Antidiarrheal activity of *Solanum asterophorum* in mice

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Abstract: Several species of *Solanum* are used in folk medicine to treat diarrhea. Therefore, the aim of this study was to investigate and compare possible antidiarrheal activity of methanol extracts from roots (Sast-MeOH R) and leaves (Sast-MeOH L) of *Solanum asterophorum* Mart., Solanaceae, in mice. Sast-MeOH R was shown to significantly and dose-relatedly inhibit the frequency of both solid (ED50 309.6±28.5 mg/kg) and liquid (ED50 152.1±32.5 mg/kg) stools. Conversely, Sast-MeOH L significantly inhibited solid stool frequency only when dosed at 500 and 750 mg/kg (48.7±7.4 and 42.3±9.8%, respectively), but also significantly and dose-relatedly inhibited liquid stools (ED50 268.4±35.2 mg/kg). Thus, Sast-MeOH R was twice as potent as Sast-MeOH L in diarrhea inhibition. Neither extracts (when dosed up to 500 mg/kg) inhibited intestinal transit. However, both extracts significantly and dose-relatedly inhibited intestinal fluids, and Sast-MeOH R (ED50 38.3±10.4 mg/kg) was again twice as potent as Sast-MeOH L (ED50 78.6±6.4 mg/kg). Results suggest that antidiarrheal effects of Sast-MeOH R and Sast-MeOH L involve changes on intestinal secretion. In addition, active metabolites with antidiarrheal activity may be more concentrated in the roots of this species. However further studies are needed to elucidate the action mechanism involved in this activity.

Keywords: antidiarrheal effect in vivo study leaves roots *Solanum asterophorum*

Introduction

Diarrhea is one of the most common diseases in children worldwide and is characterized by an increase in the number of stools (3 or more/24 h), increased fluidity of stools and/or the presence of blood/mucus in these (Mathan, 1998). It affects people of any age, but is in childhood that this condition may cause higher mortality. It is the third most common cause of disease in children in developing countries and is responsible for about one third of all hospitalizations among children under five years (Souza, 2000). Many plants are used as antidiarrheal in folk medicine, such as *Psidium guajava* (“guava”), *Stachytarpheta cayenensis* (“Brazilian tea”), *Eugenia uniflora* (“Brazilian cherry”), *Anacardium occidentale* (“cashew”), *Mangifera indica* (“mango”) and *Egletes viscosa* (“macela”) (Almeida et al., 1995; Agra et al., 2008).

There are many Angiosperm families in Northeast Brazil, among which Solanaceae A. L. Jussieu stands out, with 96 genera and 3000 species (Hunziker, 2001). The genus *Solanum* L. is the most representative of this family, with circa 1400 species (Bohs, 2005) and 5000 described epithets (Nee, 1999), inhabiting tropical and subtropical regions of the world (Agra & Bhattacharyya, 1999). Several species of *Solanum* are known in folk medicine for showing antidiarrheal activity, such as *Solanum marginatum* (Abebe, 1986), *Solanum khasianum* (Jain & Puri, 1984), *Solanum torvum* (Dominguez & Alcorn, 1985) and *Solanum xanthocarpum* (Jain & Puri, 1984).

*Solanum asterophorum* Mart., Solanaceae, is a shrubby species, popularly known as “jurubeba-de-fo”. In Brazil, it is found in Paraíba and Bahia, with indications of folk medicine use against liver dysfunctions (Agra & Bhattacharyya, 1999). Pharmacological studies show that methanol extracts obtained from aerial parts and roots of *Solanum*
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*S. asterophorum* are bioactive for *Artemia salina* but have no bioactivity for mollusks (Silva et al., 2007). When studying acute toxicity, Oliveira (2006) showed that the methanol extract obtained from leaves (Sast-MeOH) of this species did not produce nor induce behavioral changes and death in mice, while Oliveira et al. (2006) demonstrated that the same extract (Sast-MeOH) and the alkaloid isojuripidine isolated from its aerial parts (Silva et al., 2005) did not induce erythrocyte hemolysis in rats, but induced relaxation in guinea pig ileum through voltage-dependent calcium channels type 1.2 (Ca 1.2).

Since many *Solanum* species show antidiarrheal activity and *S. asterophorum* have shown spasmolytic activity in guinea pig ileum, i.e., reduced its contractibility, we decided to investigate and compare possible antidiarrheal activity of methanol extracts from roots (Sast-MeOHR) and leaves (Sast-MeOHL) of this species, using castor oil-induced diarrhea models in mice.

**Material and Methods**

**Plant material**

*Solanum asterophorum* Mart., Solanaceae, was collected in the municipality of Areia, Paraíba, Brazil, and identified by Dr. Maria de Fátima Agra at the Laboratório de Tecnologia Farmacêutica “Prof. Delby Fernandes de Medeiros”, Universidade Federal da Paraíba. The voucher specimens are deposited in the Herbarium Prof. Lauro Pires Xavier (JPB), UFPB, and duplicates are deposited in the Reference Collection of LTF/UFPB under the identification code “Agra 6002”. Powdered aerial parts and roots were extracted with MeOH in a Soxhlet apparatus. The extract was concentrated under vacuum in rotaevaporator. The crude residue was subjected to experiments.

**Animals**

Female and male Swiss mice (*Mus musculus*) from the Central Animal Laboratory, Universidade Federal de Alagoas, weighing between 25 and 35 g, were used. Previous to the experiments, the animals were kept under strict control feeding with a balanced diet based on food pellet type and had free access to water, both room ventilation and temperature (21±1 °C) were controlled and constant. All experimental procedures with *Solanum asterophorum* were submitted and approved by the Ethical Committee in Research of UFAL (No. 027241/2008 11).

**Substances**

Atropine and Cremophor® were obtained from Sigma-Aldrich(USA). Carboxymethylcellulose or castor oil were obtained from Formula, Brazil. Loperamide was obtained from Janssen Cilag Farmacêutica Ltda, Brazil, and the activated charcoal obtained from Proquímios, Brazil.

**Preparation of extracts for pharmacological tests**

The crude methanol extracts were obtained from roots (Sast-MeOH) and leaves (Sast-MeOHL) of *Solanum asterophorum*. These were solubilized in Cremophor® (3%) and diluted in distilled water to obtain stock solutions (500 mg/mL). Upon completion of the experimental protocols the solutions were diluted with distilled water or saline, according to the requirement of each protocol.

**Pharmacological behavioral screening and determination of LD50 of Sast-MeOH**

The Sast-MeOH extract was administered orally (p.o.) or intraperitoneally (i.p.) to groups of five males and five females (one dose per group), which fasted for 12 h. Simultaneously, control animals received 10 mL/kg plus Cremophor®. General signs and symptoms of toxicity, such as analgesia, contortions, aggression and diarrhea, were recorded for 3 h. The animals were also evaluated at 24, 48 and 72 h in order to observe if there was any death.

**Evaluation of antidiarrheal activity of Sast-MeOH**

Male mice were divided into groups (n=6) and gavaged with 0.9% saline solution 10 mL/kg plus Cremophor® (negative control), loperamide 10 mg/kg (positive control) and Sast-MeOH or Sast-MeOHL (doses ranging from 62.5 to 750 mg/kg). After 1 h, 0.01 mL of castor oil per gram was administered orally to each animal to induce diarrhea and the consistency of dung pats was assessed for 3 h, classifying them according to consistency in solids or liquids. ED50 values (dose of a substance that produces 50% of its maximum effect) were obtained by nonlinear regression and expressed as a percentage of the mean±SEM.

**Intestinal transit in mice**

Male mice were divided into groups (n=6) and gavaged with 0.9% saline solution 10 mL/kg plus Cremophor® (negative control), atropine 2 mg/kg (positive control) and Sast-MeOH or Sast-MeOHL (doses ranging from 62.5 to 750 mg/kg). After 1 h, 0.01 mL of castor oil per gram was administered orally to each animal to induce diarrhea and the counting and consistency of dung pats was assessed for 3 h, classifying them according to consistency in solids or liquids. ED50 values (dose of a substance that produces 50% of its maximum effect) were obtained by nonlinear regression and expressed as a percentage of the mean±SEM.
obtained from roots (Sast-MeOH R) and leaves (Sast-MeOHL) of

0.9% saline solution 10 mL/kg plus Cremophor® (negative
control) and Sast-MeOH R or Sast-MeOHL (doses ranging
from 25 to 125 mg/kg). After 30 min, 2 mL of castor oil was
administered per animal. After other 30 min, the
mice were euthanized, the small intestine was dissected
from pylorus throughout cecum and the contents were
collected into a beaker to assess the volume of fluid.
ED50 values were calculated as previously.

Statistical analysis

All results were expressed as a percentage of
the mean±SEM, and differences between means were
statistically analyzed through t-test (non-paired) or
one-way analysis of variance (ANOVA) followed by
Bonferroni correction with p<0.05 values considered
significant. ED50 and Emax values were assessed
through nonlinear regression. All data were analyzed
using GraphPad Prism version 3.03 software (GraphPad
Software Inc., San Diego, CA, USA).

Results and Discussion

In this study we investigated and compared a
possible antidiarrheal activity of crude methanol extract
obtained from roots (Sast-MeOH R) and leaves (Sast-
MeOH L) of Solanum asterophorum Mart., Solanaceae, in
mice. For the first time, the strong antidiarrheal activity
displayed by Sast-MeOH R and Sast-MeOH L extracts was
demonstrated.

Since many species of Solanum are used in
diarrhea treatment (Abebe, 1986; Jain & Puri, 1984;
Dominguez & Alcorn, 1985) and Solanum asterophorum
leaf extracts exhibit antispasmodic activity (Oliveira et
al., 2006), we hypothesized that these extracts might
contain antidiarrheal active metabolites.

The most frequent metabolites in Solanum
species are from the alkaloid family, the steroidal
alkaloids solasonine and solamargine being found in
more than 100 of its species (Blankemeyer et al., 1998;
Mesia-Vela et al., 2002), and toxic activity is attributed
to these alkaloids (Jadhay et al., 1981; Blankemeyer et
al., 1998). Due to the great interest by folk medicine in
compounds from Solanaceae plants, many studies have
been developed trying to evaluate the toxicological
activity of Solanum species (Shakunda & Chester, 1976;
Maruo et al., 2003; Heo & Lim, 2005).

Oliveira (2006) showed that Sast-MeOH R
extract did not produce behavioral changes nor did it
induce death in mice in an acute toxicity study. Since
this is a pioneer work on S. asterophorum roots, the
acute toxicological potential of Sast-MeOH R extract was
assessed and a pharmacological monitoring on diarrhea
experimental models was conducted.

The administration of Sast-MeOH R extract
orally at doses between 2500 and 5000 mg/kg and
intraperitoneally at doses between 1000 and 2000 mg/
kg in mice of both sexes also did not produce toxicity-
characteristic behavioral changes such as sedation,
agression, urination, diarrhea, convulsion, loss of
corneal reflex and ear, piloerection and writhing, nor
induced death. Thus, it was not possible to assess LD50
for the tested extracts. The fact that both the root and leaf
extracts are devoid of systemic toxicity allowed us a safe
choice of the doses used in the experimental protocols
performed in order to investigate the antidiarrheal effect
of these extracts.

Diarrhea results from rapid movement of fecal
matter through the large intestine (Guyton & Hall,
2006). To restore this process, many patients require
antidiarrheal therapy in order to increase flow resistance
and mucosal absorption and decrease secretion
(Akindele & Adeyemi, 2006). To verify the antidiarrheal
effect of Sast-MeOH R and Sast-MeOH L, castor oil was
used since it is widely employed for screening drugs
with this property. One of the advantages of this model
is the high reproducibility of the evacuation of stool
formed in less than one hour after the administration
of laxatives (Borrelli et al., 2006). The results of oral
administration of loperamide (10 mg/kg) and Sast-
MeOH R (250, 500 and 750 mg/kg) showed that there
was significant dose-related inhibition of defecation
frequency (97.9±2.1, 34.3±5.6, 47.8±9.7 and 70.1±6.3,
respectively) when compared to the negative control
(0.9% saline solution+Cremophor®) (Figure 1A).
Regarding liquid stools, both loperamide (10 mg/
kg) and Sast-MeOH R (125, 250, 500, and 750 mg/kg)
showed significant, dose-related inhibition percentages
of 43.6±4.4, 58.0±8.5, 80.6±12.6 and 100, respectively
(Figure 1B). On the other hand, Sast-MeOH L extract
(Figure 2A) showed significant inhibitory effect on
defecation frequency only when dosed at 500 and
750 mg/kg (48.7±7.4 and 42.3±9.8%, respectively).


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Regarding liquid stools, Sast-MeOH extract showed inhibitory effect when dosed at 250, 500 and 750 mg/kg, with significant dose-related inhibition frequencies of 32.9±10.1, 80.6±11.1 and 88.4±5.2%, respectively (Figure 2B). Regarding defecation frequency, only Sast-MeOH produced dose-related inhibition, with 309.6±28.5 mg/kg ED50. However, for liquid stools, Sast-MeOH extract (EDSO 152.1±32.5 mg/kg) was more effective than Sast-MeOH extract (EDSO 268.4±35.2 mg/kg) in inhibiting diarrhea. Therefore, both Sast-MeOH and Sast-MeOHH produced significant inhibitory effect on diarrhea in mice in this induction model, as observed by reducing both solid and liquid stool frequencies, suggesting that both extracts have active metabolites with antidiarrheal activity. Conversely, the root extract showed stronger antidiarrheal effect than that from the leaves, suggesting that the active antidiarrheal metabolites are more frequent in the former than in the latter.

Clinically, diarrhea is a consequence of intestinal function disorders which lead to excessive intestinal secretion, impaired intestinal absorption and/or rapid transit (Menezes et al., 1994). Therefore, the antidiarrheal effect of the extracts was studied with experimental castor oil-induced diarrhea models and castor oil-induced intestinal transit and intestinal fluid accumulation in mice. Drugs that inhibit intestinal transit in pathophysiological states may be effective in relieving diarrhea (Borrelli et al., 2006). Activated charcoal, employed in the gastrointestinal transit model as a marker, has been used for over 60 years as a simple tool for evaluating the effect of laxatives (Gaginella et al., 1994). This method is an indicator of the maximum distance traveled by the marker (activated charcoal) through its administration and the assessment of its path in the small intestine over a period of time (Gurgel, 2000). According to this model, the results of oral administration of atropine (2 mg/kg) show that there was significant inhibition of the distance traveled by charcoal in the intestine of mice of circa 32.7±3.6%, when compared to the negative control (0.9% saline+Cremophor®). Still, neither Sast-MeOH R nor Sast-MeOH L extracts reduced the distance travelled by the marker when dosed at 125, 250 or 500 mg/kg (data not shown).

Though it is common belief that the small intestine plays only an absorptive role, it also secretes water and electrolytes. This enhanced secretory capacity is clearly shown through the administration of large amounts of castor oil (Gurgel, 2000). Thus, we assessed the effects of Sast-MeOH and Sast-MeOHH extracts in mice using a castor oil-induced intestinal fluid accumulation model and evaluated the fluid amount in comparison to the control group. Loperamide (69.2±1.5%), Sast-MeOHH and Sast-MeOHH extracts inhibited significantly and dose-relately the intestinal fluid content induced by castor oil dosed at 25, 50, 75, 100 and 125 mg/kg in comparison to negative control (0.9% saline+Cremophor®). When compared to the negative control, inhibition values for the Sast-MeOHH extract were 21.6±7.5, 30.4±4.9, 42.2±2.4, 60.8±3.0 and 71.1±2.4%, respectively, and 37.2±6.4, 52.9±4.0, 51.0±4.5 and 59.3±2.5% for Sast-MeOHH, respectively (Figure 3). Sast-MeOHH extract (EDSO 38.3±10.4 mg/kg) was about twice as effective as Sast-MeOHH (EDSO 78.6±6.4 mg/kg) in inhibiting intestinal fluid content, reinforcing the idea that the secondary metabolites with antidiarrheal activity are more frequent in the roots than in the leaves.

The most important conclusions of this study are the inedited demonstrations that Solanum asterophorum Mart. displays antidiarrheal activity, which surely brings great contribution to the pharmacology of this species, as well as showing that the root extract has more effective antidiarrheal activity than the leaf extract. Since steroidal alkaloid isojuripidine was isolated from the aerial parts of S. asterophorum, part of this activity can be attributed to it. Further studies are needed, including verifications of the amount and type of root alkaloids.

Figure 1. Antidiarrheal effect of Sast-MeOHH extract on castor oil-induced diarrhea model in mice (n=6). A. Percentage frequency of defecation, and B. Percentage of liquid stools. Columns and vertical bars represent the percentage of mean and SEM, respectively. One-way ANOVA followed by Bonferroni, *p<0.05 and **p<0.001 (Saline vs. Loperamide/Sast-MeOHH).
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Figure 2. Antidiarrheal effect of Sast-MeOH$_A$ extract on castor oil-induced diarrhea model in mice (n=6). A. Percentage frequency of defecation, and B. Percentage of liquid stools. Columns and vertical bars represent the percentage of mean and SEM, respectively. One-way ANOVA followed by Bonferroni, *p<0.05 and **p<0.001 (Saline vs. Loperamide/Sast-MeOH$_L$).

Figure 3. Effect of Sast-MeOH$_A$ A and Sast-MeOH$_B$ B extracts on castor oil-induced intestinal fluid accumulation in mice (n=6). Columns and vertical bars represent the percentage of mean and SEM, respectively. One-way ANOVA followed by Bonferroni, *p<0.05 and **p<0.001 (Saline vs. Loperamide/extracts).

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