**Antiulcerogenic Effects of Selected African Nightshades**  
* (Solanum nigrum Linn.) * Genotypes on the Rat Stomach:  
A Morphologic and Morphometric Study

Efectos Antiulcerogénicos de Genotipos Seleccionados de Solanáceas Africanas *(Solanum nigrum Linn.)* en el Estómago de Ratas: Un Estudio Morfológico y Morfométrico

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**SUMMARY:** *Solanum nigrum* (SLN), commonly known as African nightshade, is used as a vegetable as well as in the management and treatment of various ailments including gastric ulcers. We analyzed, both grossly and microscopically using H&E, Masson’s trichrome and PSA staining methods, the protective effects of aqueous leaf extracts of three Kenyan SLN genotypes namely *S. scabrum* (SSB), *S. sarachoides* (SSR) and *S. villosum* (SVL) on ethanol-induced gastric lesions in rats. There was evidence of gastro-protection by all the three genotypes with the SSB showing the highest ulcer inhibition score (76.37 %) followed by SSR (72.51 %) and SVL (63.30 %). SLN-pretreated rats showed less areas of gastric mucosal surface erosion. Additionally in the pretreated animals, the depth of the ulcers were markedly reduced, reaching only the gastric pit region except in those treated with SVL where the ulcers penetrated slightly more deeply to affect the gastric glands. Compared with controls, the mean microscopic ulcer index decreased 5.07, 3.55 and 2.37-fold in rats pretreated with SSB, SSR and SVL extracts respectively. Results of this work show extracts of the three SLN genotypes to have antiulcerogenic potential but at varied strengths, thus confirming earlier reports that phytoconstituents and hence the efficacy of a medicinal plant may be influenced by genetic factors.

**KEY WORDS:** Gastric ulcer; Inhibition; Kenyan; *Solanum nigrum.*

**INTRODUCTION**

Gastric ulceration refers to lesions on the glandular part of the gastric mucosa induced by exposure to excessive hydrochloric acid and pepsin activity (Sabiu et al., 2015). Common causes of gastric lesions include *Helicobacter pylori* infection, stress, nutritional deficiencies and chemical injury associated with the use of NSADs, which are commonly used to manage musculoskeletal diseases such as arthritis (Moore et al., 2014). Ulceration studies show the rat to be uniquely susceptible to gastric lesions (Greaves, 2012), which in the current study were analyzed in the glandular region of the stomach (part that is exposed to the highly corrosive acidity of gastric juice). Four main types of epithelial cells cover the surface of the glandular stomach and extend down into the gastric pits and glands. These include mucus-secreting cells, parietal (oxyntic) cells that produce HCl, peptic (chief) cells that secrete proteolytic enzyme pepsin and G (gastrin) cells that synthesize peptide hormone gastrin. Damage to the stomach mucosa leads to impaired function and pain and has often been associated with complications such as gastric bleeding, perforation and obstruction (Milosavljevic et al., 2011).

Gastric ulceration is a common cause of morbidity and mortality, with close to 53 million people developing gastric ulcers in the world each year (Global Burden of Disease Study 2013 Collaborators, 2015). The conventional treatment of gastric ulcers involves use of proton pump inhibitors such as omeprazole and histamine-2 receptor antagonists like ranitidine, both of which act by reducing gastric acid secretion (Fernandes & Marinho, 2019). These drugs are often used in combination with antibiotics, particularly those that have activity against *H. pylori* (Gisbert
et al., 2000). Antibiotic resistance is on the rise and as such, management of gastric ulcers through the above methods is a serious challenge (Wang & Peura, 2011). Other drugs used to treat gastric ulcers are prostaglandin analogues, antimuscarinics and antacids (LoIudice et al., 1981). Use of the aforementioned drugs are associated with adverse effects including arrhythmias (Marcus et al., 2010) and hematopoietic changes (Odou et al., 1999) which appear to limit their use and to trigger the search for alternative drugs.

Medicinal plants are used in the management and treatment of many disease conditions and their use dates back to human civilization (Khan, 2014). The widespread use of medicinal plants has been linked to their ease of accessibility and the common belief that they are less toxic than allopathic drugs (Ernst & Hung, 2011). In developing countries, approximately four billion people rely on herbal products for their healthcare needs (Bodeker et al., 2005). *Solanum nigrum* (SLN) belongs to the family Solanaceae and is characterized by alternate leaves with smooth margins, small and white or pale violet flowers, and green or black berries that grow in bunches (Edmonds & Chewya, 1997). In Kenya and other parts of Africa, this plant is widely used as a vegetable and in managing a host of ailments including gastric ulcers (Edmonds & Chewya, 2005). SLN grown and utilized in Kenya occur in different genotypes/ ecotypes. Thus this study analyzes, through morphologic and morphometric means, the antiulcerogenic potential of some of the Kenyan genotypes in preventing gastric ulceration in the rat.

**MATERIAL & METHOD**

**Plants collection:** Three SLN genotypes namely, *S. scabrum* (SSB), *S. sarrachoides* (SSR) and *S. villosum* (SVL), were collected from the Kenya Agricultural Research Institute (KARI), Muguga, which is located 27 km North West of Nairobi and lies 2096 m above sea level. The rainfall here is bimodal and ranges between 900 and 1000 mm annually. Long rains of about 550 mm are experienced between March and June while short rains of about 400 mm fall between October and December. The temperature ranges between 7 °C and 20 °C with a mean of 15 °C. The area has well drained reddish brown to dark red soil (FAO, 2006). Authentication of the SLN genotypes collected from KARI was performed by a botanist at the Botany Section of the School of Biological Sciences, University of Nairobi.

**Preparation of plant extracts:** Leaves from the various SLN genotypes were harvested green and dried in shed, with constant turning over to avert fungal growth. After seven days of drying, the leaves were pulverized into fine powder using an electric blender. 100g of each of the powdered leaves was boiled in 500 ml of water for 15 minutes. The aqueous extract was then sieved, dried under vacuum in a VirTis Freeze Dryer and the residue weighed and stored at -20 °C until required for testing. Oral doses for the animals were prepared (on the basis of body weight) by dissolving the dried extract in 10 ml of distilled water just before administration.

**Experimental animals.** A total of 30 adult male Wistar rats (220-250 g) were used for this study. These rats were from a stock bred at the Department of Veterinary Anatomy animal facility, where housing of the rats was done in cages with raised floors of wide wire mesh (to prevent coprophagy). Here, the animals were raised under conditions of 12L: 12D cycle, temperature 23±2 °C and humidity 55±15 % and with provision of a balanced diet and free access to water. All protocols for experimentation of the animals were approved by the Biosafety, Animal Use and Ethics Committee of the Faculty of Veterinary Medicine, University of Nairobi (FVM BAUEC/2016/1120) and strictly conformed to the Animals (Scientific Procedures) Act 1986.

**Administration of substances to the animals.** SLN extract dosage used in this study was selected on the basis of an acute toxicity test earlier carried out in mice (Khazaei & Salehi, 2006). To ensure an empty stomach before administration of the different substances, rats in all groups were fasted for 48 hours, but with provision of a nutritive solution of 8 % sucrose in 0.2 % NaCl (until 1hr before administration of the substances) to avoid excessive dehydration during the fasting period (Glavin & Mikhail, 1976). All substances were administered to the animals orally using a stainless intubation needle.

The rats were randomly allotted to five groups of six animals each. Group 1 was the positive control and rats in this group were administered 10ml/kg distilled water prior to ulcer induction. Groups 2, 3 and 4 were the test animals and these received SLN extracts from SSB, SSR and SVL, respectively, at a dose rate of 500 mg/kg and at three different intervals: two doses on the first day at 08:00 h and 16:00 h and a third dose on the second day, 1.5 h before ulcer induction. Group 5 rats were kept as negative/ normal controls (i.e. ulcer induction was not done on them) and these were administered distilled water of equivalent volume to that of the extract administered to the test animals, and only at the same time intervals as the test animals (Glavin & Mikhail).

**Gastric ulcer induction and harvesting of the stomachs.** Induction of gastric ulceration was done on positive control rats (group 1) and the test rats (groups 2-4) using ethanol
(Merck) 50 % (v/v) (in distilled water) at a dose rate of 10 ml/kg given orally via an intubation needle (Glavin & Mikhail). One hour after ethanol-ulcer induction, all animals were sacrificed by intraperitoneal injection of lethal doses of pentobarbital sodium (140 mg/kg bwt). This was followed by perfusion fixation via the heart with 10 % formaldehyde and subsequent removal of the stomach, which was opened along its greater curvature and gently rinsed in physiological saline.

**Macroscopic analysis of the stomachs.** The glandular parts of stomachs of rats in all groups were grossly examined and measurements of lesions done under an illuminated magnifying microscope (at 10x) in order to determine the ulcerative index and the percentage ulcer inhibition/prevention scores as detailed in Alkofahi & Atta (1999). To this end, long lesions were counted and measured along their greater lengths while petechial lesions were counted with the aid of a 1-mm square grid, with each five petechial lesions being considered as 1 mm of ulcer. The sum of the total length of ulcers in each group of rats was divided by its number to obtain the ulcer index (in mm). Where MaUI denotes macroscopic ulcer index, macroscopic ulcer prevention (MaUP) ratio was worked out and expressed as a percentage using the formula:

\[
\text{MaUP} = \frac{[\text{Control MaUI} - \text{Test MaUI}]}{\text{Control MaUI}} \times 100
\]

**Microscopic analysis of the stomachs.** Soon after gross evaluation, the stomachs were further fixed by immersing them in 10 % formaldehyde after which small pieces of the glandular parts were harvested from the antral region. These were further sliced into smaller subsegments and sampled randomly for routine histologic processing. 5mm sections were cut from the selected subsegments and stained by H&E, Masson’s trichrome and PAS methods. The stained sections were then analyzed on a Leica DRM light microscope connected to a monitor. Microscopic Ulcer Index (MiUI) was obtained on a scale of 0 to 3 as described in Khazaei & Salehi. Where 0= normal tissue, 1= local damage to gastric pit cells, 2= local damage to gastric glands and 3= deep damage to gastric glands, the MiU (in mm) was calculated as follows:

\[
\text{MiUI} = (\text{no. lesions 1}) + (\text{no. lesions 2}) \times 2 + (\text{no. lesions 3}) \times 3
\]

**Data analysis.** Data were analyzed using one way analysis of variance (ANOVA) and Dunnett’s pair wise test for intergroup differences. Differences were regarded as significant when p< 0.05. In all cases, data were presented as mean ± SD.

**RESULTS**

**Gross anatomical findings.** Figure 1 shows the magnitude of lesions on the mucosa after ulcer induction and harvesting of the rat stomachs. Positive control rats (those that received distilled water before ulcer induction) presented the most numerous and extensive mucosal lesions. Additionally in this group, the stomachs were markedly congested, hemorrhagic and edematous. Rats pretreated with SLN extracts from SSB, SSR and SVL showed less gastric lesions than controls (Fig. 1).

![Fig. 1. Macrographs of stomachs of rats from the five different groups. A: Negative control rats that received only distilled water before euthanasia. B: Positive control rats administered distilled water before ulcer induction. C-E: Pretreated respectively with SSB, SSR and SVL before ulcer induction. Notice the absence of lesions in A and the massive gastric ulcerations (arrows) in B. There is evidence of ulcer protection by all SLN genotypes as revealed by the marked reduction in number of ulcerative areas (arrows), with the most effective extract being SSB followed by SSR and then SVL, scale bar= 1cm.](image)

In Table I, morphometric data on ulcerative index and ulcer inhibition values are provided in positive control and SLN-pretreated animals. In the control rats, the
Table I. Protective effects of extracts from *S. scabrum*, *S. sarrachoides* (SSR) and *S. villosum* (SVL) on ethanol induced gastric lesions in rats. Values are means ±SD, n= 6/group.

| Treatment                        | Macroscopic Ulcer Index (mm) | Microscopic Ulcer Index (mm) | Ulcer inhibition (%) |
|----------------------------------|------------------------------|------------------------------|----------------------|
| +ve control- water then ulcer induction | 13.46± 2.03                  | 14.5± 2.17                   |                      |
| SSB then ulcer induction         | 3.18± 0.64                   | 2.86± 0.44                   | 76.37                |
| SSR then ulcer induction         | 3.70± 0.73                   | 4.08± 0.75                   | 72.51                |
| SVL then ulcer induction         | 4.94± 0.81                   | 6.12± 0.83                   | 63.30                |

Light microscopic findings. Microscopic view of the rat stomach in negative (normal) controls is presented in Figure 2. These illustrations serve to guide the reader on how depths of penetrating ulcers were traced and graded when working out ulcer index values at light microscopy. The macroscopic ulcer index was estimated at 13.46± 2.03 mm while in the SLN-pretreated group, the magnitude of gastro protection against ulceration varied with genotype, with the percentage ulcer inhibition scores being 76.37, 72.51 and 63.30 in SSB, SSR and SVL, respectively.
The stomach wall is typified, like in other mammals, by a tunica mucosa, submucosa, muscularis and serosa, and with the mucosa having glands that presented distinct pit, neck and body regions. Strands of muscularis mucosa were observed to extend apically in between individual gastric glands (Fig. 2).

In Figure 3, the degree of glandular lesions is demonstrated at histological level. Positive control rats (those administered distilled water before ulcer induction) showed closely spaced glandular lesions on the mucosal surface. Penetrating ulcers in this group of rats pierced through the gastric pits and glands to terminate at the tunica submucosa. In rats pretreated with SSB, SSR and SVL extracts, gastric mucosal surface erosions were notably few and the penetrating ulcers only affect the gastric pits except for SVL where the ulcers extended to the upper region of the gastric glands (Fig. 3). Quantitatively (Table I), microscopic ulcer index in positive controls was 14.5 mm, a value that decreased to 2.86 in SSV, 4.08 in SSR and 6.12 in SVL- treated rats.

Figure 4 is a higher magnification focusing on ulcerated areas of gastric mucosa of positive control rats (i.e. those that received distilled water before ulcer induction). Here, the type of lesions encountered included: (i) mucosal erosion whose interval of occurrence was relatively short (also in Fig. 3), (ii) penetrating ulcers that dug through the pits, neck and body of the gastric glands to reach the muscularis mucosa and to a lesser extent, the tunica submucosa, (iii) death of gastric gland cells, including peptic and parietal cells, in areas of the mucosa penetrated by the ulcers and (iv) degenerative changes involving parts of the muscularis mucosa that were penetrated by ulcers.
more effective (ulcer inhibition = 76.37%) compared to SSR (71.51%) and SVL (63.30%). Our results on antiulcerogenic capacities of the three genotypes of SLN compare favorably with those of earlier investigators. In the study by Saravanan et al. (2011) in which antiulcerogenic effects were analyzed for methanolic extracts of SLN grown in India using aspirin and cold restraint stress as ulcer inducers, respective ulcer inhibition rates were 77.85% and 66.67%. The study by Jainu & Devi (2004) assessed methanolic extracts of berries of an Indian SLN for gastro protection against aspirin-induced ulcerations in rats and reported ulcer inhibition scores of 49.30, 70.12 and 72.70% for extract doses of 250, 500 and 1000 mg/kg respectively.

In the literature, data have also been provided on allopathic drugs ranitidine and famotidine, both H2-antihistamines that act by reducing HCl secretion by parietal cells (Sachs & Scott, 2016). In rats, the percentage inhibition of gastric ulceration by ranitidine in ethanol-induced ulceration is 58.51 (Khazaei & Salehi) while that of famotidine in aspirin induced ulcers is 88.91 (Saravanan et al.). The mechanism of ulcer induction on the gastric mucosa by ethanol entails a host of events including massive production of free radicals and consequential increase in lipid peroxidation which lead to damage of cells and their membranes (Shetty et al., 2000). Tannins, terpenes and fatty acids are some phytoconstituents identified in plants that show gastro protective activity against ulcers (Leite et al., 2009). Hydroxyl radical scavenging and the resultant reduction in lipid peroxidation to offer cytoprotection is the most probable mechanism of action of the aforementioned phytochemicals (Prashanth Kumar et al., 2001). Indeed, we observed here that extracts of the three Kenyan SLN genotypes inhibited ethanol-induced gastric injuries including congestion, hemorrhage, edema and ulcerations in the rat. In the SLN-pretreated animals, light microscopy using H&E, Masson’s trichrome and PAS revealed less penetrating ulcers as well as minimal degeneration of specific gastric gland cells.

In conclusion, this study established that extracts of SLN genotypes SSB, SSR and SVL provide protection against gastric ulceration. The antiulcerogenic potential, when accessed in terms of ulcer index and percentage ulcer inhibition showed genotype-dependent differences. The differences in ulcer inhibition scores may be attributed to genetic factors, which reportedly influence the nature and composition of bioactive ingredients synthesized in medicinal plants (Sharma & Sarkar, 2013). Plausibly, data generated from this work provide additional

DISCUSSION

This study provides, for the first time, experimental data on the impact of aqueous leaf extracts of SLN genotypes SSB, SSR and SVL in a rat model. Our findings revealed that extracts of the above SLN genotypes protect against ethanol-induced gastric ulceration at varied strengths, with SSB being...
knowledge for the popular use of this plant as an antiulcer remedy in Kenyan folk medicine. Future follow up studies should incorporate more SLN genotypes and focus on identifying toxic chemical moiety (-ies) of the extracts, if any, and their specific adverse effects, both short and long term.

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RESUMEN: Solanum nigrum (SLN), comúnmente conocida como la solanácea africana, se usa como vegetal, para el tratamiento de diversas dolencias incluyendo las úlceras gástricas. Analizamos de forma macro y microscópica, de forma macroscópica y microscópica, utilizando para ello tinciones de H&E, tricrómico de Masson y PSA los efectos protectores de extractos acuosos de hojas de tres genotipos SLN de Kenia: S. scabrum (SSB), S. sarrachoides (SSR) y S. villosum (SVL) en lesiones gástricas inducidas por etanol en ratas. Hubo evidencia de gastroprotección por parte de los tres genotipos con el SSB mostrando el punto de mayor inhibición de la úlcer 76,37 % en ratas pretratadas con extractos de SSB, SSR y SVL, respectivamente. Los resultados de este trabajo muestran que los extractos de los tres genotipos de SLN tienen potencial antiulcerogénico en diferentes concentraciones, lo que confirma informes anteriores que los fitoconstituyentes y la eficacia de una planta medicinal pueden estar influenciados por factores genéticos.

PALABRAS CLAVE: Úlcera gástrica; Inhibición; Kenia; Solanum nigrum.

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