Cissampelos sympodialis Eichl. (Menispermaceae), a medicinal plant, presents antimotility and antidiarrheal activity in vivo

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Abstract

Background: Cissampelos sympodialis (Menispermaceae), known as “Milona” has a specific verified medicinal use for the treatment of diarrhea and respiratory tract diseases. This work aims to evaluate the antimotility and antidiarrheal activity of crude ethanolic extract (EtOHE-Cs), and the total alkaloid fraction (TAF-Cs) obtained from aerial parts of C. sympodialis.

Methods: Normal intestinal transit and gastric emptying were used to evaluate antimotility activity. Castor oil-induced diarrhea and castor oil-induced enteropooling were used to evaluate antidiarrheal activity.

Results: The results indicated that EtOHE-Cs has no antimotility activity, but did demonstrate antidiarrheal activity (at 500 mg/kg), and this activity is related to reduction of intestinal fluid accumulation. The TAF-Cs (at 250 and 500 mg/kg) showed antidiarrheal activity by reducing gastrointestinal motility (gastric emptying and normal intestinal transit).

Conclusions: The antidiarrheal activity of C. sympodialis can be attributed to the chemical compounds already isolated and quantified in this species, mainly alkaloids.

Keywords: Gastrointestinal motility, Diarrhea, Cissampelos sympodialis

Background

Diarrhea is characterized by an increase in the frequency of bowel movements, abdominal pain, and bowel discharge of semisolid or watery fecal matter (three or more times in a day) [1,2]. Control of gastrointestinal motility is very complex involving multiple signals, such as nitric oxide (NO), gastrin, ghrelin, prostaglandins, 5-hydroxytryptamine (5-HT), dopamine, catecholamine, and acetylcholine [3,4]. The main causes of diarrhea are imbalances in the pathways just mentioned, as well as infectious agents, plant toxins, and inflammatory problems [5]. Worldwide, the disease affects around 2.2 million people annually, and those most affected are children under the age of five years [6]. Changes in gastric emptying coordination cause not only poor nutrient digestion and absorption, but also the development of diseases. Fully 25-40% of patients with functional dyspepsia report delays in gastric emptying. The development of duodenal ulcers is often related to the stimulation of gastric emptying, since acid content is not completely neutralized in the duodenum [7-10].

Medicinal herbs constitute the majority component of traditional medicines as they are practiced worldwide; this is due to their economic viability, accessibility, and ancestral experiences [11]. Thus, for gastrointestinal disorders, (such as diarrhea) researching medicinal herbs has become very important in developing new therapies.

Cissampelos sympodialis Eichl, belongs to the family Menispermaceae, and is known as “bindweed”, and in Brazil as “milona”, “jarrinha”, or “orelha-de-onça”. It is widely used by Indian tribes, and in folk medicine to treat various diseases such as diarrhea, diseases of the

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genitourinary tract, and especially respiratory tract dis-
eases such as asthma [12]. The species was selected for
study because of ethno-pharmacological and chemotaxo-
nomic criterion.

Several chemical compounds belonging to the alka-
loids class have been isolated from the specie’s leaves
and roots, as examples the: bisbenzylisooquinolinic (warif-
teine, methylwarifteine, roraimine, and simpodialine);
morphinic (milonine); aporphinic (laurifolin) and oxo-
aporphinic (liriodenine) alkaloids [13,14]. Quality control
studies have shown that both alcoholic fractions of the
leaves (AFLs), and alcoholic fractions of the roots (AFRs)
present alkaloids as their principal compounds, being
warifteine the chemical marker of both AFL and AFR
[15]. Crude ethanolic extract (EtOHE-Cs) was standard-
ized using warifteine and methylwarifteine as markers
that were also found in the total alkaloid fraction (TAF-
Cs) [16]. The aqueous fraction of the EtOHE-Cs has
shown spasmyloic activity on tracheal smooth muscle,
and this activity involving inhibition of phosphodiester-
ase (PDE) and increased levels of cyclic adenosine
monophosphate (cAMP) in guinea pig trachea. Warif-
teine, a bisbenzylisooquinolinic alkaloid obtained from C.
sympodialis also has shown spasmyloic activity by in-
hibition of calcium channels (in the rabbit thoracic
aorta) and activation of potassium channels (in the rat
thoracic aorta) [13, 17, 18]. Thus, considering the studies
cited above and other studies reporting the antidiarrheal
activity of alkaloids [19] probably alkaloids of C. sympo-
dialis can be used in diarrhea therapy. Based on its
popular use and its spasmyloic activity “in vitro”, the
aim of this study was to evaluate the antimotility and
antidiarrheal activities of EtOHE-Cs and TAF-Cs
“in vivo” obtained from aerial parts of C. sympodialis.

Methods

Materials

Metoclopramide hydrochloride 10 mg (SANOFI-
AVENTIS®; Brasil); loperamide hydrochloride 2 mg
(JANSSEN-CILAG®; Brasil); phenol red (VETEC®;
Brasil), charcoal meal (VETEC®; Brasil); and Tween 80
(MERCK®; Germany).

Plant material and extraction

Aerial parts of C. sympodialis were collected from the
garden of the “Centro de Biotecnologia” (CBIOTec/
UFPB) in March 2013, and identified by Dr. Maria de
Fátima Agra (Laboratório de Farmacobotânica - CBIOT-
ec/UFPB). A voucher specimen was deposited in the
“Herbario Lauro Pires Xavier” Herbarium, No. 1456. To
obtain the crude ethanolic extract, dried and pulverized
material from aerial parts of C. sympodialis (4000 g)
were subjected to maceration in 95% ethanol for
72 hours. After extraction, the extractive solution was
concentrated in a rotary evaporator under reduced
pressure at a temperature of 45°C, yielding 300 g of

![Image](image.png)

**Figure 1** Effect of oral administration of EtOHE-Cs (a) and TAF-Cs (b) on gastric emptying in mice. (a) ANOVA: F(5,27) = 6.803 (p <0.05) (n = 5–7) followed by Dunnett’s test (** p <0.01 compared to the 12% Tween solution group). (b) ANOVA: F(5,20) = 40.29 (p <0.05) (n = 5–7) followed by Dunnett’s test (*** p <0.001 compared to the 12% Tween solution group).
crude ethanolic extract (EtOHE-Cs). A 100 g aliquot of the EtOHE-Cs was solubilized in an acid solution (3% HCl), and filtered with filter paper. The resulting acid solution was subjected to liquid-liquid partition with dichloromethane. The dichloromethane layer was discarded, and the acid layer was basified with NH₄OH to pH = 9, with a subsequent extraction into chloroform. The chloroform phase was filtered, anhydrous sodium sulfate was added, and the liquid was concentrated in a rotary evaporator resulting in the TAF-Cs (4.8 g).

Experimental animals
Swiss adult male mice (Mus musculus) weighing between 25-35 g were used for the experiments. The animals from the “Biotério Professor Thomas George” (UFPB) were kept at temperatures between 23-25°C, with a 12-hour light/dark cycle in the animal house. The animals were fed Labina, and water ad libitum. For the experiments, they were randomly distributed into different experimental groups. All experiments were started in the morning, and the experimental procedures were approved by the “Comitê de Ética em Pesquisa Animal” (CEPA/CBIOTec/UFPB), and recorded as No. 0705/06, in accordance with international principles for research with laboratory animals [20].

Effect of C. sympodialis on gastric emptying
Adult mice, (fasted for 12 h) were randomly divided into seven groups (n = 7). Mice in the first group received 12% Tween 80 solution - vehicle (10 mL/kg), the second group received metoclopramide (30 mg/kg), and the other groups received 62.5, 125, 250, and 500 mg/kg of EtOHE-Cs, or TAF-Cs, the seventh group (zero time control) received saline solution 0.9%, all by via oral (v.o.). After 1 h of administration (extract, fraction and drugs), a suspension of phenol red marker (0.05%) in carboxymethylcellulose (1.5%) (10 mL/kg) was also given to each animal (v.o.). The zero time control group was euthanized (by cervical dislocation) immediately after the administration of the marker, and the other groups, after 30 min. The abdominal cavity was opened, the pylorus and the distal portion of the esophagus were clipped, the stomach was removed and opened, and its contents were washed with 7 mL of distilled water. The gastric contents collected were centrifuged at 450 g for 15 min, and 1 mL from the supernatant was mixed with 1 mL of 0.025 M NaOH (pH = 12). Afterwards, 150 μL of the homogenate was pipetted in duplicate in a 96 well plate, and a spectrophotometric reading was made using a 560 nm filter. The results were expressed as the percentage of gastric emptying.
compared to the zero time control using the formula below [21].

\[
\% \text{ gastric emptying} = 100 - \frac{\text{mean absorbance of sample}}{\text{mean absorbance of the zero time control group}} \times 100
\]

**Effect of *C. sympodialis* on normal intestinal transit**

The method previously described by Stickney and Northup [22] was used with modifications. Adult mice were fasted for 24 hours, and were randomly divided into six groups \((n = 7)\). Mice in the first group received 12% Tween 80 solution - vehicle (10 mL/kg); the second group received metoclopramide (30 mg/kg) or loperamide (5 mg/kg); and the other groups received 62.5, 125, 250, and 500 mg/kg of EtOHE-Cs, or TAF-Cs, respectively, all by the oral pathway (v.o.). At 1 hour from these administrations, a suspension of charcoal meal (0.5%) in carboxymethylcellulose (0.5%) (10 mL/kg) was given to each animal (v.o.). After 30 minutes, the animals were euthanized by cervical dislocation to remove the small intestine. The transit percentage was calculated on the basis of distance traveled by the charcoal meal as divided by the total length of the intestine, using the formula below.

\[
\% \text{ transit} = \frac{\text{distance traveled by charcoal meal}}{\text{total length of the intestine}} \times 100
\]

**Effect of *C. sympodialis* on castor oil-induced diarrhea**

For the evaluation of antidiarrheal activity was used as a basis the methodology described by Awouters and collaborators [23] with some local modifications. Adult mice were fasted for 12 h and were randomly divided into six groups \((n = 5–8)\). Mice in the first group received 12% Tween 80 solution - vehicle (10 mL/kg), the second group received loperamide (5 mg/kg), and the other groups received 62.5, 125, 250, and 500 mg/kg of EtOHE-Cs, or TAF-Cs, respectively, all by oral pathway (v.o.). At 1 hour from these administrations, castor oil was given to each animal (v.o.) 10 mL/kg. Following the administration of castor oil, the animals were placed in separates cages containing transparent blotting papers for observation of the total number of feces, and their quantification (liquid, semi-solid, and solid) during 4 h. After this, all the animals were euthanized by cervical dislocation. The following parameters were monitored: evacuation classification – 1 (normal stool), 2 (semi-solid stool), and 3 (watery stool) and evacuation index (EI), expressed according to the formula:

\[
\text{EI} = 1 \times (\text{no. of type 1 stools}) + 2 \times (\text{no. of type 2 stools}) + 3 \times (\text{no. of type 3 stools})
\]

**Table 1 Effect of oral administration of EtOHE-Cs, and loperamide on castor oil induced diarrhea in mice**

| Treatment (v.o.)       | Dose (mg/kg) | Evacuation Index (EI) |
|------------------------|--------------|-----------------------|
| 12% Tween 80 solution  | -            | 13.0 (12.0 - 18.0)    |
| Loperamide             | 5            | 0.0 (0.0 - 1.0)**     |
| EtOHE-Cs               | 62.5         | 12.0 (7.0 - 14.0)     |
| EtOHE-Cs               | 125          | 13 (9.0 - 19.0)       |
| EtOHE-Cs               | 250          | 11.5 (8.0 - 17.0)     |
| EtOHE-Cs               | 500          | 4.0 (2.0 - 5.0)*      |

Data are presented as median (minimum value – maximum value). Kruskal-Wallis test followed by Dunn’s multiple comparison test (*) \(p < 0.05\), ** \(p < 0.005\) compared to the 12% Tween solution group. The EtOHE-Cs reduced evacuation index (500 mg/kg) compared to control group (12% Tween 80 solution).

**Effect of *C. sympodialis* on castor oil-induced enteropooling**

Animals (adult male mice) were fasted for 12 hours and were randomly divided into three groups \((n = 7)\). Mice in the first group received 12% Tween solution 80 - vehicle (10 mL/kg), the second group received loperamide (5 mg/kg), and the third group received 500 mg/kg of EtOHE-Cs, (the only dose that showed antidiarrheal activity), all by oral pathway (v.o.). At 1 h from the administrations, the animals were euthanized by cervical dislocation, laparotimized, and then the pyloric and caecal ends of the small intestine were tied and the intestines were removed. The content of each intestine was measured in a graduated measuring cylinder, and the volume was noted according to Ezeja and Anaga [24].

**Statistical analysis**

The parametric data were expressed as the mean ± standard deviation (SD) and non-parametric data were expressed as median (minimum value – maximum value).

**Table 2 Effect of oral administration of TAF-Cs, and loperamide on castor oil induced diarrhea in mice**

| Treatment (v.o.)       | Dose (mg/kg) | Evacuation Index (EI) |
|------------------------|--------------|-----------------------|
| 12% Tween 80 solution  | -            | 15.0 (10.0 - 23.0)    |
| Loperamide             | 5            | 0.0 (0.9 - 2.0)***    |
| TAF-Cs                 | 62.5         | 13.5 (6.0 - 16.0)     |
| TAF-Cs                 | 125          | 6.0 (4.0 - 8.0)       |
| TAF-Cs                 | 250          | 1.0 (0.0 - 5.0)**     |
| TAF-Cs                 | 500          | 1.0 (0.0 - 2.0)**     |

Data are presented as median (minimum value – maximum value). Kruskal-Wallis test followed by Dunn’s multiple comparison test (** \(p < 0.05\), *** \(p < 0.001\) compared to 12% Tween solution group). The TAF-Cs reduced evacuation index (250 and 500 mg/kg) compared to control group (12% Tween 80 solution).
value). This data was subjected to variance analysis (ANOVA), followed by a Dunnett’s test (parametric) or Kruskal-Wallis test followed by Dunn’s multiple comparison test (non-parametric). The minimum level of significance was p < 0.05 in all analyses. For the data processing, INSAT (GraphPad Software® Inc., San Diego, CA, USA) software was used.

Results and discussion

Effect of *C. sympodialis* on gastrointestinal motility: normal intestinal transit and gastric emptying

The antimotility activity of EtOHE-Cs and TAF-Cs was investigated by measuring gastric emptying, and normal intestinal transit in mice. Earlier studies have reported that antimotility and antidiarrheal properties of medicinal plants are due to tannins, alkaloids, saponins, and sterols [25]. Normal intestinal transit and gastric emptying were not altered for any dose (62.5, 125, 250 and 500 mg/kg) evaluated for EtOHE-Cs, however TAF-Cs (at 250 and 500 mg/kg) reduced gastric emptying and normal intestinal transit, as expressed in Figures 1a, 1b, 2 and 3. The results suggest that *C. sympodialis* alkaloids affect gastrointestinal motility, and are likely to present spasmolytic activity “in vivo”. Though showing antimotility activity, we commenced investigations of antidiarrheal activity, because diarrhea is not only caused by motility deregulation. It is also caused by microorganism infections, hyper secretion of intestinal fluids, and inflammatory bowel disease [26].

Effect of *C. sympodialis* on castor oil-induced diarrhea and enteropooling

Diarrhea is a disease that has many clinical signs such as hyper-propulsive motility of gastrointestinal tract, and hyper-secretion throughout the intestinal mucosa. Animal models are commonly used to induce experimental diarrhea [26, 27], and to study a plants’ mechanisms of action and active principles. Castor oil, prostaglandin E$_2$ (PGE$_2$), and heat-labile enterotoxin are commonly used to induce diarrhea in animals. The induction of diarrhea by castor oil is recommended for study of the anti-secretory and antimotility potential of medicinal plants. “Castor oil diarrhea” is due to its active metabolite, ricinoleic acid that is released by the action of intestinal lipases [28].

The liberation of ricinoleic acid results in irritation and inflammation of the intestinal mucosa, and is associated with the release of nitric oxide, prostaglandin, and other autacoids [28, 29]. The enteropooling induced by castor oil, probably occurs through stimulation of cyclic AMP/GMP production, and phosphorylation of cystic fibrosis transmembrane conductance regulators (CFTRs). This consequently leads to intestinal motility stimulation and to increased secretion of fluids and electrolytes (mainly Cl$^-$ and Na$^+$) [30,31].

In castor oil induced-diarrheic animals, the EtOHE-Cs in its highest dose (500 mg/kg) significantly reduced the evacuation index as can be seen in Table 1, and TAF-Cs showed similar antidiarrheal effects at doses of 250 and 500 mg/kg, as can be seen in Table 2.

The EtOHE-Cs does not have antimotility activity. Due to this, in another step we investigated if the extract had intestinal anti-secretory activity. For the enteropooling study, the EtOHE-Cs (at 500 mg/kg) significantly reduced the intraluminal fluid volumes of the intestinal contents (Figure 4), suggesting that the EtOHE-Cs antidiarrheal activity involves the reabsorption of water and electrolytes (such as Na$^+$), and probably, the involvement of prostaglandins [28].

Conclusions

The antidiarrheal activity of *C. sympodialis* can be attributed to chemical compounds already isolated and quantified in this species, such as flavonoids and principally the alkaloids warifteine and methylwarifteine [32]. These compounds are known to inhibit autacoid and prostaglandin release [33]. Inhibition of prostaglandin E$_2$
(PGE$_2$) is known to reduce secretory response in the intestine, and to inhibit gastrointestinal motility [34].

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

LMB directed and IRPs designed the study. IRPs, FDFM AFM, ASSCL, JMFB and LMB performed the experiments. IRPs and FDFM drafted the manuscript. JMFB and LMB corrected the manuscript. All authors have read and approved the final manuscript.

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