Attenuation of pain and inflammation induced in mice treated orally with crude extract of *Aloysia polystachya* (Griseb.) Moldenke (Verbenaceae)

Mariela Ortiz¹, Isabelle Wilson², Yenny Montalbetti¹, Olga Heinichen¹, Wilfrido Arrúa¹, Nelson Alvarenga³, Derlis A. Ibarrola¹, María del Carmen Hellión-Ibarrola¹

¹Departamento de Farmacología. Facultad de Ciencias Químicas. Universidad Nacional de Asunción Paraguay
²Department of Psychological and Brain Sciences. Dartmouth College, Hanover, NH USA
³Departamento de Fitoquímica. Facultad de Ciencias Químicas. Universidad Nacional de Asunción Paraguay

**ABSTRACT**

The objective of this work was to determine the analgesic and anti-inflammatory activity of *Aloysia polystachya* (Griseb.) Moldenke (Verbenaceae) in experimental models of acute pain and inflammation in mice. Methods of pain induced by caudal pressure (Randall-Selitto), chemical stimulation (acetic acid or the writhing test), and thermal stimulation (hot plate) were used to study analgesic effects. Additionally, edema of the paw induced by injection of 1% carrageenan was used to evaluate the anti-edema activity of *A. polystachya*. Oral doses of 100 and 200 mg/kg of crude extract of *A. polystachya* (CEAp) significantly reduced the sensibility to painful stimuli induced by the application of pressure in the tail comparable with an analgesic effect (*p* < 0.05) in a non-dose dependent manner. Additionally, the number of abdominal contortions was significantly reduced in comparison with the control group and with similar strength to the group treated with Indomethacin 10.0 mg/kg. Similarly, in the model of pain induced by thermal stimulation, it was observed that the groups treated with CEAp presented statistically significant analgesic activity in comparison with the control group and with a similar intensity to the group treated with morphine 6.0 mg/kg. Finally, a statistically significant reduction of edema induced by 1% carrageenan was observed with oral administration of 100 mg/kg of CEAp in comparison to the positive control of edema in a manner similar to the group treated with Indomethacin 10 mg/kg. Based on these results, it was concluded that the CEAp possesses the capacity to increase pain threshold in three pre-clinical models of pain induced (mechanical pressure, chemically and thermally) in mice, compatible with an analgesic effect. Also, CEAp demonstrated antiedematous capacity in carrageenan-induced paw edema in mice, concordant with anti-inflammatory effect using the plethysmography method. These pharmacological effects are potentially due to the presence of verbascoside in CEAp. Additionally, these experimental results are correlated with the popular use of CEAp and present a variety of opportunities for pharmaceutical research such as the development of innovative phytopharmaceuticals.

**INTRODUCTION**

Pain presents a major threat to survival and requires humans to perceive their environment effectively, identify hazards, act to prevent damage to one’s body, and promote effective recovery in the event of damage (Tabor *et al.*, 2017). Pain is a fundamental experience associated with the perception of actual or potential harm to oneself. It can result in an infection, inflammation, peripheral tissue damage, or neural damage. In general, it is recognized that neurotransmitters, sodium channels, and neuromodulators mediate painful responses in the clinic, and in each of the different experimental models of nociception they may act differently (Bannon, 2001; Kopach *et al.*, 2012). The
evidence follows that a large proportion of patients with chronic pain can develop major depressive disorder compared to patients with other chronic medical conditions (Tenti et al., 2021).

Inflammation is a response of the organism against a harmful stimulus and is always associated with pain (da Silva et al., 2011). It takes place in vascularized connective tissue and involves vascular changes, cellular events, and the production of chemical mediators of inflammation. All these components of the system are closely linked and involve a complicated series of events including arteriolar dilation, increased vascular permeability in venules and capillaries, fluid exudate (including plasma proteins), and migration of leukocytes to the inflamed area (de Morais Lima et al., 2011; Trivellatograssi et al., 2013).

The existence of a relationship between chronic pain and depression is now widely accepted among researchers (Rojas-Corrals et al., 2003). Chronic pain and depression are two very frequent and debilitating concurrent conditions. They present a bidirectional longitudinal relationship due to the high probability of developing depression secondary to chronic pain and vice versa (Maalø, 2021). Various antidepressant drugs used to treat depression are effective for treating pain associated with damage to the nerves (neuropathic pain). At least a third of patients with neuropathic pain that received traditional antidepressants (like amitriptyline) had relief of moderate pain. There is also evidence that venlafaxine, a newer antidepressant, has similar effectiveness to traditional antidepressants. Nevertheless, approximately a fifth of the patients that receive these drugs for pain must stop treatment due to adverse effects. Neuropathic pain can be treated with antidepressants, and its effect is independent of any effect on depression (Gregory et al., 2014).

Aloysia polystachya (Griseb.) Moldenke (Verbenaceae), known commonly as “burrito,” is a perennial shrub that grows in Paraguay, northwestern Argentina, Bolivia, and Brazil. The leaves infusion of A. polystachya is used as a tonic, digestive, and carminative agent, for stomach pains and slow digestion, against liver disorders, for the treatment of “empacho,” and as a tonic of the nerves (Consolini et al., 2011). In previous studies, we have described, among other effects, the sedative, anxiolytic, and antidepressant effects of the hydroalcoholic extract of A. polystachya leaves in mice and rats (Hellión-Ibarrola et al., 2006; Hellión-Ibarrola et al., 2008; Mora et al., 2005).

The presence of glycosides phenylethanoids, triterpenes/steroids, phenols/tannins, flavonoids, and traces of saponins and lactones has been reported as chemical components of the hydroalcoholic extract (Carmona et al., 2019; Pereira et al., 2019). Additionally, compounds such as terpenes, monoterpenes (carvacrol, carvone, eucarvone, limonene (−), limonene, α-pinene, sabinene (+), sabine (−), thujone, α-thujone, α- (−) thujone, β-thujone, iso-thujone, y (+) iso-thujone, and sesquiterpenes are presents in the essential oil (Carmona et al., 2019). Data not published by our lab accounts for the presence of zinc, copper, calcium, and vitamins B1 and B2 in significant amounts.

Considering that the preclinical studies with A. polystachya demonstrated its anxiolytic and antidepressant activity (Hellión-Ibarrola et al., 2006; Hellión-Ibarrola et al., 2008; Mora et al., 2005) as well as its clinical validation as an anxiolytic in the treatment of anxiety (Carmona et al., 2019) and that antidepressant drugs such as amitriptyline are effective in the treatment of chronic pain, it is reasonable to evaluate the influence of A. polystachya on experimental pain as a potential alternative or complementary therapy with the use of medicinal plants. Therefore, this work aims to determine the influence of the oral administration of the crude extract of A. polystachya (CEAp) on different preclinical models of acute pain induced by harmful stimuli (mechanical pressure, 0.8% acetic acid, and thermal) and its influence on carrageenan-induced inflammation in mice.

MATERIALS AND METHODS
Plant material and extraction procedure
Cultivated samples of A. polystachya were collected in Colonia Potroco Sur y Mafussi de Pedro Juan Caballero, Paraguay. A voucher sample was authenticated and deposited at the Herbarium of the Faculty of Chemical Sciences UNA (Code C. Céspedes 953). The collected samples were air-dried, protected by sunlight, and conditioned in appropriated labeled containers. In traditional Paraguayan medicine, all medicinal plants are milled with a hardwood stick inside a wooden container and later placed in a container filled with tap water and used, unfiltered, as a beverage or for drinking a traditional “terere” (Martinez and Barboza, 2010). Consequently, in addition to the supernatant liquid, small particles are swallowed, whose components can potentially be absorbed from the gastrointestinal tract. Based on the latter, for this study, we selected a more powerful extraction system and closer to the popular procedure, using the ethanol: water mixture (60:40). The dried material was subjected to grinding to obtain a fine powder, which was extracted with ethanol: water (60:40) using the method of conventional reflux for 1 hour in a bath and at 50°C. The procedure of extraction was repeated two times, and the filtered hydroalcoholic extracts were mixed, homogenized, and evaporated at low pressure. The concentrated CEAp was frozen and finally lyophilized for its use in all biological experiments.

Drugs
All the drugs used were of analytical quality. Carrageenan, indomethacin, and acetic acid were obtained from Sigma (USA), and sodium chloride was acquired from Wako (Japan). Verbasoside (purity > 98%) was purchased from ABCAM Inc. (Discovery Drive, Cambridge Biomedical Campus, Cambridge, CB2 0AX, UK) Morphine, ethanol, and propylene glycol for pharmaceutical use were obtained locally.

Animals
Swiss Albino female mice (20–30 g, b.w.) obtained from the Animal Facility of the Department of Pharmacology at the Faculty of Chemical Sciences (National University of Asunción) were used. All animals were maintained in a controlled environment (19°C–23°C temperature and 55% ± 5% relative humidity) with 12 hours of light/dark cycle. The animals received commercial foods and were fasted overnight before corresponding experiments, with free access to water. The experimental procedures, handling, and treatment of animals were conducted in accordance with international standards of animal welfare established by the Ethics Committee of the European Community (Real Decreto, 2005). The experimental protocol was submitted and approved by the institutional Ethics Research Committee of the Faculty of Chemical Sciences on October 6, 2016 (Code 238/16).
UPLC-ESI-MS (Ultra Performance Liquid Chromatography-Electrospray Ionization-Mass Spectrometry) analysis

The analyses were carried out using a Waters (Milford, MA) Acquity ultra-performance liquid chromatograph coupled with a Xevo TQD triple quadrupole mass spectrometer used as the detector. A Phenomenex KINETEX core-shell EVO-C18 (2.1 × 100 mm, 1.7 μm) column was used for separation purposes. The temperature of the column was kept at 40°C. The mobile phase used was methanol (phase A) and water (phase B) (LC-MS grade, Merck KGaA, Darmstadt, Germany), with 0.1% formic acid and 10 mM ammonium formate for both. The flow rate was 0.3 ml/minute in gradient mode, as follows: 0–0.7 minutes, 80%–80% A; 0.7–3.2 minutes, 80%–60% A; 3.2–7.6 minutes, 60%–20% A; 7.6–8.3 minutes, 20%–0% A; 8.3–10.4 minutes, 0%–80% A, 10.4–13 minutes, 80%–80% A. The samples were dissolved in LC-MS MeOH, filtered through 0.22 μm nylon syringe filters, and injected at 5 mg/ml.

The MS spectra were acquired in full scan mode (m/z 80–800, at a speed of 2,000 m/z per second) with ESI as the ionization source in negative and positive modes. The mass spectrometer conditions were as follows: electrospray capillary voltage 2.5 kV, source temperature 150°C, desolvation temperature 350°C, cone gas flow 80 l/hour, and desolvation gas flow 900 l/hour. The cone voltage was set at 30 V. The system was controlled using the Waters MassLynx V4.1 software.

Evaluation of the analgesic activity of the CEAp

Mechanical pressure-induced painful stimulus in mice (Randall-Selitto Test)

Swiss Albino female mice (20–30 g b.w.) were used and distributed randomly in five groups with six mice per group. One group was treated orally with the vehicle (0.1 ml for each 10 g, b.w.), and a second group was treated with indomethacin (10 mg/kg, p.o.). The other three groups received CEAp (100, 200, and 500 mg/kg, p.o.) dissolved in a 0.9% saline solution. After 1 hour of receiving treatment, the animals were immersed in the subplantar (s.p.) region of the right hind legs with 40 μl of carrageenan (1%) as described by Kim et al. (2013). A similar volume of a normal saline solution was injected into the contralateral foot. Paw volume measurements were recorded immediately before, every 30 minutes, and up to 3 hours after carrageenan injection. Briefly, the procedure consisted of immersing the rear legs up to the lateral malleolus inside the digital plethysmograph vessel (LE 7500 Panlab, Harvard Apparatus, Spain). The displaced volume is automatically recorded and tabulated as an individual value (Muhammad et al., 2012). The difference in volumes between the legs was considered as the edema value for each animal.

Statistical analysis

The results were expressed as mean ± standard deviation. Analysis of variance (ANOVA) followed by Tukey’s multiple comparison test was performed as statistical analysis using the GraphPad Prism 5.0 software. The level p < 0.05 was considered to be statistically significant.

RESULTS

Verbascoside identification

Verbascoside was identified in the crude extract by LC-MS (Fig. 1). The chromatogram of the standard showed a peak at 7.37 minutes with a deprotonated molecular ion of m/z 623.89 [M-H]-. The extract showed a peak at 7.46 minutes, with the same molecular ion of m/z 623.89 [M-H]- as the standard (Fig. 2). This fact confirmed the presence of the compound in the extract.

Effect of CEAp on mechanical pressure-induced pain behavior in mice (Randall-Selitto test)

As depicted in Figure 3, groups of animals treated with doses of 100 (241.4 ± 47.41 g; p < 0.001) and 200 mg/kg (210.0 ± 81.20 g; p < 0.01) of CEAp revealed a statistically significant analgesic effect in comparison to the group treated with the vehicle (104.0 ± 25.80 g). The pain threshold increased by 233%, 136%,
and 102% in groups treated with indomethacin, 100 and 200 mg/kg of CEAp in comparison to the control group, respectively. Nevertheless, the dose of 500 mg/kg (147 ± 35.80 g) did not induce a statistically significant response compared to the control group. As was indicated above, indomethacin at 10 mg/kg (346.0 ± 105.5 g; \( p < 0.001 \)) demonstrated a statistically significant effect which validated the method used.

Effect of CEAp on chemically induced (acetic acid 0.8%) pain behavior in mice (Writhing test)

The groups of animals treated orally with the doses of 100 (30.90 ± 20.66; 51%), 200 (30.80 ± 11.55; 51%), and 500 mg/kg (19.88 ± 11.51; 68%) of CEAp caused a very significant reduction \( (p < 0.001) \) in the number of abdominal contortions compared to the group treated with the vehicle (62.78 ± 14.85), compatible with an analgesic activity of the sample (Fig. 4). In the same sense, administration of indomethacin denoted a significant reduction of 51% in the number of contortions (30.78 ± 6.92) elicited, in comparison to the group treated with the vehicle, validating the utilized method. These results indicated that the dose of 500 mg/kg is more potent than the positive control indomethacin 20 mg/kg in the reduction of pain behavior induced chemically with acetic acid. The observed effects in this test are not dose dependent in nature.

The analysis of variation in the number of chemically induced writhing behavior as a function of time denotes that the speed of reduction of the painful behavior in groups of animals treated with indomethacin (10 mg/kg) and CEAp (500 mg/kg) begins at 10 minutes after the injection of 0.8% acetic acid, respectively. Undoubtedly, the reduction in the number of
writhing behavior is more intense, as a function of time, with all doses of CEAp than the vehicle-treated group. Furthermore, the indomethacin curve shows a lower number of writhing behavior than the negative control group, demonstrating its effectiveness as an analgesic and thus validating the method used (Fig. 5). Linear regression analysis between indomethacin-treated and CEAp groups (100, 200, and 500 mg/kg) differs significantly when compared to the vehicle-treated group.

**Effect of CEAp on thermally induced pain behavior in mice (Hot Plate Test)**

Figure 6 shows the influence of oral administration of CEAp on reaction time (latency) to heat-induced pain behavior in mice. There is a significant increase in the reaction time of the group treated with 100 mg/kg of CEAp (20.52 ± 4.16 seconds; \( p < 0.01; 171\% \)) in comparison to the group treated with the vehicle (7.57 ± 2.56 seconds). Nevertheless, it should be noted that doses of 200 (13.50 ± 3.96 seconds) and 500 (13.73 ± 7.70 seconds) mg/kg of CEAp do not raise the latency time compared to the group treated with the vehicle. Additionally, a statistically significant increase in morphine-induced latency (28.04 ± 3.46; \( p < 0.001; 270\% \)) was observed compared to the negative control group, which validates the method used.

**Anti-inflammatory effect of the CEAp on carrageenan-induced acute paw edema in mice**

Figure 7 shows the influence of oral administration of CEAp on carrageenan-induced acute paw edema in mice. There is a significant decrease of 56% in the paw edema provoked by carrageenan (10.63 ± 2.93; \( \mu l \)) by treatment with 100 mg/kg de CEAp (6.00 ± 1.55; \( \mu l; p < 0.05 \)). In the same sense, 10 mg/kg of
indomethacin (5.56 ± 2.88; µl; p < 0.01) provoked a statistically significant decrease of 52% in the inflammatory response elicited by carrageenan, thus validating the phlogistic method used. Nevertheless, it should be noted that groups treated with doses of 200 (7.20 ± 1.92; µl) and 500 (7.00 ± 2.53; µl) mg/kg of CEAp did not significantly reduce the level of edema. Concurrently, the statistically significant increase in the volume of edema induced by carrageenan (10.63 ± 2.93; µl; p < 0.001) in comparison with the untreated basal group (2.75 ± 2.50; µl) was observed. Finally, in the group treated with 10 mg/kg of indomethacin (5.56 ± 2.88; µl; p < 0.01), an efficient reduction of the volume of edema was observed and globally validates the method used.

**DISCUSSION**

This work evaluated the antinociceptive and anti-inflammatory activity of CEAp, with previously demonstrated antidepressant activity in our lab (Hellión-Ibarrola et al., 2006; Hellión-Ibarrola et al., 2008; Mora et al., 2005). Usually, in the validation of plants used for medicinal purposes in Paraguay, the first step is to select plants with a long history of use, the procedure for obtaining the extract, followed by the selection of the preclinical method to be used (induced pain and inflammation...
in rodents), consistent with popular use. The next step is the oral treatment of the groups of animals with different doses of the sample to determine their safety and efficacy. Then, if the result deserves efficacy, it is considered to carry out refined studies to determine the components, molecular mechanisms of action of the effects caused by the extract.

Based on the International Association for the Study of Pain, Maallo et al. (2021) explain pain as “an unpleasant sensory or emotional experience associated with actual or potential tissue damage.” These pain components have their own pathways, centers, and regulatory mechanisms. In general, pain can be classified as nociceptive pain provoked by the normal activation of peripheral nociceptors (“alarm” and protection of the organism against harmful stimuli, with high threshold pain and often accompanied by a withdrawal reflex), inflammatory pain provoked by tissue damage and activated by invasion of inflammation-mediating cells (macrophages, neutrophil mast cells, and granulocytes) with low pain threshold, and pathological pain (neuropathic and dysfunctional) that appears as a consequence of a lesion to the nervous system (central or peripheral) or abnormal central processing (Woollf, 2010).

In the same context, the pain-depression or depression-pain couple is accepted as very common comorbidities in the general adult population with chronic pain. 19% of European adults and 20.4% of American adults have chronic pain, and depression is the most frequently associated comorbidity. The prevalence of depression in the European and American societies is 7% and 7.5%, respectively (Tenti et al., 2021). Severe chronic pain is still poorly managed due to the continuous imbalance between pharmacological analgesia and tolerability, which leads to a poor treatment outcome (Barroso et al., 2021). The experimental models of pain are separated as acute, persistent, and neuropathic. Experimental acute pain is induced by mechanical, chemical, and thermal stimuli. Persistent pain is represented by the formalin test and neuropathic pain by nerve ligation assays (Bannon and Malmberg, 2007). Therefore, A. polystachya has sufficient merits to be evaluated for its potential analgesic and anti-inflammatory activity in preclinical models.

The oral administration of CEAp in mice denotes an efficient analgesic and anti-inflammatory capacity in the models used. In the mechanical pressure-induced pain assay (Randall-Selitto test), an effective non-dose-dependent effect was observed. The pain threshold was increased by 136% and 102% in the groups treated with 100 and 200 mg/kg of CEAp, respectively, compared to the control. This indicates a major analgesic potential with the lowest dose of CEAp, while a decreasing degree in the intensity of the analgesic effect was found with increasing dosage until it resulted in nonsignificant values with the dose of 500 mg/kg of CEAp. Additionally, the analgesia induced with the extract was moderate in intensity and 30% lower than that obtained with indomethacin (analgesic reference drug) which increased the pain threshold by 233% compared to the control. It should be noted that the behavioral measures of mechanical sensitivity are commonly used to measure alldynia (a painful response to nonharmful stimuli) and hyperalgesia. Systems of application of mechanical pressure (caudal, plantar, etc.) have been developed to measure pain responses provoked by mechanical stimuli. Tail withdrawal or vocalization of the animal (fastened) is used as the endpoint of the test (Gregory et al., 2014). Assessing withdrawal response to pressure applied to tissue using the Randall-Selitto model reproduces similar results in the reduction of pressure pain threshold, commonly observed in clinical pain conditions (fibromyalgia, myofascial pain, or osteoarthritis). Considering the decreasing pain sensitivity (analgesia) that was observed with all doses when pain was inflicted by chemical methods, CEAp may have clinical resemblance and utility for acute pressure pain such as in fibromyalgia (Gregory et al., 2014). Additionally, the intensity of the observed effects was similar to that of the indomethacin. This model was characterized by the induction of acute pain of longer duration (compared to caudal pressure) due to peritoneal irritation, involving tissue damage in which inflammatory signaling mechanisms of higher intensity and sensitivity operate. In agreement with Carvalho et al. (2006), the principal mediators are eicosanoids and sympathomimetic amines preceded by the release of TNF-α (nociceptive cytokine). Nevertheless, the obtained results show the efficacy of CEAp as an analgesic and its probable anti-inflammatory effect.

In addition, in the heat-induced pain assay, an efficient increase in the latency time of the hot plate jump or paw lick compatible with the analgesic effect of CEAp (100 mg/kg) was observed. The act of jumping or licking involves supraspinal structures indicating that it goes beyond nociceptive reflexes because it involves information processing in higher structures (Gregory et al., 2014). Accordingly, previous work with CEAp has shown antidepressant and anxiolytic activity in mice (Heilón-Ibarrola et al., 2006; Heilión-Ibarrola et al., 2008; Mora et al., 2005) treated via the oral route. There is a possibility that analgesic activity is mediated by regulatory pathways at higher levels because it increases the latency time in the hot plate assay. Efforts
to carry out studies to elucidate the regulatory mechanism of the analgesic effect must be reinforced to advance this work.

Likewise, CEAp was active against the acute inflammation caused by subplantar injection of carrageenan. Carrageenan-induced paw edema in the mouse is also associated with nociceptive changes in the paw and migration of inflammatory cells to the injection site (Gregory et al., 2014; McCarson, 2015) and is sensitive to nonsteroidal anti-inflammatory drugs (Lapa et al., 2002). Recently, it has been demonstrated that the phase of the inflammatory response after the injection of carrageenan in the paw of the mice is associated with the production of cyclooxygenase 2 (COX-2), high prostaglandin production (by the action of COX-1 and COX-2), free oxygenated radicals (Nitric Oxide derivatives from endothelial nitric oxide synthase, and inducible nitric oxide synthase), and neutrophil infiltration (Duarte et al., 2016; Fehrenbacher et al., 2012; McCarson, 2015). As demonstrated, CEAp shows anti-inflammatory activity by reducing the edema by an unknown mechanism. However, the presence of acteoside or verbascoside, among other components of CEAp, is highly probably accountable for analgesic/anti-inflammatory activity. According to Gutiérrez-Rebolledo et al. (2016), the verbascoside is a well-known COX-2 inhibitor with antioxidant capacity in addition to its MAO-A (Monoamine oxidase A) inhibitory effects (Pereira et al., 2019), related at least partially to antidepressant/anxiolytic properties (Carmona et al., 2019). Interestingly, drugs able to block inflammatory cytokines or associated inflammatory pathways (COX-2) are effective in reducing depressive symptoms in patients with chronic inflammation (rheumatoid arthritis, autoimmune disease, cancer, etc.) and those patients suffering from major psychiatric disorders (Pereira et al., 2019). Indeed, we have no knowledge about the molecular mechanism of action of CEAp, and with the available results, it is not possible to make a complete hypothesis to address which of the pathways mentioned above and if verbascoside or which other component(s) are involved in analgesia and inflammation. Consequently, this study is not yet focused on determining molecular mechanisms involved in increasing pain threshold or correcting tissue damage caused by inflammation. The presence of verbascoside in CEAp accounts for the huge potential to be responsible for these analgesic and anti-inflammatory activities. Really, the aim of this study is to determine the influence of the oral administration of the CEAp on different preclinical models of acute pain induced by harmful stimuli (mechanical pressure, 0.8% acetic acid, and thermal) and its influence on carrageenan-induced inflammation in mice. Certainly, the pharmacological study of the specific molecular mechanism is the prospective core to be examined experimentally.

It should also be mentioned that components obtained from the essential oil (carvone and α-thujone) or from CEAp (phenolic compounds, glycosides phenylethanoids, and terpenes) are potentially responsible for the wide profile of pharmacological effects observed in animals and ultimately clinical trials of this noble and appreciated natural product (Carmona et al., 2019; Consolini et al., 2011; Hellión-Ibarrola et al., 2008; Pereira et al., 2019). Among the pharmacological effects studied are described low acute toxicity and anxiolytic and antidepressant activity in mice and rats. Additionally, antispasmodic effects were depicted from the aqueous extract and dyes of A. polystachya (Griseb.) Moldenke probably mediated by blocking Ca$^{2+}$ channels (Consolini et al., 2011). The most remarkable result is that CEAp proved to be clinically effective in moderate anxiety (Carmona et al., 2019) and its principle component, acteoside (verbascoside), also present in other species, proved to be effective as an anti-inflammatory treatment (Gutiérrez-Rebolledo et al., 2016). Therefore, the results of this work are correlated with available literature and validate its popular use as an analgesic and anti-inflammatory treatment. Consequently, in addition to the popular procedure, these outcomes reinforce the real possibility of developing innovative products such as the formulation of standardized phytotherapy from semipurified or isolated components from A. polystachya for a more precise and effective treatment of pain and inflammation associated with depression or vice versa.

Limitations of this study include the restricted knowledge about the profile of components and antioxidant properties of the crude extract. Also, the pain or inflammation mechanism affected is unknown, and molecular/docking studies to shed light on these questions are required. Therefore, the next steps with A. polystachya include solving the chemical components, antioxidant properties, and the deep examination of the molecular/docking mechanism involved in analgesic, anti-inflammatory, and antidepressant activities.

CONCLUSION

The present work demonstrated that the oral administration of the CEAp produces moderate analgesic activity using the models of induced pain in mice through three harmful stimuli: a) caudal mechanical pressure, b) chemical pain of the abdominal contortions, and c) thermal pain with indomethacin as an analgesic and anti-inflammatory positive control. The presence of verbascoside was determined by chromatographical methods.

Based on these results, it was shown in the Randall–Selitto test that the doses of 100 and 200 mg/kg of CEAp increase the pain threshold by 136% and 102% compared to the group of animals treated with the vehicle, respectively. Also, it was determined that 100, 200, and 500 mg/kg of CEAp have the capacity to reduce considerably the contortions induced by 0.8% acetic acid in a dose-dependent manner by 51%, 51%, and 68% compared to the vehicle. These results are compatible with the analgesic activity of the CEAp. The potency of 100 and 200 mg/kg CEAp was similar to indomethacin, while that of 500 mg/kg was much better than the positive control in the decrease of contortions induced by intraperitoneal administration of 0.8% acetic acid solution. Likewise, the pain threshold was increased with 100 mg/kg of CEAp by 171% using the hot plate, reinforcing the potential effectiveness of this natural product as an alternative analgesic treatment.

Likewise, the experimental edema induced by carrageenan was reduced in over 50% with 100 mg/kg of CEAp whose potency was very similar to that caused by indomethacin (10 mg/kg). We conclude that the results are correlated with the popular use of CEAp and open a variety of possibilities for pharmacotoxicological investigation and the potential development of innovative phytopharmaceuticals with relevant health-commercial importance. Complementary pharmacological and chemical works are underway to determine the possible mechanism of action and a full component, in addition to verbascoside, involved in the analgesic-anti-inflammatory effect.
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AUTHOR CONTRIBUTIONS
All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work. All the authors are eligible to be an author as per the international committee of medical journal editors (ICMJE) requirements/guidelines.

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DATA AVAILABILITY
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