The antioxidant, anticancer and anticoagulant activities of *Acanthus ilicifolius* L. roots and *Lumnitzera racemosa* Willd. leaves, from southeast coast of India

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**ABSTRACT**

The present study was designed to evaluate and determine the phytochemical composition, antioxidant, anticancer and anticoagulant activities of the aqueous extracts of *Acanthus ilicifolius* roots and *Lumnitzera racemosa* leaves from Pichavaram mangrove forest, Tamil Nadu, India based on folklore knowledge. The preliminary phytochemical screening revealed the presence of the following classes of bioactive compounds: phenols, flavonoids, alkaloids, terpenoids, sterols, tannins, carbohydrates, cardiac glycosides, saponins and quinones. The total phenolic content was reported much higher as compared to the total flavonoid content mainly in the leaf extract of *L. racemosa*. This mostly contributes to the antioxidant power of the extracts, which is affirmed by the IC$_{50}$ values of the crude extracts in the DPPH assay, which was lower than the ABTS assay. The FRAP assay also exhibited a consistent increase in reducing ability with increase in the concentration which is indicative of the extract’s antioxidant potential. The extracts were also reported to exhibit *in vitro* cytotoxicity and apoptosis inducing ability in Hep G2 cancer cells. And the anticoagulant study conducted provided a first hand report for the plants exhibiting the property. However, further studies must be conducted for secondary metabolite profiling to decipher and clarify the compound(s) responsible for the reported activities of the plant crude extracts.

**INTRODUCTION**

Mangroves are a category of plants, the use of which for medicinal purposes dates back to the year 1230 (Bandaranayake, 1998). The current study aims at documenting a maiden report on the antioxidant, anti-cancerous and anti-coagulant properties of specific parts of the mangrove species: *Acanthus ilicifolius* L. (Acanthaceae) and *Lumnitzera racemosa* Willd. (Combretaceae), based on folklore knowledge, from the Pichavaram mangrove forest situated in the state of Tamil Nadu, India. Prominent survey reports from India suggests the use of roots of *Acanthus ilicifolius* and leaves of *Lumnitzera racemosa* for treatment of snake bites, rheumatism, skin allergies, blood purifier, asthma, diabetics etc. (Bandaranayake, 1998; Pattanaik et al., 2008) And as such, the specific plant parts were collected to primarily investigate their phytochemical & antioxidant properties and their role as anti-cancer agents. They were also tested for blood anti-coagulating property, which in sum would scientifically validate their folklore functionality.

**MATERIALS AND METHODS**

**Sample collection and authentication**

Fresh samples of mangroves (*Acanthus ilicifolius* and *Lumnitzera racemosa*) were collected from Pichavaram mangrove forest located along the southeastern coastline, in the state of Tamil Nadu, India. Guided by the knowledge of folklore use of plants for different medicinal purposes, specifically, the roots of *Acanthus ilicifolius* and the leaves of *Lumnitzera racemosa* were collected from this region during the month of April-May 2013. The plant sample specimens were identified and preserved in the herbarium under the collection number AUCASMBTR01 and AUCASMBTR09.
Sample preparation and extraction

The plant samples collected were then thoroughly washed with distilled water for the removal of contaminants, mud and dirt. After shade drying at room temperature for 3-4 days, the plant samples were pulverized for use in the extraction process.

For the preparation of extracts (root and leaf), 5g of the powdered plant parts (root and leaf) were soaked in 50ml of water in a beaker and was kept on a magnetic stirrer for 24h at room temperature. The extracts were then filtered using Whatman No. 1 filter paper and the filtrates were concentrated using rotary vacuum evaporator. The dried extracts were finally stored at -20 °C until further use (Solomon Charles Ugochukwu et al., 2013).

Chemicals, reagents and cell lines

All the chemicals and reagents used in the phytochemical and antioxidant assays were obtained from the certified suppliers and were of the highest analytical grade. The Hep G2 (human liver hepatocellular carcinoma) cell line was procured from the cell repository of National Center for Cell Science (NCCS), Pune, India. Liquicelin-E and Liquiplastin reagents, for the anticoagulation studies were purchased from Tulip Diagnostics Pvt. Ltd., India.

Phytochemical screening of the extracts

Using standard protocol (Solomon Charles Ugochukwu et al., 2013), the freshly prepared aqueous extracts were subjected to qualitative phytochemical screening for detecting the presence of the following bioactive chemical constituents: phenols, flavonoids, alkaloids, terpenoids, sterols, tannins, proteins and amino acids, carbohydrates, cardiac glycosides, saponins and quinones.

Determination of total phenolic content (TPC)

The standard protocol (Ainsworth and Gillespie, 2007; Barku et al., 2013) for estimating the total phenol content of the extracts using Folin-Ciocalteu reagent was adapted with little modifications. Gallic acid (20-100 µg/ml) was used as a standard. To a volume of 1 ml (100 µg/ml) of aqueous plant extract, 5 ml of Folin-Ciocalteu reagent (diluted to 10 folds) and 4ml of sodium carbonate solution (7.5%) were added. The reaction mix was then allowed to stand in dark at room temperature for 30 min and absorbance of the blue color developed was recorded at a wavelength of 765 nm using a Shimadzu UV-1800 UV-VIS spectrophotometer (Shimadzu, Kyoto, Japan). The TPC of the extracts were determined using the linear regression equation acquired by plotting gallic acid standard curve. The results were calculated as mean ± SD (n=3) and expressed as µg/ml of gallic acid equivalent (GAE) of dry extract.

Determination of total flavonoid content (TFC)

The aluminum chloride colorimetric method (Barku et al., 2013) was used for determining the total flavonoid content of the plant aqueous extracts. Quercetin (20-100 µg/ml) was used as the standard. Each plant extract (1 mg/ml, 0.25 ml) was added to 1.25 ml of distilled water and then 0.075 ml of sodium nitrite solution (5%) was added. The reaction mix was incubated at room temperature for 5 min, following which 0.15 ml of 10% aluminum chloride solution was added and the mix was again allowed to stand for another 6 min at room temperature before adding 0.5 ml of 1M sodium hydroxide solution and finally diluting the reaction mix with 0.275 ml of distilled water. The absorbance was recorded at 510 nm in a Shimadzu UV-1800 UV-VIS spectrophotometer (Shimadzu, Kyoto, Japan). The TFC of the plant extracts were calculated as mean ± SD (n=3) using the linear regression equation obtained by plotting quercetin standard curve and expressed as µg/ml of quercetin equivalent (QE) of dry extract.

In vitro antioxidant assays

DPPH radical scavenging activity

The scavenging activity of the plant aqueous extracts against DPPH (2,2'-diphenyl-2-picrylhydrazyl) radical was determined by the standard method (Banerjee et al., 2008; Zheleva-Dimitrova et al., 2010), with few modifications. An aliquot of 200 µl of different concentrations (20-100 µg/ml) of each of the extracts was mixed with 3.9 ml of freshly prepared DPPH solution (25 mg/L) in methanol. The reaction mixture was mixed and incubated for 30 min at room temperature in dark and its absorbance was recorded at 517 nm. Ascorbic acid was used as the reference standard. The DPPH scavenging capability was calculated using the following equation:

\[
\text{DPPH radical scavenging activity (\%)} = \left( \frac{\text{Abs}_{\text{control}} - \text{Abs}_{\text{sample}}}{\text{Abs}_{\text{control}}} \right) \times 100
\]

where \(\text{Abs}_{\text{control}}\) is the absorbance of the DPPH radical in methanol; \(\text{Abs}_{\text{sample}}\) is the absorbance of the DPPH solution in presence of the aqueous extract or standard. The antioxidant value was expressed as IC\(_{50}\), which is defined as the concentration in µg of the dry extract per ml that inhibits the formation of DPPH radical by 50%. Each value was determined from the slope of the linear regression equation (\(y = mx + c\)), obtained by plotting the ascorbic acid standard curve. All results were calculated as mean ± SD (n=3).

ABTS radical scavenging assay

For ABTS [2,2'-azino-bis (3-ethylbenzothiazoline-6-sulphonic acid)] assay the standard protocol (Floegel et al., 2011; Shalaby and Shanab, 2013) was followed with slight modifications. Initially, two stock solutions of 7 mM ABTS and 2.4 mM potassium persulfate were prepared. The working stock solution was then prepared by mixing the two solutions in equal volume and was allowed to react for 16 h at room temperature in the dark. After incubation, the solution was diluted by mixing 1 ml of ABTS+ with 60 ml methanol to obtain an absorbance of 0.706 ± 0.01 units at 734nm. For each assay the ABTS+ solution was freshly prepared. Then, 1 ml of aqueous extract was added to 1 ml of ABTS+ solution and was allowed to react for 7 min at room temperature. After the incubation, absorbance of the reaction mix
was recorded at 734 nm. Ascorbic acid was used as the experimental standard with which the ABTS radical scavenging capacities of the extracts were compared. The percentage inhibition value was calculated as:

\[
\text{ABTS radical scavenging activity} (\%) = \frac{\text{Abs}_{\text{control}} - \text{Abs}_{\text{sample}}}{\text{Abs}_{\text{control}}} \times 100
\]

where Abs\(_{\text{control}}\) is the absorbance of the ABTS radical in methanol; Abs\(_{\text{sample}}\) is the absorbance of the ABTS in presence of the aqueous extract or standard. The antioxidant value was expressed as IC\(_{50}\), which is defined as the concentration in µg of the dry extract per ml that inhibits the formation of ABTS radical by 50%. Each value was determined from the slope of the linear regression equation \((y = mx + c)\), obtained by plotting the ascorbic acid standard curve. All results were calculated as mean ± SD (n=3).

**Ferric reducing antioxidant power (FRAP) assay**

The ferric reducing power method (Oyaizu, 1986; Chen et al., 2015) was followed for deducing the reducing power of the aqueous extracts. 200 µl of extract was added to 500 µl of phosphate buffer (0.2 M, pH 6.6) and 500 µl of 1 % potassium ferricyanide. The reaction mix was then incubated at 50 °C for 20 min, after which 10 % trichloroacetic acid was added and was subjected to centrifugation at 3000 rpm for 10 min. 700 µl of the supernatant was then taken into a fresh tube and to it 700 µl of distilled water and 140 µl of freshly prepared 1 % ferric chloride solution were added. The absorbance was then recorded at 700 nm. Ascorbic acid was used as the reference standard.

**Cytotoxicity and apoptosis studies on Hep G2 cancer cell line**

For the MTT assay (Nguyen et al., 2015), the Hep G2 (human liver hepatocellular carcinoma) cells were harvested during the logarithmic growth phase and seeded in 96-well plates when the cell density in the culture flask had attained 70-80 % confluency. In each well, a cell density of 3 × 10\(^4\) cells in a volume of 100 µl was maintained and the plate was incubated for 24 h in a CO\(_2\) incubator. The crude extracts were prepared as a stock of 1 mg/ml in the Dulbecco’s Modified Eagle’s Medium (DMEM). The cells were then treated with increasing concentrations of the crude extracts (10, 25, 50, 100 µg/ml) for 48 h in a CO\(_2\) incubator. 10 µl of MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide] solution (5 mg/ml in PBS, pH 7.2) was then added to each well and the plate was further incubated for 4 h in an incubator. After incubation, the medium was carefully decanted to air-dry the formazan crystals in dark. Later, 100 µl of DMSO was added to the wells followed by gentle shaking to solubilize the formazan dye for 15 min at room temperature. Absorbance was then measured using a Synergy H4 microplate reader at 570 nm and the percent cell viability was calculated.

In order to determine the level of apoptosis in Hep G2 cells, DAPI (4’,6-diamidino-2-phenylindole) staining method was performed (Lian et al., 1998). The Hep G2 cells were treated with respective crude extracts at its IC\(_{50}\) concentration and incubated for 24 hours. The cells were then harvested and pelleted by centrifugation at 1100 rpm for 10 min. The medium was aspirated and cells were re-suspended in 1 ml PBS. Again the cells were pelleted and 1 ml of permeabilization buffer (PBS/0.01 M glycine/0.1% Triton X-100) was added followed by incubation for 10 min with gentle rocking. The cells were then washed with 1 ml PBS and fixed with 1 ml of 2% paraformaldehyde in PBS buffer containing 10 µg of DAPI [2µl from stock: 5 mg/ml DAPI solution in N,N-Dimethyl formamide (DMF)] for 10 min in the dark. The stained cells were then washed in 1 ml PBS for 5 min. Finally an aliquot was put onto a clean slide, mount with a coverslip and apoptotic cells were observed using a fluorescent microscope at an excitation wavelength of 350nm. The Image J (Version 2.1) software was used for calculating the pixel value statistics of the stained cells that helps quantify the score of DNA damage in the form of a density histogram which represents normalized values of the treated cells against control.

**Determination of the anticoagulant activity**

The crude extracts for the assays were prepared by dissolving the dried extract in normal saline (0.9% sodium chloride) at different concentrations. Blood samples were collected from 4 healthy volunteers using disposable polypropylene syringe and 3.8% tri-sodium citrate (9 parts of blood to 1 part of trisodium citrate solution) as the anticoagulant. The blood was immediately centrifuged at 4000 x g for 15 min. The freshly prepared plasma was separated, pooled and was stored at 4 °C for use in the assay. The steps followed in this assay had been adapted from (Kathiresan et al., 2006), with slight modifications.

In this study, the action of the crude extracts in the intrinsic and common blood coagulation pathway is evaluated by the APTT assay and that of the extrinsic pathway is determined by the PT assay. In the APTT assay, 0.1 ml of the blood plasma was added to 0.1ml of Liquicelcin-E (a phospholipid preparation derived from the rabbit brain with ellagic acid as an activator) and shaken briefly to mix them. The reaction tube was incubated at 37 °C for 20 min. After incubation, 0.1 ml of pre-warmed calcium chloride solution was forcibly added into the reaction tube. Finally, 0.1 ml of each of the already prepared saline plant extracts of different concentrations (100, 500, 1000 µg/ml) were added to separate reaction tubes and kept at 37 °C. The tubes were shaken gently and tilted back and forth to allow the mixing of the reaction and the time taken to coagulate was recorded in seconds using a stopwatch. As soon as the clot formation begins the stopwatch is stopped. The assay was carried out in triplicates for each sample and the average value was noted and the activity was expressed as clotting time ratio in relation to the control taken, which is heparin.

In the PT assay, 0.1 ml of the blood plasma was taken in a tube and was placed on a water bath at 37 °C for 3-5 min. To this 0.2 ml of pre-warmed (37 °C) Liquiplastin reagent (liquid calcified Thromboplastin reagent, derived from rabbit brain) was added. And then 0.1 ml of the already prepared saline plant extracts of different concentrations (100, 500, 1000 µg/ml) were added to
separate tubes and kept at 37 °C. The time required for the formation of clot was recorded similar to that in the APTT assay.

**Statistical analysis**

The results of all series of experiments were performed in triplicates and were expressed as mean ± SD (standard deviation). Quantitative and graphical data were analyzed using GraphPad Prism 5 software and Microsoft Excel 2011 Package. The data in apoptosis study was analyzed using ImageJ (Version 2.1) package, a public domain, JAVa based image processing software developed at the National Institutes of Health (NIH).

**RESULTS AND DISCUSSION**

**Phytochemical screening of aqueous extracts**

The aqueous extracts of roots of *Acanthus ilicifolius* and leaves of *Lumnitzera racemosa* showed the presence of several classes of bioactive compounds (Table 1). These are the chemical compounds produced in plant tissues as a result of their defense mechanism and are known to have several health benefits (Gavamukulya *et al*., 2014). The presence of the above class of compounds in the crude extracts validates the presence of molecules that are extensively used in the field of medicine (traditional and pharmaceutical industry) and also weighs on the importance of traditional knowledge for use of plants in drug discovery process.

**Table 1:** Results of preliminary phytochemical screening of the aqueous plant extracts.

<table>
<thead>
<tr>
<th>Phytochemicals</th>
<th>Name of Tests</th>
<th><em>Acanthus ilicifolius</em></th>
<th><em>Lumnitzera racemosa</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenols</td>
<td>Ferric chloride test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Gelatin test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>Ferric chloride test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Alkaline reagent test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Alkaloids</td>
<td>Wagner’s test</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Terpenoids</td>
<td>Salkowski’s test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Sterols</td>
<td>Libermann-Burchard test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Tannins</td>
<td>Braymer’s test</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Proteins and Amino acids</td>
<td>Biuret test and Ninhydrin test</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td>Molisch’s test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Cardiac glycosides</td>
<td>Keller-Kiliani test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Saponins</td>
<td>Foam test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Quinones</td>
<td></td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

+: Presence and -: Absence

**Total phenolic content (TPC) & Total flavonoid content (TFC)**

The total phenolic content and total flavonoid content of the extracts were expressed as µg/ml of gallic acid equivalent (GAE) and quercetin equivalent (QE), respectively (Table 2). The TPC in the extracts were determined from the gallic acid calibration curve with a regression equation of $y=0.0046x+0.0891$ and $R^2=0.995$ (Figure 1). And the TFC in the extracts were determined from the quercetin calibration curve with a regression equation of $y=0.0096x+0.0521$ and $R^2=0.993$ (Figure 2). The values clearly indicate and confirm the presence of phenolic compounds in the leaf extract of *Lumnitzera racemosa* as well as in the root extracts of *Acanthus ilicifolius* and literature data abounds in reports that supports the fact that higher phenolic content directly contributes to the overall antioxidant activity of the extract (Luximon-Ramma *et al*., 2002). The antioxidant activity of the phenolic compounds is due to their redox properties, which can play an important role in absorbing and neutralizing free radicals, quenching singlet and triplet oxygen, or decomposing peroxides (Saha *et al*., 2009).

**Table 2:** TPC and TFC of the aqueous plant extracts (n=3). Values are mean ± SD.

<table>
<thead>
<tr>
<th>Crude Extract</th>
<th><em>Acanthus ilicifolius</em></th>
<th><em>Lumnitzera racemosa</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Total phenol content (µg/ml of GAE)</td>
<td>200.98±0.004</td>
<td>476.37±0.007</td>
</tr>
<tr>
<td>Total flavonoid content (µg/ml of QE)</td>
<td>17.28±0.005</td>
<td>24.96±0.004</td>
</tr>
</tbody>
</table>

**In vitro antioxidant assays**

**DPPH radical scavenging activity and ABTS radical scavenging activity**

Figure 3 clearly shows a decrease in the concentration of DPPH due to the scavenging capability of the plant aqueous extracts. The percentage scavenging effect increases with the simultaneous increase in the concentration of extracts. This concludes that the leaf extracts of *Lumnitzera racemosa* is an efficient DPPH-free radical scavenger, exhibiting a lower IC$_{50}$
value of 38.89 µg/ml which is in fair proximity to the IC₅₀ value of the experimental standard, ascorbic acid (21.71 µg/ml), thus proving to have good antioxidant power.

Similarly in the ABTS radical scavenging activity assay too, the scavenging effect increases with the increase in the concentration of the plant extracts (Figure 4) and the IC₅₀ values of the leaf extracts of Lumnitzera racemosa and the root extracts of Acanthus ilicifolius were calculated as 44.38 µg/ml and 60.89 µg/ml respectively. The IC₅₀ value of the experimental standard ascorbic acid was only 19.93 µg/ml.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Ascorbic Acid IC₅₀</th>
<th>Acanthus ilicifolius root extract IC₅₀</th>
<th>Lumnitzera racemosa leaf extract IC₅₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>DPPH radical scavenging assay</td>
<td>21.71</td>
<td>59.85</td>
<td>38.89</td>
</tr>
<tr>
<td>ABTS radical scavenging assay</td>
<td>19.93</td>
<td>60.89</td>
<td>44.38</td>
</tr>
</tbody>
</table>

The present antioxidant study thus reflects that the plant aqueous extracts, particularly the leaf extract of Lumnitzera racemosa was a potent antioxidant. It showed 85% DPPH-free radical scavenging activity at 100 µg/ml. Also the IC₅₀ values of DPPH assay was lower than the ABTS assay, which might be due to the reason that the aqueous extracts had compounds that are better capable of scavenging the DPPH radicals than the ABTS radicals. According to Mukherjee et al., 2011, various factors like (i) stereo-selectivity of the radicals, (ii) solubility of the extract in different testing systems, (iii) polarity of the solvent, (iv) functional groups present in the bioactive compounds, have been reported to affect the capacity of extracts to react and quench different radicals. In sum, the results of the current antioxidant study comprehends and establishes the possibility of the use of these plant extracts for treatment of free radical induced pathological damages (Aiyegoro and Okoh, 2010).

**Ferric reducing antioxidant power (FRAP) assay**

In the FRAP assay, Figure 5 clearly indicates a steady increase in absorbance with the increase in the concentration of plant aqueous extracts which signifies that the ferric reducing ability of the extract increases with increasing concentration. And this reducing capacity is the indication of the extracts having potential antioxidant activity (Babu et al., 2001; Ravikumar and Gnanadesigan, 2011; Firdaus et al., 2013). However, in comparison to the experimental standard ascorbic acid, the plant extracts exhibited moderately lower ferric reducing capability.

**Cytotoxicity and apoptosis studies on Hep G2 cancer cell line**

The present study in addition to previously reported studies (Bandaranayake, 2002; Huo et al., 2005; Raut and Khan, 2012; Singh and Aeri, 2013; Huang et al., 2014; Nguyen et al., 2015) suggest that mangroves are a rich source of secondary metabolites probably because they grow and survive under very hostile environmental conditions (high salinity, low nutrition, low oxygen conditions of waterlogged mud and high solar radiation during low tide) (Kathiresan and Bingham, 2001). These naturally occurring bioactive constituents (eg- phenolic compounds) are majorly known to play an important role in the mechanism of anticancer (Babu et al., 2002; Bunyapraphatsara et al., 2003; Van Kiem et al., 2008; Ravikumar and Gnanadesigan, 2011; Firdaus et al., 2013). This is a maiden report of the mangrove plant samples from Pichavaram mangrove forest, Tamil Nadu, India that states the cytotoxicity of the aqueous root extract of Acanthus ilicifolius and leaf extract of Lumnitzera racemosa. In this study the cytotoxic effect of the crude plant extracts were tested against Hep G2 cancer cell line using MTT assay. The IC₅₀ values for root extract of Acanthus ilicifolius and leaf extract of Lumnitzera racemosa were reported as 39.76 µg/ml and 26.05 µg/ml respectively. Also, the percent cell viability (Figure 6) of the leaf extract of Lumnitzera racemosa exhibited more potent cytotoxicity on Hep G2 cell lines at different concentrations. The higher
cytotoxicity of the crude extracts may be due to their higher antioxidant potentials or due to the synergistic effect of the multiple bioactive components present in the extract. The antioxidants are capable of scavenging the free radicals that are known to damage healthy cells and is a prime reason for the occurrence of cancer and other free radical induced conditions.

Determination of anticoagulant activity

Heparin is majorly used for limiting blood coagulation in the field of medicine. But owing to some of its side effects like hemorrhage there has been an urge to look for an alternate to it and hence researchers have been trying to find anticoagulant from natural sources (Edemeka and Ogwu, 2000; Félix-Silva et al., 2014; Khouya et al., 2015). The anticoagulant study involving the APTT and PT assays using normal citrated plasma is an attempt to check the anticoagulation effect of these mangrove plant extracts. Studies however suggest that the anticoagulant property of mangrove plant extract varies over a great range depending on the species and also on the part of the plant (Kathiresan et al., 2006). This assay was performed just to check and report if the plants exhibited anticoagulant property. This is the maiden report for the plants used in this study.

The results show that both the assays vary slightly in the anticoagulant activity and the prolongation of the APTT is slightly higher than the PT in case of both the plant extracts. Table 4 indicates that aqueous extracts of A.ilicifolius and L.racemosa had only slightly prolonged the coagulation times compared with the control sample treated with PBS, suggesting that extracts inhibited the common pathways but isn’t too effective compared to Heparin. A previous research however showed that certain other mangrove species like Avicennia marina, Aegiceras corniculatum exhibited a higher anticoagulant property (Kathiresan et al., 2006), which further proves that the property is totally independent of the plant genera and varies greatly with regional location.

Table 4: Clotting time ratio in mangrove extracts assayed by using APTT and PT methods. The clotting ratio is the time taken for blood clotting in the mangrove extracts treated to the control. The clotting time for control was 80 s and 32 s for APTT and PT methods respectively.

<table>
<thead>
<tr>
<th>Mangrove Species</th>
<th>APTT Assay (µg/ml)</th>
<th>PT Assay (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>100</td>
<td>500</td>
</tr>
<tr>
<td>Acanthus ilicifolius</td>
<td>1.3</td>
<td>1.35</td>
</tr>
<tr>
<td>Lumnitzera racemosa</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td>Heparin</td>
<td>315</td>
<td>-</td>
</tr>
</tbody>
</table>

CONCLUSION

We have investigated the aqueous extracts of the mangroves for their phytochemical constituents, in vitro antioxidant, anti-cancer and anticoagulant activities to scientifically validate their folklore use in treatment of diseases. This is a first hand report that provides sufficient evidence for carrying out further research on the selected plants to decipher the exact mechanism involved in anticaner and anticoagulant activity. Thereby suggesting in vitro, in vivo and secondary metabolite profiling studies to unravel and identify the bioactive compound(s) responsible, and ultimately provide alternative treatment strategies.
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