression of these cytokines may be an valuable therapeutic strategy for preventing inflammatory reaction [27, 28], turning LDT11 a potential candidate for the formulation of new drugs.

In this study, two different colorimetric methods, the WST-8 assay, and the neutral red uptake test, were performed to evaluate if the LDT11 used in the experiments would show some degree of cytotoxicity. The analysis of its in vitro cytotoxicity allowed the determination of the concentrations needed to obtain the best performance of its biological activity without compromising the cellular viability. Viability and cytotoxicity results were similar for both tests at all concentrations used (25 \( \mu \)M, 50 \( \mu \)M, 75 \( \mu \)M, 100 \( \mu \)M, 125 \( \mu \)M, and 150 \( \mu \)M). At concentrations of 25 \( \mu \)M and 50 \( \mu \)M, cell viability was greater than 80%. However, the increase of the concentrations to 75 \( \mu \)M, 100 \( \mu \)M, 125 \( \mu \)M and 150 \( \mu \)M caused more than 60% decrease in cell viability, being consequently considered cytotoxic. Others studies found that the assay shows cytotoxic activity when cell death exceeds 50% [18, 31]. In the present research values above 60% were considered citotoxic. Therefore, the concentration chosen to continue the experiments of gene expression analysis was 50 \( \mu \)M of LDT11.

The TNF-\( \alpha \) gene has the main physiological effect of promoting the inflammatory immune response through the recruitment and activation of neutrophils and monocytes. Consequently, TNF-\( \alpha \) is responsible for a several effects on the body, promoting vasodilation and acting on endothelial cells, stimulating the secretion of a group of cytokines that have chemotactic action on leukocytes. TNF-\( \alpha \) is also the cytokine responsible for septic shock, in addition to inhibiting the appetite and inducing fever through the release of the adrenocorticotrophic hormone (ACTH) [30, 32]. Additionally, it is also a major inducer of the transcription factor NF-\( \kappa \)B, and degrades the inhibitor of NF-\( \kappa \)B (I\( \kappa \)B). Degradation of I\( \kappa \)B allows NF-\( \kappa \)B to be translocated to the nucleus. [21, 33]. Pretreatment with LDT11 showed suppression of the TNF-\( \alpha \) gene in LPS-stimulated cells greater than the suppression observed in cells treated with commercial anti-inflammatory drugs. Consequently, a similar suppression was observed in NF-\( \kappa \)B and TNF-\( \alpha \), being greater in LDT11-treated cells than in the cells treated with commercial drugs.

NF-\( \kappa \)B is responsible for the transcription of innumerable genes related to pro-inflammatory activity, such as IL-1, IL-6, iNOS, COX-2 [34–36]. Consequently, a decrease in the gene expression of these inflammatory mediators was also observed. Compared with LPS-stimulated cells, the
COX-2 and IL-1β genes showed more than 80% gene suppression after treatment with LDT11. The suppression of iNOS and IL-6 genes was practically total in LDT11 pre-treated cells. The evaluation of the treatment after 24 h showed that the relative amount of iNOS decreased in all the tests performed, including in the LPS treated cells. Consequently, the gene suppression was less expressive than that observed in experiments conducted after six hours. Gene expression of iNOS after 24 h was the only outcome in which treatment with LDT11 was inferior to the results obtained with the commercial drugs. Although, treatment with LDT11 was also capable of reducing the iNOS gene expression when compared to LPS treated cells.

Nitric oxide (NO), an important molecule produced in the process of oxidative stress, is synthesized by the enzyme nitric oxide synthetase, that is a product of the iNOS gene. It acts as a biological mediator similar to neurotransmitters and can regulate the tonus of blood vessels. On the other hand, it is an oxygen free radical that can function as a cytotoxic agent in pathological processes, especially in inflammatory diseases [37–39].

Interleukin-6 (IL-6) is one of the leading mediators of the acute phase of inflammation, with a crucial activity on eosinophil chemotaxis to the inflammation site, and has a vital role on coagulation [20, 40]. IL-6 is known as a multifunctional cytokine, which in addition to its pro-inflammatory and sclerosing functions, also affects the activity of neoplastic cells [30, 41]. In addition to its critical local effects, this cytokine has systemic activity, which contributes to the defense of the organism. One of these effects is the elevation of body temperature, causing fever from an endogenous source [21].

Therefore, NO and IL-6 were measured to confirm the results found with the analysis of the transcripts obtained by qPCR. The assays were performed using the supernatants of cell cultures submitted to the treatment already described on 2.4 item. The potential immune-protective activity of LDT11 was evidenced by the results obtained, both at 6 LPS treated and at 24 h post-treatment, which
demonstrated a significant reduction in the production of NO and IL-6 when compared to LPS treated cells.

The evaluation of the expression of the genes TNF-α, COX-2, iNOS, NF-κB, IL-1β and IL-6 at 6 and 24 h after treatment of the cells with LDT11, makes evident that the LDT11 was most effective in protecting against the inflammation when compared with results obtained with the use of commercial drugs.

In addition to a significant reduction in the expression of inflammatory genes, LDT11 also had a faster and superior anti-inflammatory action than the commercial products, and its response was already evident in the test carried out six hours after the treatment of the cells, suggesting that this molecule can be used as an anti-inflammatory drug.

**Conclusion**

LDT11, a phenolic derivative of CNSL, showed potential immunoprotective and anti-inflammatory properties, having a rapid and effective activity. Treatment with LDT11 decreased the expression of the TNF-α, COX-2, iNOS, NF-κB, IL-1β and IL-6 inflammatory genes. Additionally, LDT11 protective effect on inflammation was confirmed by the decreased of NO and IL-6 production. The anti-inflammatory activity of LDT11 was superior to commercial drugs, ASA and DEX.

**Abbreviations**

ANOVA: Analysis of variance; ASA: acetylsalicilic; ATCC: American Type Culture Collection; cDNA: complementary DNA sequence; CNSL: cashew nut shell liquid; CO₂: Carbon dioxide; COX-2: cyclooxygenase isomor 2; CT: Crossing threshold; DEX: Dexamethasone; DMEM: Dulbecco’s Modified Eagle Medium; DMSO: Dimethyl Sulfoxide; DNA: Deoxyribonucleic acid; DNase: Deoxyribonuclease; dNTP: Deoxynucleotides 5’-triphosphate; ELISA: Enzyme Linked Immuno Sorbent Assay; FC: Fold-change; GAPDH: Glyceraldehyde-3-phosphate dehydrogenase; iNOS: Nitric oxide synthase induced; LDT11: semisynthetic Medium; LPS: Lipopolysaccharide; MGMT: Dimethyl Sulfoxide; DNA: Deoxyribonucleic acid; NO: Nitric oxide; IL-6: Interleukin-6; iNOS: Nitric oxide synthase induced; LDT11: semisynthetic

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**Availability of data and materials**

The statistical analysis from the current study are available from the main researcher on reasonable request. Please allow ten business days for data to be emailed.

**Authors’ contributions**

MQS: responsible for the gene expression tests and NO and IL-6 dosages. IMSNT: accountable for the primers efficiency tests used in the study for evaluation of gene expression. FCA: Responsible for cell culture and cytotoxicity testing. RP: Advisor responsible concept for the project. GSH: Responsible for the purification and organic modification for the synthesis of LDT11. LASR: Coordinator of the Laboratory of Therapeutic Strategies Development, where LDT11 was obtained for this research. YKMN: Coordinated the project, responsible analysis of gene expression data. CBP: Assisted in the conceptualization of the study, revised the protocol, corresponding author. All authors read and approved the final version of the manuscript.

**Ethics approval and consent to participate**

Not applicable.

**Competing interests**

The authors have declared that there are no conflict of interest or competing interests to declare.

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