In vitro ovicidal effect of a *Senecio brasiliensis* extract and its fractions on *Haemonchus contortus*

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**Abstract**

**Background:** Haemonchosis affects sheep husbandry and its treatment is often compromised due to the development of anthelmintic resistance. Plant-derived bioactive compounds have been studied as alternative to control *Haemonchus contortus*. The objective of this study was to evaluate the effect of *Senecio brasiliensis* extracts on *H. contortus* egg hatching and infective larvae migration.

**Results:** The aqueous extract from dried and fresh plant and alkaloid-enriched fraction of the previously dried leaves of *S. brasiliensis* inhibited *H. contortus* egg hatching. The main plant compound in alkaloid fraction was integerrimine, a pyrrolizidine alkaloid (PA). However, the aqueous extract from dried plant displayed higher efficacy when compared to their alkaloid enriched or non-polar fractions, meaning that, although PAs contributed to the ovicidal effect, other compounds in the plant can also contribute to their effect. Furthermore, the aqueous extract from dried plant also had higher efficacy than aqueous extract from fresh plant in larvae migration inhibition. Finally, extract from dried plant presented low in vitro cytotoxic effect.

**Conclusion:** Taken together our results suggest a good anthelmintic effect of *S. brasiliensis*, especially when aqueous extract is prepared from dried plant. Further in vivo studies should be performed focused on forms of administration of this extract in rearing sheep.

**Keywords:** Anthelmintic, Egg hatching, Infective larvae migration, Plant extract, Pyrrolizidine alkaloids, Small ruminants

**Background**

Endoparasitic diseases are a high concern in sustainable sheep production [1]. *Haemonchus contortus* [2] (barber’s pole worm), a blood-sucking nematode and member of the order Strongylida and family Trichostrongylidae, can cause anemia, anorexia, diarrhea, gastritis, and even death in animals [3]. This parasitic infection results in direct economic losses related to decreased animal performance and/or death as well as indirect economic losses linked to the high cost of anthelmintic drugs and the labor and equipment required for the control of parasitosis [4, 5].

*H. contortus* control is mainly based on the use of commercial anthelmintics; however, these drugs have lost their effectiveness due to the development of drug-resistant parasite strains [6, 7]. To aid the control of nematodes, in conjunction with the use of anthelmintics, alternative strategies have been developed, such as pasture management, nutritional adjustments, genetic selection, use of nematophagous fungi, and the development of plant-derived anthelmintic compounds [8, 9].

The genus *Senecio* (Asteraceae) comprises approximately 1500 species [10]. Plants belonging to this genus are known for the production of compounds such as alkaloids, sesquiterpenes, and flavonoids [11] and for their anti-inflammatory, vasodilator, antiemetic, antimicrobial, and parasiticide activities [12–14].

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Senecio brasiliensis is a native species found in south and southeast of Brazil [15]. It is toxic to the livestock [16, 17] due to the presence of pyrrolizidine alkaloids (PAs), which are widely found in the Asteraceae, Boraginaceae, and Fabaceae families [18, 19]. However, the leaves and inflorescence of S. brasiliensis are used in traditional medicine for the treatment of stomach pain [20], a practice justified by the antiulcerogenic and cytoprotective effect of PAs [21, 22].

Based on reports of the pharmacological effects of the genus Senecio and the species S. brasiliensis, we aimed to evaluate the effect of aqueous extract of this plant and its non-polar and alkaloid fractions on H. contortus egg hatching and infective larvae migration, besides evaluating the cytotoxic effect of this plant.

**Results**

The GC-MS analysis showed the presence of four constituents for the alkaloid fraction (AF) from dried plant (DP) and eight constituents for AF from fresh plant (FP) (Table 1). In AF from DP, all peaks were attributed to PAs. On the other hand, in AF from FP, in addition to the alkaloids, carboxylic acids and ketones were found. For both extracts, the major compound was integerrimine (~91% in AF from DP and ~79% in AF from FP).

The effective concentrations to inhibit H. contortus egg hatching (Fig. 1) by 50% (EC50) were 0.660 mg.mL\(^{-1}\) for DP and 2.596 mg.mL\(^{-1}\) for FP, respectively (Table 2). DP showed high inhibitory activity at the highest concentration tested (94%) (Fig. 2a), whereas FP showed moderate inhibitory activity at the highest concentration tested (73%) (Fig. 2b). The percentage of inhibition of egg hatching ranged from 1 to 94% for DP and from 4 to 73% for FP. For DP aqueous extract, inhibition rates above 50% were observed for concentrations higher than 0.625 mg.mL\(^{-1}\) (Fig. 2a). On the other hand, for FP, significant egg hatching inhibition was observed only for concentrations above 2.5 mg.mL\(^{-1}\) (Fig. 2b).

The estimated EC50 was 2.435 mg.mL\(^{-1}\) for the AF from DP. Non-polar enriched fraction (NPF) from DP showed EC50 of 4.356 mg.mL\(^{-1}\). The EC50 values for AF and NPF from FP could not be determined, due to low effect, since the eclosion inhibition was less than 10% in all tested concentrations. At the highest evaluated concentrations, AF (Fig. 2c) and NPF (Fig. 2d) from DP exhibited egg hatching inhibition of 98 and 70%, respectively. The average inhibition for AF from DP ranged from 1.7 to 98% and started at 2.5 mg.mL\(^{-1}\), while the average inhibition for NPF from DP ranged from 0.5 to 70% and started at 5 mg.mL\(^{-1}\). The egg hatching inhibition in the positive control (thiabendazole 5 mg.mL\(^{-1}\)) was 98%.

Concerning larval migration inhibition, the EC50 were 10.15 mg.mL\(^{-1}\) for DP and 1712 mg.mL\(^{-1}\) for FP (Table 2). The maximal larval migration inhibition, observed when high concentrations of plant extract were used, was 15% for FP and 82% for DP. The percentage of inhibition migration ranged from 6 to 82% for DP and from 2 to 15% for FP. For DP, migration inhibition rates above 50% were observed only in concentrations higher than 12.5 mg.mL\(^{-1}\) (Fig. 3a). On the other hand, no migration inhibition above 50% for FP was observed (Fig. 3b). The larval migration inhibition of the positive control (levamisole 6.25 \(\mu\)g.mL\(^{-1}\)) was 96%.

A dose-dependent pattern is observed for cytotoxic activity of DP aqueous extract (Fig. 4). However, the effect of this extract in mammal cell lethality was pronouncedly less intense than for egg hatching or larval migration inhibition, as demonstrated by the SI that was 68.63 and 4.48, respectively. The cytotoxic EC50 of DP was estimated in 45.43 mg.mL\(^{-1}\) (Table 2). Maximal cell

**Table 1** GC/MS analysis of Senecio brasiliensis alkaloid fraction from dried and fresh plant

| Peak | tR  | IK  | Name of the compound                                      | Peak area (%) |
|------|-----|-----|-----------------------------------------------------------|---------------|
| 1    | 17.020 | 1464 | Senecionine                                               | 3.34          |
| 2    | 18.114 | 1436 | Integerrimine                                             | 91.14         |
| 3    | 18.400 | 1456 | Platyphylline                                             | 1.46          |
| 4    | 19.704 | 1444 | 3(2H)-isoquinolinolone, octahydro-, (4ar-trans)           | 4.06          |
| 1    | 4.830  | 1381 | Cyclohexanone, 5-methyl-2-(1-methylethylidene)             | 4.15          |
| 2    | 5.721  | 1491 | 4-amino-2-nitro-benzaldehyde oxime                        | 0.84          |
| 3    | 7.178  | 1648 | 1,2-cyclopropanedicarboxylic acid, 3-(1-methylethenyl)-, diethyl ester | 4.12          |
| 4    | 8.959  | 1406 | 2-(4-nitrobutyryl) cyclo-octanone                         | 1.19          |
| 5    | 17.071 | 1363 | Senecionine                                               | 4.45          |
| 6    | 18.254 | 1446 | Integerrimine                                             | 79.26         |
| 7    | 19.068 | 1500 | 2-anilino-4-methylquinoline                               | 0.82          |
| 8    | 19.73  | 1445 | Neo-triangularine                                         | 1.5           |

GC/MS: Gas Chromatography-Mass Spectrometry, tR: retention time, IK: Kovats Index
lethality was 54.94% when 50 mg.mL\(^{-1}\) of DP extract was used.

**Discussion**

Integerrimine, found in plant extracts, is among the main PAs in *Senecio* [22–24]. Although integerrimine was the most abundant component in both AFs, its concentration was higher in AF from DP as compared to AF from FP, possibly because the drying procedure at 40 °C increased the concentration of some compounds while caused the loss of others, what could also be a plausible explanation for the reduction of compounds diversity in AF from DP when compared to AF from FP. Previous studies have shown that hot-drying can affect the chemical composition of plant extracts when compared with those obtained by freeze-drying (lyophilization) [25, 26].

An anthelmintic is considered effective if it exhibits at least 90% of inhibitory activity [27]. Considering this cut-off, DP and AF from DP from *S. brasiliensis* showed efficacy against egg hatching of *H. contortus* isolated from sheep feces. In our experiment we observed the phenomenon ‘failing eclosion’, described by Vargas-Magaña et al., [28] and Chan-Pérez et al., [29] in studies with plant extracts. These authors used this term to differentiate larval eggs from morulated eggs, which were classified as dead. In our work we observed the presence of morulated eggs only in the test with the positive ovicidal control Thiabendazole. We believe that the extract of *S. brasiliensis* prevents the larva exit of the egg, by mechanisms still unknown. Regarding larval migration inhibition, it is not possible to conclude an effectiveness of *S. brasiliensis* extracts, since at the tested concentrations; the highest inhibitory activity observed was 82.6%, observed for DP aqueous extract. Taking into consideration the effect of DP on egg hatchability, we believe that, in further studies, *S. brasiliensis* extract should be considered to be used in reduction of pastures contamination, acting as an environmental control, as observed by Niezen et al., [30].

Among the set of alkaloids identified in AF from DP, PAs were the most abundant, suggesting their participation in the observed effect. In addition, the efficacy of those substances in combating plant nematodes has already been reported [31]. Surprisingly, the effect of DP (EC\(_{50} = 0.660 \text{ mg.mL}^{-1}\)) was higher than AF from DP (EC\(_{50} = 2.435 \text{ mg.mL}^{-1}\)), suggesting that PAs are not the only responsible components for anthelmintic effects observed, which has already been reported for other plant extracts. In the study conducted by Brandão et al. [32], the total extract of *Bidens* sp. (Asteraceae) was more effective in reducing *Plasmodium* sp. parasitemia than its fractions. Gomes et al. [33] also recorded a similar event in their in vitro study with *Zizyphus joazeiro* against *H. contortus*; the saponins found in a plant fraction acted together with other substances present in the total extract. Saponins, tannins, phlobatannins, phenols, anthraquinones, flavonoids, glycosides, steroids, terpenes, cardenolides, and chalcones [34, 35] have been identified in other *Senecio* species and some of these compounds have been associated with anthelmintic effects [36–38].

Even according to the traditional medicine, whole plants are used rather than isolated compounds. This phenomenon can be attributed to the synergic effect, what occurs when the combination of components of a plant extract or mixture of plant extracts effect is greater than the sum of individual effects [39]. Although synergy is a well-documented effect, there is a lack in acknowledgment.
concerning its mechanism of action [40]. An example of synergy can be found in the work of Klongsiriwet et al., [41], who detected a greater inhibition of the exsheathment of *H. contortus* infective larvae when tannins and flavonoids were mixed, in relation to the effect of these compounds in the isolated form. However, these authors only speculated how the synergy occurred, as Williamson [40] and Wagner and Ulrich Merzenich [42] cite in their works. Although we have not performed experiments with isolated components and their possible mixtures, we suppose that DP is more potent than its fractions due to some degree of synergy between the alkaloids and the other components present in aqueous extract of the plant.

Despite *S. brasiliensis* having oxicidal effect, this plant may be toxic to animals [43], which may be attributed to liver metabolism of pyrrolizidine alkaloids. Three metabolic ways may occur with PA in liver: dehydrogenation, hydrolysis and n-oxidation [44, 45]. According to these authors, dehydrogenation forms pyrroles that are toxic, whereas hydrolysis and n-oxidation leads to non-toxic metabolites. The pyrroles originating from dehydrogenation of PAs can bind with the hepatocyte DNA, inhibiting cellular mitosis and thereby causing hepatic dysfunction [46, 47]. However, it is known that sheep, the targets of our study, are more resistant to *Senecio* intoxication than cattle due to their ruminal microbiota [48] and liver metabolization system [49].

It worth mentioning that in vitro anthelmintic and cytotoxic effects of *S. brasiliensis* were conducted with plant extracts in a non-metabolized form. The hypothesis of animal toxicity after liver metabolization...
of the compounds corroborate our results, in which in vitro cytotoxicity was low [50, 51] and which confirms previous researches that claim that alkaloids are not toxic in the non-metabolized form [44, 52]. Thus, future studies involving new formulations in order to avoid liver metabolization or to protect that compounds against organism degradation need to be carried out.

Conclusions

DP of S. brasiliensis displays good anti-H. contortus egg hatchability and, less pronouncedly, anti-larvae activity, presenting low in vitro cytotoxicity to Vero cells. Future studies are required to clarify the mechanism of action on H. contortus eggs and the pharmacological mechanisms of the plant extract in animals to validate the DP uses in integrated nematodes control programs.

Methods

Reagents

Sodium hydroxide (NaOH), sulfuric acid (H₂SO₄), dichloromethane (CH₂Cl₂), sodium sulfate (Na₂SO₄), and methanol (MeOH) (High Performance Liquid Chromatography grade) were purchased from Vetec® (Brazil). For the parasitological tests, thiabendazole, tween® 80, levamisole, sodium hypochlorite (NaOCl) and sodium chloride (NaCl) were purchased from Sigma-Aldrich (USA). Rezasurin used in cytotoxicity evaluation was also purchased from Sigma-Aldrich (USA).

Plant material

S. brasiliensis was collected from fields that belong to Embrapa Pecuária Sul, located in the city of Bagé, Rio Grande do Sul, Brazil (31°21´13.3″S 54°00´36.2″W). A voucher specimen (103517) was stored in the herbarium of the Embrapa, Brasília, Distrito Federal, Brazil.

Extraction and fractionation

Aqueous extract from fresh plant (FP)

Leaves of S. brasiliensis were collected, immediately macerated and mixed with water (100 mg.mL⁻¹) at room temperature. The material was filtered through cotton, concentrated by freeze-drying using a lyophilization apparatus (LP510, Liotop®, Brazil), and stored at −20 °C.

Aqueous extract from dried plant (DP)

Collected leaves of S. brasiliensis were dried at 40 ºC for 2 days. Subsequently, the material was macerated, powdered, and mixed with distilled water (40 °C for 30 min) to make a 100 mg.mL⁻¹ stock solution. The solution was filtered through cotton, concentrated by lyophilization, and stored at −20 ºC.
Alkaloid and non-polar fractions

The alkaloid and non-polar organic fractions (AF and NPF, respectively) were prepared according to the procedure described by Torras-Claveria et al. [53] and Andrade et al. [54] with some modifications. Lyophilized samples were dissolved in 40 mL of MeOH by sonication (Sanders, Brazil). Next, the sample was acidified to pH 2 with 2% H₂SO₄, and the organic fraction was removed by mixing with CH₂Cl₂ followed by decantation. This procedure was repeated three times. The lower fraction was enriched with non-polar compounds and was called NPF. The upper fraction (polar) was basified up to pH 11 with 10% NaOH, and the alkaloids were extracted by liquid-liquid partition with CH₂Cl₂ three times. Afterwards, the lower fractions were enriched with alkaloids compounds and called AF. Finally, the NPF and AF were concentrated in a rotary evaporator (Tecnal, Brazil).

GC-MS analysis of alkaloid extracts

The dried alkaloid-enriched extracts were dissolved in CHCl₃ and directly injected into the GC-MS (Gas Chromatography-Mass Spectrometry) apparatus (injection volume: 1 μL) consisting of a Hewlett Packard 6890 coupled with a mass spectroscopy device (5975 GC/MS, Hewlett Packard, USA) operating in EI (Ionization by Electron) mode at 70 eV. An HP-5 MS column (30 m × 0.25 mm i.d., film thickness 0.25 m) was used. The temperature gradients were as follows: 100–180 °C at 15 °C/min, 180–300 °C at 5 °C/min, 10 min hold at 300 °C and 2 min at 100 °C. The injection temperature was 250 °C, and the flow rate of the carrier gas (helium) was 1 μL/min. A split ratio of 1:5 was applied. Furthermore, the arithmetic retention indexes were calculated by linear interpolation relative to the retention times (tR) of a series of n-alkanes (C7–C30). The obtained values were compared with the published retention index (Kovats index) values [55, 56]. Mass spectra were deconvoluted using AMDIS® 2.64 software (NIST) [54].

Parasitological tests

Animals

Eggs were obtained from the feces of sheep infected and maintained with a monospecific culture of H. contortus. These animals belong to the Embrapa Pecuária Sul, which has own sheep farming, and where the research was carried out. The recovery of eggs was performed according to the method described by Coles et al. [57]. Feces were homogenized in water and filtered through sieves of diminishing pore diameter (180, 90, 68, 38 μm), and the eggs were separated by centrifugation (3000 rpm for 5 min) in a saturated NaCl solution. Immediately, the eggs were washed with distilled water and used in experiments. The experimental protocol was in accordance with the directives of Brazilian National Experiment Control Council (Ethics Committee on Use of Animals – Embrapa Pecuária Sul, protocol under registration number 01/2017). To obtain third stage larvae, the eggs obtained in the same manner as described above were incubated under aerobic conditions in fecal cultures for 7 days at 28 °C and 80–85% humidity. Larvae obtained from these cultures were isolated using a Baermann funnel system and larval cultures were stored at 4–8 °C. After experiments, the animals were treated with anthelmintics and returned to the Embrapa Pecuária Sul fields.

Egg hatching test

The in vitro egg hatching test (EHT) was based on the method described by Coles et al. [58] and standardized by von Samson-Himmelstjerna et al. [59] with modifications. DP and FP were diluted in distilled water and AF and NPF were diluted in 5% tween® 80 aqueous solution. Diluents alone were used as negative controls. Thiabendazole (5 mg.mL⁻¹) was used as positive control, diluted in 5% tween® 80. Approximately 120 H. contortus eggs diluted in water or 5% tween® 80 were incubated with final concentration of 10, 5, 2.5; 1.25, 0.625, 0.312, and 0.156 mg.mL⁻¹ of DP, FP, AF or NPF in a BOD incubator; for 24 h at 28 °C and 80% humidity. Next, 1% lugol’s iodine was added to each well, and the eggs and larvae at the first stage (L1) were microscopically quantified. The test was only validated when hatching in the negative control was greater than 90%. To calculate the percentage of inhibition of larval hatching [58], the formula [(A)/(A + B)] × 100 was used, where A = number of eggs and B = number of larvae. The data were corrected using the Abbott formula [60]. Tests were performed in quadruplicate.

Larval migration inhibition test

Larval migration inhibition test (LMIT) was performed according to Demeler et al., [61, 62] with some modifications. DP and FP were diluted in distilled water, which alone was used as negative control. Levamisole (6.25 μg.mL⁻¹) was used as positive control, diluted in distilled water. About 120 third stage larvae, previously exsheathed with a solution of sodium hypochlorite (0.3%), were incubated in seven different concentrations of plant extract (25; 12.5; 6.25; 3.12; 1.56; 0.78 and 0.39 mg.mL⁻¹) in 24 well plates in a final volume of 1000 μL per well. After 24 h at 28 °C, the total volume (extract + larvae) was deposed into and 25-μm sieves mounted on wells and incubated for more 24 h at 28 °C. After the second incubation, sieves were carefully removed and the remaining non-migrated larvae were washed with distilled water into another well. 1% lugol’s iodine was added to each well and migrated and non-migrated larvae were quantified at 100x magnification. The percentage of non-migrated larvae to the total amount of larvae was calculated for the controls and every concentration tested, as mentioned in EHT, with A = number of non-migrated larvae and B = number of migrated larvae.
and Abbott formula [60] also were used. Tests were performed in quadruplicate.

**Cell citotoxicity assay**

Vero cells were maintained in RPMI media (Cultilab, Brazil) supplemented with penicillin (100 U.L⁻¹) (Sigma-Aldrich), streptomycin (0.01 mg.mL⁻¹) (Sigma-Aldrich), gentamycin (10 µg.mL⁻¹ Sigma-Aldrich), and 10% bovine calf serum (Cultilab). Confluent cells were trypsinized and seed on 96 well plates, totalizing 1.5 x 10⁴ cells/well. Four hours later, when cells were completely adhered, test solutions were added. Two fold serial dilutions were carried out with DP diluted in distilled water to obtain final concentrations ranging from 50 to 0.098 mg.mL⁻¹. The microplates were sealed and incubated at 37 °C in normal atmosphere during 24 h. After this period, resazurin solution (Sigma-Aldrich) was added to each well, to a final concentration of 0.3 mM and the plates were incubated at 37 °C for additional 18 h. The absorbance values were read by dual wavelength using a microplate spectrophotometer (Versa-Max, Molecular Devices) at 570 and 600 nm. All experiments were performed three times each in triplicate. Percentage of viable cells (VC) was calculated as described by Rolón et al., [63]. Percentage of citotoxicity was calculated as 100-VC and was used to calculate EC₅₀. Selectivity index (SI), the ratio between cytotoxicity EC₅₀ and egg hatching or larval migration inhibition EC₅₀, was calculated to evaluate the safety of the extract tested [50].

**Statistical analysis**

An analysis of variance (ANOVA) followed by Tukey’s test was performed on the test results. A p-value ≤0.05 indicated statistical significance. A logistic equation with a variable slope was used to fit the dose-response data by non-linear regression and find the EC₅₀. All analyses were performed after transforming the concentrations into their logarithms (X = log X) and constraining the bottom value to 0% and top value to 100%. All statistical analyses were performed using GraphPrism® version 6.07 (GraphPad Software, USA).

**Abbreviations**

°C: Celsius degree; AF: Alkaloid fraction; DNA: Deoxyribonucleic acid; DP: Dry plant; EC: Effective concentration; EHT: Egg hatching test; FP: Fresh plant; GC: MS: Gas Chromatography-Mass Spectrometry; H. contortus: Haemonchus contortus; LMVT: Larval migration inhibition test; mg.mL⁻¹: Milligram per mL; mM: Millimolar; NaCl: Sodium chloride; nm: Nanometers; NPF: Non-polar fraction; PA: Pyrrolizidine alkaloid; DNA: Deoxyribonucleic acid; DP: Dry plant; VC: Viable cells; SI: Selectivity index

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**Availability of data and materials**

The data generated or analyzed during this study are included in this published article (and its supplementary information files). Data that are not in this article are available from corresponding author on reasonable request.

**Authors’ contributions**

APM, RD, SMS and MIBV conceived the studies. The laboratory experiments and analyses were performed by RD, RD, PAdS, EBG and KMC. The statistical analyses were performed by RD, SMS and EBG. APM and MIBV were responsible for funding acquisition. This work was under supervision of APM and MIBV. The original draft and review of writing were performed by SMS, EBG, APM and MIBV. All authors read and approved the manuscript.

**Ethics approval**

The experimental protocol was in accordance with the directives of Brazilian National Council for the Control of Animal Experimentation (CONCEA) (Ethics Committee on Use of Animals – Embrapa Pecuária Sul, protocol under registration number 01/2017).

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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