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1. Introduction

“Diarrhea is the passage of 3 or more loose or liquid stool per day, or more frequently than is normal for the individual. It is usually a symptom of gastrointestinal infection, which can be caused by a variety of bacterial, viral and parasitic organisms, infection is spread through contaminated food or drinking-water, or from person to person as a result of poor hygiene” (WHO). Diarrheal diseases affect all races, sexes, ages and geographic areas, has high impact on mortality and morbidity worldwide, an estimated 2-4 billion episodes of infectious diarrhea occurred each year and are especially prevalent in infants (Hodges and Gill 2010; Farthing 2002). In 2005, 1.8 million people died worldwide from diarrheal diseases (WHO, 2007). In México, in the past 6 years, the gastrointestinal infection has been a serious health problem and was the second cause of morbidity among all age groups (SS, 2008).

2. Pathophysiology classification of diarrhea

- Osmotic; is caused by poorly absorbable solutes (eg. sorbitol, magnesium salts) remaining in the gastrointestinal lumen retain water and electrolytes resulting in reduced water reabsorption
- Altered Motility; caused slowing of the motor function of the small intestine as with narcotic use, scleroderma, diabetic autonomic neuropathy and amyloidosis
- Exudative; the intestinal epithelium’s barrier function is compromised by loss of epithelial cells or disruption of tight junctions (eg. E. coli, Salmonella, Shigella, Yersinia, Campilobacter, Mycobacterium tuberculosis, Clostridium difficile y Entamoeba histolytica), inflammatory disease process as in ulcerative colitis and Crohn’s disease
- Secretory; is caused by an increase in water and electrolytes (Chloride or bicarbonate) movements to the intestinal lumen, the final effect is the increase of secretion and decrease of absorption of net sodium and water (Navaneethan and Giannella, 2010).

2.1 Secretory diarrhea

Secretory diarrhea occurs when the balance between absorption and secretion in the small intestine is disturbed by excessive secretion caused by bacterial enterotoxins, is a net movement from mucous intestinal to lumen, the volume exceed 10 mL/Kg/day, and the osmolarity is similar with plasma. It is the leading cause of death in infants in developing
countries and currently accounts for an estimated of three million deaths each year among under 5 years old children (Casburne-Jones and Farthing, 2004; Filbin 2004). Most causes of secretory diarrhea alter the second messenger system through alteration in cAMP, cGMP or intracellular calcium regulated ion transport pathways, alterations in these mediators cause CFTR-mediated Cl⁻ secretion an inhibition of small intestinal-coupled Na⁺-Cl⁻ transport (Navaneethan and Giannella, 2010)

2.1.1 Secretory diarrhea noninfectious
Some of these include
• Tumors (pancreatic islet, which secrete vasoactive intestinal peptide (VIP), carcinoid which elaborate serotonin, bradykinin, substance P and prostaglandins, medullary carcinoma of thyroid-secreting calcitonin)
• Neurotransmitters are also potent secretory stimuli, such as histamine in systemic mastocytosis and inflammatory cytokines
• Malabsorbed bile salts and fatty acids (hydroxyl fatty acids also stimulate colonic secretion)
• The congenital absence or alterations in the numerous transporters that maintain the constant flux of the ions and water
• Rare congenital syndromes: congenital chloridorrhea, there is a defect in brush border Cl⁻/HCO₃⁻ exchange in the ileum and the colon and hence impaired absorption of chloride, congenital sodium diarrhea results from a congenital defect in Na⁺-bile acid absorption in the colon (Navaneethan and Giannella, 2010; Filbin 2004)

2.1.2 Secretory diarrhea caused by pathogens
Microbial causes include rotavirus, norovirus, Cryptosporidium, its affects the absorptive villi inhibiting sodium absorption. Enterotoxigenic E. coli (ETEC), V. cholera elaborate enterotoxins that stimulate intestinal chloride secretion along with impaired sodium absorption, Giardia lambia adhere to the mucosa disrupting the absorptive/secretory process of enterocyte producing active secretion (Navaneethan and Giannella, 2010)

2.1.2.1 Enterotoxigenic bacteria
• Vibrio cholerae
• Enterotoxigenic Escherichia coli
• Clostridium perfringes
• C. botulinum
• Campylobacter jejuni
• Klepsiella pneumoniae
• Aeromonas hydrophila
• Yersinia enterocolitic

2.1.2.2 Enteroinvasive bacteria
• Enteroinvasive Escherichia coli
• Salmonella typhi
• S. enteritidis
• Shigella spp
2.1.2.3 Viruses
- Group A rotaviruses, G1 and G serotypes
- Norovirus (old term of Norwalk virus)
- Parvoviruses (Havai, Colorado, Ditchillign
- Enteric adenoviruses 40 and 41
- Coronaviruses
- Calciviruses
- Astroviruses

2.1.2.4 Parasites
- Gardia lamblia
- Entamoeba histolityca
- Cryptosporidium parvum
- Isospore belli
- Sarcosystis sp
- Cyclospore cayetanensis
- Blastocystis hominis
- Microsporidie

2.2 Diarrhea caused by enterotoxins
A number of several bacteria cause diarrhea by the production of potent enterotoxins, such as enterotoxigenic Escherichia coli, Salmonella typhi, S. typhimurium, clostridium difficile, C. freundii, Aeromonas hydrophila, Yersinia enterocolic, Camphyllobaster jejuni and Vibrio cholera. Enterotoxins have their effect on the enterocyte functions by stimulating the secretion of transepitelial electrolytes, increasing the osmotic flux of water and ions to the intestinal lumen, specifically, heat-labile (LT) and heat stable (ST) enterotoxins from E. coli, V. cholera and C. jejuni increase net fluid secretion by affecting the enzymes adenylate cyclase or guanilate cyclase by activation of the cAMP (cyclic 3',5'-adenosine monophosphate) in the mucosal epithelium which induces an increase of intestinal secretion and causes diarrhea. (Casburn-Jones and Farthing 2004, Amstrong and Cohen, 1999).

2.2.1 Vibrio cholerae enterotoxin
Vibrio cholerae enterotoxin is an oligomeric protein which is composed by two subunits, A subunit of 27.2 KDa composed as A1 and A2 subunits, and B subunit composed by five subunits B of 11.6 KDa each one, AB₅ complex Fig 1. (Sixma, 1991)

2.2.2 Vibrio cholerae enterotoxin mechanism
Mechanisms proposed to secretory diarrhea caused by V. cholera enterotoxin involves the union of subunits B to the oligosaccharide portion of the receptor GMI, present in the apical surface of enterocytes, this union lend the entrance of A subunit of toxin to the enterocyte for acidic endosomes, which pick up the golgi apparatus and endoplasmic reticulum,
inside of enterocyte disulphure bond between A1-A2 is dissolved by protein disulfide isomerase, it causes release of A1 which is capable of binding NAD and catalyzing the NADP-ribosylation of $G_{\alpha}$, a GTP-binding regulatory protein associated with adenylate cyclase. A1 subunit stimulates increasing of 1000 times production of cAMP second messenger, cAMP active protein cinase A, which phosphorile and activate transmembranal chloride channels of the enterocytes located on intestinal crypts, it causes massive secretion of water and electrolytes to intestinal lumen, in villus cell there is a decrease of absorption of $Na^+$ and $Cl^-$ ions (Kopic 2010). Fig 2. shows the proposed mechanism to the action of *Vibrio Cholerae* enterotoxin. At 5 to 10 minutes to the exposure of V. cholera toxin cause intestinal hypersecretion of water and electrolytes for several hours (Thiagarajah, 2005; Spangler, 1992). The symptoms are manifested as severe cramp and the copious “rice water” diarrhea characteristic of the disease.

### 2.3 Dehydration

During diarrhea there is an increased loss of water and electrolytes (sodium, chloride, potassium and bicarbonate) in the liquid stool; dehydration occurs when these losses are not replaced adequately and a deficit of water and electrolytes develops. The degree of dehydration is graded according to signs and symptoms that reflect the amount of fluid lost:

- **In the early stages of dehydration,** there are no signs or symptoms
- **As dehydration increases,** signs and symptoms develop. Initially these include: thirst, restless or irritable behavior, decreased skin turgor, sunken eyes, and sunken fontanel (in infants).
- **In severe dehydration,** these effects become more pronounced and the patient may develop evidence of hypovolemic shock, including: diminished consciousness, lack of urine output, cool moist extremities, a rapid feeble pulse, low or undetectable blood pressure, and peripheral cyanosis.
- **Death follows soon if rehydration is not started quickly** (WHO, 2005).

### 2.4 Secretory diarrhea treatment

#### 2.4.1 Oral rehydration

To control diarrhea disease, a sufficient hydration of the patient should be procure and provide the necessary ions to maintain electrolyte balance, the treatment of choice is oral
Management of Secretory Diarrhea

Fig. 2. Cholera toxin mechanism proposed by Velázquez et al., A1,2 B5 (subunits), GM1 (ganglioside receptor), Gsα (G protein), cAMP (cyclic AMP), CFTR (cystic fibrosis transmembrane conductance regulator).

Rehydration solution (ORS), it has reduced the levels of mortality in children and elderly by dehydration, but not morbidity (Turvill et al., 2000), the treatment is based by active absorption of glucose by smooth intestine, during the intestinal infection lend to the co-transport of Na+ ions and water absorption. WHO and UNICEF guidelines recommend their use, is important to notice that ORS, is useful to treat dehydration caused by diarrhea, but it not decrease the amount and duration at the same. Depends of severity of diarrhea, in some cases ORS is not enough and antibiotic, spasmytic, and antiprotozoal drugs should be used. WHO recommended use of secure and effective drugs to the pediatrics (Marion et al., 2010).
2.4.2 Drugs used to treat secretory diarrhea

To treat the secretory diarrhea there are some drugs which reduce the intestinal movement such as codeine (1), loperamide (2), diphenoxylate (3), lidamidine (4), bismuth subsalicylate (5), racecadotril (6) and clonidine (7) Table 1.; which are capable to stimulate absorption direct and reduce secretion of water and electrolytes in gastrointestinal tract, to decrease propulsion, contact time of intestinal content with mucosal surface increase, it favors the absorption. They act not premised release of prostaglandins too (Marion et al., 2010; Martindale 2009).

<table>
<thead>
<tr>
<th>Structure</th>
<th>Effect</th>
</tr>
</thead>
</table>
| ![Codeine](https://example.com/codeine.png)  
**Codeine (1)**  
Has a high antidiarrheal action but, produces secondary effects such as nauseas, dizziness and acts against central nervous system, it can be used carefully in children, continuous use can induce physical dependence and addiction |
| ![Loperamide](https://example.com/loperamide.png)  
**Loperamide (2)**  
Decrease intestinal motility and present antisecretory effect by activation of calmodulin, increase the water and electrolytes absorption to the intestinal lumen. It should not be administrated to children under six years old, patients with constipation, atony or intestinal obstruction, should avoid its use on bacterial infective severe and in acute dysentery. Frequent adverse reactions induced are hypersensitivity reactions (cutaneous eruption), gastrointestinal disorders (constipation, colic, abdominal distention, nauseas and vomit), fiver and dry mouth, is a non-prescription drug for children because can cause CNS depression |
| ![Diphenoxylate](https://example.com/diphenoxylate.png)  
**Diphenoxylate (3)**  
Inhibits intestinal propulsion and fecal excretion velocity, cause decrease of intestinal transit, to therapeutic doses it induces adverse reactions in central nervous system (confusion, sedation, depletion, cephalae), allergic reactions (anaphylaxis, prurite) on gastrointestinal apparatus (toxic megacolon, paralytic ileum, vomit, nauseas and abdominal pain) It can cause euphoria and has analgesic effect. Difenoxilate is contraindicated in children younger than 2 years old |
| ![Lidamidine](https://example.com/lidamidine.png)  
**Lidamidine (4)**  
Improve the absorption of water and electrolytes in intestinal velocities and reverse their secretion to level on intestinal crypts |
### Table 1. Drugs used to treat secretory diarrhea

<table>
<thead>
<tr>
<th>Structure</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Bismuth subsalicylate" /></td>
<td>Showed antisecretor effect for the inhibition of prostaglandins, reduce depositions number and reduce abdominal pain, causes adverse reactions (dizziness, cephalgia, constipation, dark stools, ataxia, tremor, encephalopathy, confusion, delirium and convulsions).</td>
</tr>
<tr>
<td><img src="image" alt="Racecadotril" /></td>
<td>Decrease intestinal hipersecretion of water and electrolytes to intestinal lumen, inhibits release of encephalinse endogenus witch act on opiateous receptors γ decreasing cAMP level (decrease water and electrolytes secretion), cause some adverse reactions such as hypokalemia, bronchospasm, fever, vomit and otitis.</td>
</tr>
<tr>
<td><img src="image" alt="Clonidine" /></td>
<td>Stimulates sodium and chloride absorption and inhibits chloride secretion by interaction with its receptor on enterocyte, causes an alteration of gut motility with effect on intestinal transport, it causes hypotension</td>
</tr>
</tbody>
</table>

On the other hand, there are some compounds that showed inhibitory properties on the intestinal secretion Fig 3. such as berberine (8), chlorpromazine (9), nicotinic acid (10), indomethacin (11), somatostatin (12) and ethacrinic acid (13) but they were not developed as antidiarrheal drugs (Fedorack and Field, 1987). Thus, the research for new antisecretory agents that should be effective and safe to treat diarrhea is still a necessary goal.
2.4.3 Potential target areas to design therapeutic agents on Vibrio cholerae toxin

During the last two decades there has been a continuous research of drugs that inhibit the secretory process in the enterocyte to help to the control of diarrhea, but only a few candidates have emerged, and none has found a place in the routine management of secretory diarrhea. Particularly to cholera toxin its mechanism of action reveal several potential target areas to design therapeutic agents such as:

a. The inhibition of adenylate cyclase
b. The blockage of the active site of the enzyme located in the A subunit
c. The disruption of the assembly of the holotoxin by interrupting the A₂-B interaction
d. The interception of the receptor binding to the bottom of the B pentamer (Guangtao Z., 2009).

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e. Inhibitors of enkephalinase and of the cystic fibrosis transmembrane conductance regulator (Thiagarajah, 2005).

f. Inhibition of transport proteins involved in cAMP activated chloride secretion

![Chemical structures](http://www.bmsc.washington.edu/WimHol/figures/figs2/WimFigs2.html)

**Fig. 4.** Synergism of structure-based drug design with combinatorial chemistry for the design of receptor antagonist of cholera toxin

http://www.bmsc.washington.edu/WimHol/figures/figs2/WimFigs2.html
2.5 Medicinal plants as a source of antidiarrheal compounds

Diarrheal diseases are a health problem because affect a large number of the population mainly children and elderly. In Mexico the use of medicinal plants to treat gastrointestinal disorders including diarrhea occupied the first place, there are few pharmacological and chemical studies which support their use. Approximately 80% of the world’s population uses medicinal plants to treat immediately health problems, is clear the importance of multidisciplinary research of our natural sources. The study of medicinal plants with the propose to provide pharmacological evidence that may explain its therapeutic use. There are some in vitro models such as isolated ileum of guinea pig, isolated jejune of rabbit, ileum and duodenum isolated of rat or rabbit and in vivo reduction of intestinal motility using charcoal meal, Castor oil, PGE$_2$, MgSO$_4$ induced diarrhea, and Enteropooling models. Antispasmodic activity has been demonstrate for some flavonoids such as quercetin, quercitrin, genistein, sakuranetin, rutin and bisabolol; terpenoids such as himachalol, coleonol, β-damascenone, e-fitol, capsidiol, β-eudemol, hinesol, huatriwaico acid, camaldulin and tymol; essential oils such as, 1,8-cineol, eugeol, timol, carvacrol, estragolnetol, α y β pinenes, nonanal, and linalool; alkaloids such as himbacine, protopine, coptisine, cantleyine, mitraginine, verteine, retuline, cavidine and metuenine (Astudillo et al., 2009).

There are a great number of natural remedies for diarrhea control, historically Papaver somniferum preparations are efficient and powerful against diarrhea, as the derivative codeine, alkaloids are ones of major substances explored form natural products and they give to pharmaceutical industry a big number of patents, another class of compound explored therapeutically are flavonoids and has been used as complement in treatment of cancer, heart diseases, venous insufficiency, venous ulcers, hemorrhoids and diarrhea. (Martindale 2009)

2.5.1 Antisecretory compounds isolated from medicinal plants

Some studies have been performed in order to find antisecretory compounds from several plants used in traditional medicine to treat several kinds of diarrheas. In this sense the extracts from Croton urucurana, C. lechleri, Berberis aristata and Guazuma ulmifolia were studied against intestinal secretion caused by V. cholera toxin, in the cases of C. lechleri, B. aristata and G. ulmifolia the isolated compounds were oligomeric proantocyanidins and berberine, respectively. From C. urucurana saponins, steroids, alkaloids, antocianidins and catechins have been isolated. Prontocianidins and catechins probably can be associated with their antisecretory activity. Steviosid (29) and dihydroisosteviosid (30) can inhibit cAMP-activated chloride secretion in human’s intestine cells by targeting CFTR (Pariwat 2008). Penta-μ-digalloyl-glucose (PDG) (31) isolated of Chinese gallnuts showed efficacy in reducing enterotoxin-induced intestinal fluid secretion in mice (Wongasmitkul et al., 2010)
Crofelemer (32) is a proanthocyanidin oligomer obtained from Croton lechleri (dragon’s blood), the sap has been used to treat diarrheas including dysentery and cholera, pharmacological studies have shown that it reduces fluid secretion in cell culture and mouse models (Gabriel et al., 1999), it has been reported that the antisecretory mechanism of action of crofelemer involves dual inhibition of The cystic fibrosis transmembrane regulator conductance (CFTR), a cAMP stimulated Cl- channel, and calcium-activated Cl- channels (CaCC) at the luminal membrane of enterocytes (preliminary studies showed that crofelemer (32) may reduce watery stool output in patients with infectious diarrhea such as cholera. But it needs further Phase 3 clinical trials are still necessary (Crutchley et al., 2010).

Penta-α-digalloyl-glucose (PDG) (31)

We continue with the research of compounds with antisecretory activity useful to treat diarrhea. Medicinal plants used in Mexican traditional medicine to treat gastrointestinal disorders could be a source of compounds with therapeutic utility. In México, the use of medicinal plants to treat gastrointestinal disorders such as diarrhea and dysentery is widespread (Aguilar et al., 1994). However most of these plants have not been investigated from a pharmacological point of view to demonstrate their antisecretory properties, which could lead to support their use as antidiarrheal and anti-dysenteric drug in traditional medicine. We screened aqueous and methanol extracts from 26 Mexican medicinal plants to assess their antisecretory activity using the cholera toxin-induced intestinal secretion in rat jejunal loops model. None of this species or their isolated compounds has been previously evaluated as antisecretory agents (Velázquez et al., 2006).
2.5.2 Material and methods

2.5.2.1 Plant materials

The plants used in that study were collected from different regions in Mexico: Mexico City, States of Hidalgo, Mexico, Sinaloa, Guanajuato and Yucatan, all of them were selected according to their use in Mexican traditional medicine to treat gastrointestinal disorders. Voucher herbarium specimens were deposited in Herbarium IMSSM of Instituto Mexicano del Seguro Social and were authenticated by MS Abigail Aguilar.

2.5.2.2 Preparation of crude extracts

The air-dried plant material (20g) was extracted by maceration with 300 mL of MeOH for 1 week. Then the macerate was filtered and concentrated under reduced pressure at 40°C. For aqueous extracts, 20 g of air-dried plant material were extracted by decoction with 100 mL of distilled water for 30 min, the solution was filtered and lyophilized.

2.5.2.3 Cholera toxin

Lyophilized powder (1mg) of Cholera toxin (SIGMA) containing approximately 220,000 units/mg of protein was suspended in 1 mL of sterile water. Aliquots of the toxin solution were dissolved in a 1x PBS (NaCl 8g, KCl 0.2 g, Na2HPO4.7H2O 0.115 g, KH2PO4 0.2 g/L) solution with 1% bovine serum albumin (SIGMA) to obtain a concentration of 3 μg/mL.

2.5.2.4 Antisecretory assay

The antisecretory activity of the extracts was tested using a method described by Torres et al., in 1993. Briefly, male Sprague-Dawley rats (200-250 g) were obtained from the animal house of the IMSS. The experimental protocols were in accordance with the official Mexican norm NOM 062-ZOO-1999 entitled technical specifications for the production, care and use of laboratory animals (SAGARPA 2001). The antisecretory effect of the extract was studied on intestinal secretion indirectly by measuring the fluid accumulation in the intestine following cholera toxin administration to rats. Two jejuna loops were prepared in the rats and inoculated with 3 μg/mL of cholera toxin dissolved in 1x PBS with 1 % bovine albumin. Rats (n=4 per group by duplicated) were treated orally with each extract (300 mg/Kg in 1 mL of a 2 % DMSO solution in water). Loperamide (10 mg/Kg) was used as antidiarrheal drug. After 4 h, the animals were sacrificed using ethyl ether. The antisecretory activity of the extracts was measured as the fluid accumulation in the loops and expressed in percentage of inhibition. Values are expressed as mean ± SEM. Statistical significance was determinate using Mann-Whitney U-test. Values with p<0.05 were considered significant.

2.5.2.5 Results

We tested 56 aqueous and methanol crude extracts obtained from 26 medicinal plants used in Mexican traditional medicine for the treatment of gastrointestinal disorders. The antisecretory activity was tested using the cholera toxin-induced intestinal secretion in rat jejunal loops model. Only the principal antisecretory activity of the extracts tested is shown in Table 2, the full list is showed in Velázquez et al., 2006. In traditional medicine since infusions or decoctions are usually taken three times per day when diarrhea occurs, our results can be related with their traditional use because the used dose is approximately one cup of plant tea which is recommended by Mexican people to treat gastrointestinal disorders (Aguilar et al., 1994). We found that both extracts from Chiranthodendron pentadactylon, Hippocratea excelsa and Ocimum basilicum were the most
active with inhibition values ranging from 68.0 to 87.6% at 300 mg/kg. Methanol extract of *Geranium mexicanum* (aerial parts) and the aqueous extract of *Bocconia frutescens* were active too with inhibition values of 93.4 and 86.0%, respectively. On the other hand, the methanol extract of *Chenopodium ambrosioides* green variety (aerial parts), *Lygodium venustum*, *Punica granatum* and *Ruta chalepensis*, the aqueous extracts of *Alloysia triphylla*, *Chenopodium ambrosioides* green variety (aerial parts), *Dorstenia contrajerva* and *Schinus molle* shown inhibitory activity with values ranging from 43.4 to 79.5%. The 87% of the extracts tested showed inhibitory activity of the intestinal secretion; only seven extracts did not show any antisecretory activity. In general, among the researched extracts, the methanol extracts exhibited the highest antisecretory activity.

<table>
<thead>
<tr>
<th>Family</th>
<th>Plant specie</th>
<th>Part used</th>
<th>Voucher number</th>
<th>Extract</th>
<th>% Inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verbenaceae</td>
<td><em>Alloysia triphylla</em> (L’Her)</td>
<td>AP</td>
<td>126110</td>
<td>Methanol</td>
<td>7.8±4.7</td>
</tr>
<tr>
<td></td>
<td><em>Alloysia triphylla</em> (L’Her)</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>80.4±22.8</td>
</tr>
<tr>
<td>Papaveraceae</td>
<td><em>Bocconia frutescens</em> L.</td>
<td>AP</td>
<td>12618</td>
<td>Methanol</td>
<td>24.1±15.4</td>
</tr>
<tr>
<td></td>
<td><em>Bocconia frutescens</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>86.0±9.8</td>
</tr>
<tr>
<td>Chenopodiaceae</td>
<td><em>Chenopodium ambrosioides</em> L., green variety</td>
<td>AP</td>
<td>14402</td>
<td>Methanol</td>
<td>43.4±6.5</td>
</tr>
<tr>
<td></td>
<td><em>Chenopodium ambrosioides</em> L., green variety</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>48.7±11.6</td>
</tr>
<tr>
<td>Sterculiaceae</td>
<td><em>Chiranthodendron pentadactylon</em> Larreat</td>
<td>F</td>
<td>14104</td>
<td>Methanol</td>
<td>87.6±15.3</td>
</tr>
<tr>
<td></td>
<td><em>Chiranthodendron pentadactylon</em> Larreat</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>84.8±17.4</td>
</tr>
<tr>
<td>Moraceae</td>
<td><em>Dorstenia contrajerva</em> L.</td>
<td>AP</td>
<td>14406</td>
<td>Methanol</td>
<td>24.4±16.4</td>
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<tr>
<td></td>
<td><em>Dorstenia contrajerva</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>44.8±5.9</td>
</tr>
<tr>
<td>Geraniaceae</td>
<td><em>Geranium mexicanum</em> H. B. &amp; K.</td>
<td>AP</td>
<td>14405</td>
<td>Methanol</td>
<td>93.4±6.7</td>
</tr>
<tr>
<td></td>
<td><em>Geranium mexicanum</em> H. B. &amp; K.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>42.1±15.2</td>
</tr>
<tr>
<td>Hippocrateae</td>
<td><em>Hippocrates exels</em> H. B. &amp; K.</td>
<td>R</td>
<td>14394</td>
<td>Methanol</td>
<td>80.3±21.3</td>
</tr>
<tr>
<td></td>
<td><em>Hippocrates exels</em> H. B. &amp; K.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>75.0±24.9</td>
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<td>Schizaeaceae</td>
<td><em>Lygodium venustum</em> Sw.</td>
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<td>1270</td>
<td>Methanol</td>
<td>51.6±15.6</td>
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<td><em>Lygodium venustum</em> Sw.</td>
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<td></td>
<td>Aqueous</td>
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<tr>
<td>Labiatae</td>
<td><em>Ocimum basilicum</em> L.</td>
<td>AP</td>
<td>14393</td>
<td>Methanol</td>
<td>68.7±9.7</td>
</tr>
<tr>
<td></td>
<td><em>Ocimum basilicum</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>68.0±20.8</td>
</tr>
<tr>
<td>Punicaceae</td>
<td><em>Punica granatum</em> L.</td>
<td>EF</td>
<td>14403</td>
<td>Methanol</td>
<td>55.9±3.6</td>
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<tr>
<td></td>
<td><em>Punica granatum</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>19.1±6.9</td>
</tr>
<tr>
<td>Rutaceae</td>
<td><em>Ruta chalepensis</em> L.</td>
<td>AP</td>
<td>14400</td>
<td>Methanol</td>
<td>73.7±0.01</td>
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<tr>
<td></td>
<td><em>Ruta chalepensis</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>23.6±9.27</td>
</tr>
<tr>
<td>Anacardiaceae</td>
<td><em>Schinus molle</em> L.</td>
<td>AP</td>
<td>14408</td>
<td>Methanol</td>
<td>79.5±17.7</td>
</tr>
<tr>
<td></td>
<td><em>Schinus molle</em> L.</td>
<td></td>
<td></td>
<td>Aqueous</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 2. Antisecretory activity of methanol and aqueous extracts of Mexican medicinal plants on intestinal secretion response to cholera toxin, AP: aerial parts, EF: fruit exocarpus, F: flowers, R: roots.

2.6 Antisecretory study of *Chiranthodendron pentadactylon*

We selected *Chiranthodendron pentadactylon* Larreat (Sterculiaceae) to perform bio-guided assay fractionation. *C. pentadactylon* know in Mexico as “flor de manita” has been used in Mexican traditional medicine since Aztecs ancient times to treat heart illness, epilepsy,
diarrhea and dysentery (Linares et al., 1988, Argueta et al., 1994). This study lends to the isolation of some compounds with in vivo antisecretory activity (Velázquez et al., 2009).

2.6.1 Isolation of active compounds
The flowers of *C. pentadactylon* were ground and extracted by maceration at room temperature with methanol, the extract was suspended in 10 % MeOH-water and successively partitioned with CH₂Cl₂ and EtOAc, the aqueous residual layer was lyophilized. The fractions were tested for antisecretory activity at doses of 50 mg/Kg. The most active fraction was AcOEt with 88.2 % of inhibition. In order to isolate the active compounds, it was subjected to column chromatography on Sephadex (Pharmacie) using CHCl₃ in EtOH, MeOH and Water to give eight secondary fractions, further chromatography led to the isolation of tiliroside (33), astragalin (34), isoquercitrin (35), (+)-catechin (36), and (-) epicatechin (37). All the isolated compounds were identified by comparison of spectroscopic data (¹H and ¹³C NMR, UV, IR, [α], and TLC and HPLC with authentic samples available in our laboratory (Kuroyanagui et al., 1978; Lee et al., 1992; Lui et al., 1999; Calzada et al., 2007).

2.6.2 Antisecretory activity of isolated compounds
Antisecretory activity of the isolated compounds from the AcOEt fraction was tested on cholera toxin-induced intestinal secretion in rat jejunal loops model (table 3). Among the isolated compounds (-) epicatechin (37) showed the best antisecretory activity on the intestinal secretion with an ID₅₀ of 8.3 µM/mL, its antisecretory activity was like of loperamide (2) (ID₅₀ 6.1 µM/mL), isoquercitrin (35) and (+)-catechin (36) showed moderate and weak antisecretory activity, respectively. Tiliroside (33) and astragalin (34) were inactive at doses tested table 3. Flavonoids such as flavan-3-ols and flavonol glycosides have been considered as the active principles of many antidiarrheal plants. Isoquercitrin isolated from *Psidium guajava* showed spasmolytic effect on guinea pig ileum (Morales et al., 1994). Tiliroside (33) and (-)-epicatechin (37) obtained from *Helianthemum glomeratum* and *Rubus corifolius*, respectively, showed antiprotozoal activity against *Entamoeba histolytica* and *Giardia lamblia* (Alanis et al., 2003; Barbosa et al., 2007). Data obtained in this investigation suggest that (-)-epicatechine (37), isoquercitrin (35) and tiliroside (33) may play an important role in antidiarrheal of *C. pentadactylon* in Mexican traditional medicine. Also, our results are in agreement and could explain the result previously obtained by Hör et al., 1995, with antise retory oligomeric proantocianidins from *Guazuma ulmifolia* which monomeric unit are (+)-catechin (36) and (-)-epicatechin (37). The antiprotozoal activity together with the antisecretory activity is evidences that support the use of these plants to treat diarrhea in Mexican traditional medicine.

Further studies are carried on in order to determinate the action mechanism of active compounds against intestinal secretion caused by *Vibrio cholerae* toxin (non published results). Additionally we are studying some medicinal plants used to treat gastrointestinal disorders in Mexican traditional medicine from Hidalgo, using intestinal propulsion (charcoal meal), castor oil induced diarrhea and castor oil induced intestinal fluid accumulation models in vivo.
tiliroside 33                                            astragalin 34

Isoquercitrin (35)     (+)-catechin (36)

(-) epicatechin (37)    loperamide (2)
<table>
<thead>
<tr>
<th>Compound</th>
<th>Doses (mg/Kg)</th>
<th>% of Inhibition</th>
<th>ID₅₀ (μM/Kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MeOH extract</td>
<td>300</td>
<td>87.1 ±14.5</td>
<td>-</td>
</tr>
<tr>
<td>EtOAc fraction</td>
<td>50</td>
<td>88.2±9.5</td>
<td>-</td>
</tr>
<tr>
<td>Tiliroside (33)</td>
<td>10</td>
<td>-</td>
<td>Inactive</td>
</tr>
<tr>
<td>Astragalins (34)</td>
<td>10</td>
<td>-</td>
<td>Inactive</td>
</tr>
<tr>
<td>Isoquercitrin (35)</td>
<td>10</td>
<td>-</td>
<td>19.2</td>
</tr>
<tr>
<td>(+)-catechin (36)</td>
<td>10</td>
<td>-</td>
<td>51.7</td>
</tr>
<tr>
<td>(-)-epicatechin (37)</td>
<td>10</td>
<td>-</td>
<td>8.3</td>
</tr>
<tr>
<td>Loperamide (2)</td>
<td>10</td>
<td>-</td>
<td>6.1</td>
</tr>
</tbody>
</table>

Table 3. Antisecretory activity of MeOH extract, EtOAc fraction and isolated compounds from Chiranthodendron pentadactylon

3. Conclusion

Some of the medicinal plants tested showed antidiarrheal activity in the model used. Both extracts of Annona cherimola, Chiranthodendron pentadactylon, Hippocratea excelsa, Ocimum basilicum, Geranimum mexicanum (aerial parts), methanol extract of Ruta chalepensis, Lygodium venustum, Punica granatum, and the aqueous extract of Bocconia frutescens, Aloysia triphylla, Dorsenia contrajerva and Schinus molle showed better antisecretory activity than loperamide. The active extracts found in this study will be an option to develop novel phytodrugs useful to treat fluid loss in diarrhea. These results allow to propose these species as a potential sources of antisecretory compounds and should be therefore subjected to further bioassay-guided phytochemical studies to obtain their active principles, the antisecretory compounds isolated from medicinal plants combined with ORS might be useful in decreasing the mortality caused by dehydration. The properties previously described of (-)-epicatechin suggest that it may be a leading compounds in the development of novel antidiarrheal agents. The results obtained give some scientific support to the use of some medicinal plants tested for the treatment of gastrointestinal disorders such as diarrhea.

4. Acknowledgment

MS Abigail Aguilar, IMSS herbarium, for authentication of plant material and MS Carlos Carrillo. This study was supported by CONACyT (grant: 3800-M); IMSS-FOFOI (FP-2001-05) and PROMEP (PROMEP/103.5/10/7313).

5. References


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The 21st Century has seen a resurgence of research of the gastrointestinal tract, especially since it was established that it plays a central role as an immune system organ and consequently has a huge impact on causation, impact and transmission of most human ailments. New diseases such as the Acquired Immunodeficiency Syndrome, hepatitis and tumours of the gastrointestinal tract have emerged and they are currently subjects of intensive research and topics of scientific papers published worldwide. Old diseases like diarrhea have become extremely complex to diagnose with new and old pathogens, drugs, tumours and malabsorptive disorders accounting for the confusion. This book has set out algorithms on how to approach such conditions in a systematic way both to reach a diagnosis and to make patient management cheaper and more efficient. "Current Concepts in Colonic Disorders" attempts to put all the new information into proper perspective with emphasis on aetiology and providing rational approach to management of various old and new diseases. As the book editor, I have found this first edition extremely interesting and easy to understand. Comments on how to improve the content and manner of presentation for future editions are extremely welcome.