Pharmacognostic analyses and evaluation of the in vitro antimicrobial activity of *Acmella oleracea* (L.) RK Jansen (Jambu) floral extract and fractions

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Received 7 November, 2014; Accepted 15 January, 2015

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**Key words:** Antimicrobial, phytotherapy, flowers pharmacognosy, *Salmonella typhi*.

**INTRODUCTION**

*Acmella oleracea* (L) RK Jansen, vernacular Jambu, has motivated the interest of researchers because of its therapeutic potential, inducing projects in different areas linked to health sciences, such as medicine, dentistry...
and pharmacy. A. oleracea is a perennial, herbaceous flowering shrub common in Northern Brazil, particularly in Para State (Coutinho et al., 2006; Silva and Santos, 2011). Its chemical constitution described in the literature includes the alkamide spilanthol, α and β-amyrinester, stigmasterol, miricilic alcohol glycosides, sitosterol, saponins and triterpenes (Lemos, 2012).

Although, the entire plant is medicinally used, it is mainly in the flowers where the highest amount of Spilanthol (Wongsawatkul, 2008) occurs (Cavalcanti, 2008; Nigrinis et al., 1986), using guinea pigs, confirmed the main biological activity of this plant organ; the local anesthetic effect. The same authors also reported the flavoring, insecticide, bactericide and healing action of the floral extract, both in oral mucosa and on the skin. No scientific report about the microscopic description of A. oleracea flowers was found in the surveyed literature. This work reports the result of the pharmacognostic investigation on the A. oleracea floral drug and its Ethanol extract. In addition, the in vitro antimicrobial activity of EEFAO and its fractions on different bacteria and fungi species were reported.

MATERIALS AND METHODS

Plant materials

Young leaves and flowers of A. oleracea (L.) RK Jansen were acquired in Manituba, Para State, metropolitan region of Belem (01° 21’ latitude South and 48° 20 longitude West). A voucher is deposited at the Herbarium of the Botany Laboratory, Embrapa Amazonia Oriental, Belem - Para, registered as IAN 188444, in October, 2012.

Macroscopic and microscopic description

Macroscopic characterization of the flowers and leaves was performed with bare eye, according to parameters described by Esau and Silva (1976). For microscopic description, semi-permanent cuts were prepared using fresh leaves and young flowers. The plant material was sliced transverse in longitudinally by hand. The samples were clarified in 10% aqueous Sodium Hypochlorite and stained with Methylene Blue and Safranin. The cuts were observed using a Nikon optical microscope (Eclipse 50i) equipped with a Motic® camera (Moticam 2300) and the pictures were processed using Motic Image Plus 2.0® software. The photomicrographs obtained at 10X and 40X were analyzed in comparison to literature data.

Preparation of extract

To obtain the Ethanol extract, 3.8 kg of cleaned A. oleracea flowers were dried under circulating hot air (Quimis Q317B) at 40°C ± 2°C until constant weight of an aliquot. The dried plant material was grinded in a Wiley knives mill using a sieve of medium mash. The obtained herbal drug weighed 426.1 g, which was macerated in 2.5 L 70% Ethanol in a stainless steel container for seven days. The obtained tincture was then filtered and concentrated under reduced pressure on a rotatory evaporator (800 Fisatom). The aqueous residue was frozen and lyophilized (Freeze dryer L101, LIOTOP). The freeze-dried extract was stored under refrigeration until use.

Pharmacognostic essays

The physical-chemical quality control of the plant drug and its extract EEFAO was performed according to the Brazilian Pharmacopeia, 5th edition, and the following tests were performed: particle size distribution, solids contents, pH, ash content and moisture content, respectively (Silva, 2008).

Fractionation

An aliquot of 4.26 g of EEFAO was fractionated by solid-liquid partition using 5 to 7x 50 ml aliquots of increasing polarity solvents: Hexane, Chloroform, Ethyl Acetate and Methanol. This procedure is established in the routine of Phytochemical Analysis Laboratory, Faculty of Pharmaceutical Sciences, Federal University of Pará. The fractions were concentrated on a rotatory evaporator at low pressure (Fisatom 800).

Chromatographic analyses

The chromatographic profile of the samples was obtained by thin layer chromatography (TLC) using silica gel (SIGMA) as stationary phase, mixture of solvents of different polarities as the mobile phase ding according to Wagner and Bladt (2001), to detect Flavonoids, Tannins, Terpene and Alkaloids (ANVISA 2010).

Evaluation of antimicrobial activity

Microorganisms and growth conditions

Bacteria: Salmonella typhi ATCC00259, Enterobacterium faecalis ATCC29212 and Staphylococcus aureus ATCC00577.

Yeast: Candida albicans ATCC0175. The Laboratory of Microbiological Control, Faculty of Pharmaceutical Sciences provided the samples of microorganisms. The bacteria grown in Mueller-Hinton broth (Himedia) at 37°C were kept in Mueller Hinton agar plates at 4°C. The yeasts were grown and maintained in broth and Sabouraud Agar (Himedia).

Antimicrobial activity

The tests were performed using broth micro dilutions techniques and MIC of EEFAO and its fractions were determined as described by Holetz et al. (2012) with modifications. Aliquots of 100 μl of broth and 10 μL of microorganisms adjusted McFarland scale (10^9 Colony Forming Units) were used to evaluate the activity of EEFAO and its fractions at a concentration of 100μg/Ml (Wagner and Bladt, 2001). minimal inhibitory concentration (MIC) was defined as the lowest concentration of the sample, which produces a marked reduction of at least 80% of the tested microorganisms (Wagner and Bladt, 2001).

Statistical evaluation

The data statistical evaluation was performed using Bioestat 5.3, and T Student test with 95% level of confidence.

RESULTS

The macroscopic analysis of A. oleracea (L) RK Jansen
leaves and fresh flowers was performed on the external surfaces of these organs (Figure 1). The microscopic analysis of *A. oleracea* leaves revealed simple membranous leaves showing wavy uniseriate epidermal cells, bicellular or tricellular trichome with basal cell showing rough cuticle and anomocytic stomata. The ventral mesophyll is well-organized showing two layers of palisade parenchyma and several layers of spongy parenchyma. The cross-section of central rib shows a concave-convex profile and two vascular bundles are present in addition to the conducting vessels (Figure 2). The presence of the undifferentiated hypanthium (HP) with various vascular bundles (FV) and differentiated laterally sepals (SP) can be microscopically observed in the longitudinal section of the immature flower buds. The central region of the hypanthium presents uniseriate epidermis, at its ends involucral bracts can be also observed in (BI), peleas (PL) and anthers (AT) (Figures 3 and 4). The pharmacognostic analyses show the following results: the granulometry of the *A. oleracea* herbal drug (dried and grinded flowers). The grinded dried flowers indicates that the sample is a coarse powder since its particles were predominantly retained on the sieve with the highest mash value (1.700 mm) reaching 99.95% of the sample weight (Table 1). The pH of the herbal drug was determined for decoction, after filtration and cooling in a calibrated potentiometer. The result, 5.33, is the middle value of three determinations with standard deviation of ± 0.24, using osmosed water as reference, pH of 6.25 (Table 1).

The determination of total ash present in the sample revealed as a middle value of three experiments 7.07%; standard deviation of ± 0.03% (Table 1). Completing the pharmacognostic analyses, moisture was determined in triplicate yielding, 11.6% standard deviation ± 0.264%. In addition, the middle value of dried residue of the extract is 2.5%; SD ±0.05 (Table 1). All the parameters were determined in triplicate. Chromatographic analyses of EEFAO and its Hexane, Chloroform and Ethyl Acetate fractions where performed using TLC on normal phase silica gel with Hexane/Acetone (80:20) as eluent. The obtained chromatograms, show orange colored bands due to reaction with Dragendorff’s reagent, at Rf value of 0.37. The antimicrobial test using EEFAO and its fractions resulted in significant reduction (p ≤ 0.05) of the colonies number of *Salmonella typhi* ATCC 00259 (Table 2); especially for the chloroform fraction, which inhibited the microorganism growth at a MIC of 31.25 µg/mL of the *Salmonella typhi* ATCC 00259.

**DISCUSSION**

As described in the literature, genus Acmella is part of the Asteraceae family, consisting of Angiosperm plants (Esau, 1976). The macroscopic description of leaves of this plant species discloses phylliform or membranous aspect of the largely oval leaves, being about 3 to 6 cm long and dark green in color (Coutinho et al., 2006; Silva and Santos, 2011). Mature capitula were macroscopically described as oval, irradiated or discoid; with conical receptacle, golden or pink to reddish paleas and obtuse to acuminate apex; ray flowers when present, are ligulate, pistillate; corolla can be white yellow or orange, with two or three lacinios at the apex. The disc flowers are perfect, tubular, white, yellow or orange corolla, four or five acute lacinia and four to five anthers. Oval or ellipsoid cypselae disc is laterally compressed, sometimes with cortical margin present when mature; pappus absent or 1 to 10 weak bristles, tricosted, ellipsoid cypselae ray are usually...
Figure 3. A, B and C - Photomicrographs of structures found in leaves. A. *oleracea* A - demonstrates the structures where the midrib with vascular tissues. B - Demonstrates the mesophyll with two layers of palisade parenchyma and C demonstrates the abaxial surface (bottom) with bay leaf stomata 40×.

Figure 4. A and B - Photomicrographs of *A. oleracea* flowers: A: highlights the common structures to flowers of the same family with SP-sepals; HP-hypanthium; FV-vascular bundles; and B: BI-Involucral bracts; PL-Paleas and AT- anthers. Safranin and Methylene Blue 40×.
Table 1. Results of the pharmacognostic tests of herbal drug and extract of *A. oleracea*.

<table>
<thead>
<tr>
<th>Sample</th>
<th>G+sd</th>
<th>pH+sd</th>
<th>TA± sd</th>
<th>LD ± sd</th>
<th>DR± sd</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>A. oleracea</em></td>
<td>99.95%±0.01</td>
<td>5.33±0.24</td>
<td>7.07%±0.03</td>
<td>11.6%±0.264</td>
<td>2.5%±0.05</td>
</tr>
</tbody>
</table>

G-Granulometry; TA-Total Ash; LD-Loss on drying; DR- Dried residual; sd- standard deviation p ≤ 0.05.

Table 2. Inhibitory activity of chloroform fraction of EEFAO on *Salmonella typhi* ATCC00259.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Microorganism</th>
<th>MIC (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chloroform fraction</td>
<td><em>Staphylococcus aureus</em> ATCC 00577</td>
<td>1000</td>
</tr>
<tr>
<td></td>
<td><em>Enterobacterium faecalis</em> ATCC 29212</td>
<td>1000</td>
</tr>
<tr>
<td></td>
<td><em>Salmonella typhi</em> ATCC 00259</td>
<td>31.25</td>
</tr>
<tr>
<td></td>
<td><em>Candida albicans</em> ATCC 0175</td>
<td>1000</td>
</tr>
</tbody>
</table>

MIC*- Minimum Inhibitory Concentration.

usually present in mature cortical margin. Similar structures were found and described in Figure 1 (Coutinho et al., 2006).

No data about the microscopic structure of the flowers of this plant species was found in the literature. However, Flowers of Flowering Plants have similar microscopical structures like hypanthium, which is the region of the flower that receives seeds after differentiation (Figures 3 and 4). Comparing the microscopic structures of the flowers of this plant, species having similar structures were observed in *Acmella brasiliensis* and *Acmella marajoensis* (Holletz et al., 2002; Coutinho et al., 2006), it is possible to infer that *A. oleraceae* also has differentiated hypanthium for seeds accumulation, as well large caliber vascular bundles another to ensure the maintenance of these reproductive structures during prolonged periods.

The analyses of EEFAO show that the results here reported are in accordance with the parameters described in the Brazilian Pharmacopoeia 5th edition (Silva, 2008). In Brazil, the National Agency for Sanitary Surveillance (ANVISA) requires pharmacognostic tests for quality control, such as chromatographic profile by TLC or phytochemical screening, as criterion for notification or registration of traditional herbal medicines and notification of herbal product (Baccarin et al., 2009). This paper reports the TLC analysis, based on the methodology described by Wagner and Bladt (2010) (ANVISA, 2010), highlighting the substance at Rf 0.37, which reacts positively to Dragendorff’s reagent. Armond (2007) performed a phytochemical investigation of *A. oleraceae* using the same phytochemical methodology, the author reports about a substance with Rf 0.36, in the same chromatographic condition (ANVISA, 2010), which reacts as Flavonoid with Sulfuric Vanillin and fail to react with Iodine and KOH like some Alkaloids he tested. In the present work the substance with Rf 0.36 reacts to Dragendorff’s reagent, as usual for substances containing nitrogen, like alkamides. This observation raises the following question, can alkamides react both as N-containing substance with Dragendorff’s reagent or as bident structure common in Flavonoids allowing the complexation of a metal ion? (Marques et al., 2012). In fact, the reaction of alkamides with Sulfuric Vanillin may involve a nucleophilic attack of Nitrogen atom on the Aldehyde Carbonyl group of Vanillin.

Among the antimicrobial tests against *Salmonella, typhi* showed the best result. This genus of bacteria is usually found in the gastrointestinal tract of domestic and wild animals, especially birds and reptiles. Numerous *Salmonella* serotypes are pathogenic for both animals to humans (Anvisa, 2014; Armond, 2007). It is estimated that 36% of dogs are asymptomatic carriers of this bacteria; In witch, the clinical signs of the disease vary depending on the number of infective organisms and the immune status of the animal as well others adverse factors such as intercurrent illnesses. Young animals or elderly one are the most susceptible to the bacteria, increasing the severity of the infection (Armond, 2007). There is evidence that besides frames of severe enteritis, bacteria can also cause this kind of generalized skin lesions redness, blistering and crusting in immuno-compromised patients (Anvisa, 2007). Considering that the antimicrobial susceptibility to plant extracts, the inhibitory action to be considered promising must show a minimum inhibitory concentrations less than or equal to 100 µg/mL. Samples with MIC extracts ranging from 100 to 500 µg/mL are considerate having moderate antimicrobial activity. However, MIC values greater than 1000 µg/mL characterize the samples microbiologically inactive (Wagner and Bladt, 2001). The crude extract of *A. oleraceae* does not show antimicrobial activity as reported by Holletz et al. (2002) (Wagner and Bladt, 2001). This work reports the antimicrobial activity of the chloroform fraction prepared from the crude extract EEFAO. Prachayasittikul et al. (2009) (Carvalho et al.,...
2003) demonstrated that the chloroform fraction of A. oleracea was able to inhibit the growth of Saccharomyces cerevisiae ATCC 2601 and Streptococcus pyogenes in a MIC 256 μg/mL. In this study Salmonella typhi ATCC00259, had their growth inhibited by a MIC of 31.25 μg/mL of chloroform fraction of EEFAO.

Conclusion

Quality parameters (particle size, total ash content, pH, loss on drying and dried residue) described in this work allow the identification and standardization of the herbal drug, the extract obtained from the flowers of A. oleracea and its fractions, where the presence of a substance reactive to the Dragendorff reagent can be detected, probably spilanthol.

The microscopic description of the flowers of this plant species is very useful because it enables the micrographic characterization of plant drug (Figures 3 and 4). No records of these data were found in the surveyed literature. The result of antimicrobial activity evaluation indicates that the chloroform fraction of EEFAO is able to reduce the visible growth of Salmonella typhi. This observation can justify the use of A. oleracea in the development of therapeutic products or conservatives for foods.

Conflict of interest

All authors declare that they have no conflict of interest.

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