Reduction of detrimental effect of soybean oil in-vivo using watermelon white rind extract

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Aim: To study the effect of white rind extract on decreasing soybean oil impact on calcium and phosphorous blood levels *In-Vivo*.

Method: Dried watermelon white rind was directed to mycotoxin and elemental determinations to assure its safe usage. Soybean oil was subjected to fatty acid and GC-MS analysis. Biological experiment was conducted using male albino rats fed diet prepared by soybean oil and supplied with aqueous watermelon white rind extract for two months' interval period. At the end of the experiment, the calcium and phosphorus in blood were determined.

Results: The rind was free from aflatoxin and ochratoxin. Watermelon white rind aqueous extract contained iron, copper, potassium, chromium and selenium at concentration ranges of 3.4, 0.53, 45.51, 0.0142 and 0.0985 ppm, respectively.

Soybean oil had free fatty acid, peroxide value, iodine number and anisidine value of 0.43%, $13.62 \text{ meg } O_2/\text{Kg}$, 132 and 0.7, respectively.

GC-MS analysis of soy oil ascertained the presence of twenty-four compounds: linoleic acid, methyl ester (25.27%), monensin (15.75%), elaidic acid (9.24%), nonadecanoic acid, methyl ester (7.04%), cis-13-eicosenoic acid (4.92%), cis-vaccenic acid (4.68%), linoleic acid (4.67%), palmitoleic acid (4.46%), 9-tetradecenal (4.42%) and cysteine (4.18%)were the most predominant.

Fatty acid profile of the oil showed that the ratio of saturated fatty acid to unsaturated fatty acids was 1:5.

Conclusion: Rats fed diet prepared by soybean oil had a decreased calcium level in comparison with negative control (p<0.05). Supplementation with watermelon white rind aqueous extract rendered calcium level to normal status as negative control. Phosphorus level wasn't affected by soya oil.

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KEYWORDS: watermelon white rind, fatty acid and GC-MS analysis, calcium and phosphorous blood levels.

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INT& ODUCTION

Weatermelon (*Citrullus lanatus* var. *lanatus*, family Cucurbitaceae) is a flowering plant4originally from southern Africa. The white rind is thrown as unused-agro waste. Rind5constitutes 30% of the weight of whole watermelon fruit.

Ola36t al. [1] cited that ethanolic and aqueous extracts of watermelon white rind possessed antibacterial activity against *E. coli and Salmonella sp.* Gas Chromatography-Masse Spectrometry analysis revealed the existence of methionine, L-Aspartic acid, Gly89l-D-asparagine, 9-Cis-Retinoic acid, Stearic acid allyl ester and Ascorbic acid permethyl that contributed to its antibacterial activity.

The rind had total antioxidant activity of 297 mg AAE/100g, total phenols content of 139.42 mg GAE/100g and total flavonoids of 40.4 mg QE/100g. FRAP assay indicated the high reducing ability of the rind. Crude protein content amounted to 13.3%, crude fibe. 44(14.7%) and fat (2.11%). The rind is a source of iron (30.4 mg/kg), potassium (6.9.45%), copper (9.4 mg/kg), chromium (85μg/100g) and selenium (542μg/100g). Unsaturated fatty acid amounted to 81.2%. Vitamins A and E valued 383.44 μg/100g and 43.72 mg/100g, respectively [2]. Wastes are source of sugars, minerals, organic acid. 48 dietary fiber, and bioactive compounds [3].

Soy be an oil affected negatively bone structure as reported by [4].

A stody investigated the adverse effect of soybean oil in rat found that oil induced signsticant fatty liver [5].

In the present work, biological experiment was designed to evaluate safety usage of watened white rind extract on decreasing soybean oil impact on calcium and phosphorous blood levels *In-Vivo*.

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MSTERIAL AND METHODS

₩atermelon white rind was cut into small pieces, dried at 40°C and pulverized into fine 58 owder.

Preparation of white rind aqueous extract:

60 ne gram of dried powder was mixed with one liter of hot water, stirred, filtered and use 6 as the sole source of fluid.

Determination of Aflatoxin and Okratoxin

Total Aflatoxin and Ochratoxin were determined according to AOAC [6]. Total aflat64 ins and ochratoxin A standards were purchased from Sigma (St. Louis, MO, USA). Stock solu65 ns of each mycotoxin were prepared by dissolving solid commercial toxin. The presence of a66 toxins was detected by high performance liquid chromatography (HPLC, Agilent 1200) usin67C18 column of LiChrospher RP-18 (5µm × 25cm). The mobile phase consisted of water: met68 nol: acetonitrile (54:29:17, v/v/v) at flow rate of 1ml/min. The excitation and emission wav69 engths for all aflatoxins were 362 and 460 nm.

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Elemental analysis of rind aqueous extract

ħ2on, copper, potassium, chromium and selenium were determined according to AO★3C [7]. Minerals in the different samples were determined using atomic absorption spec#4ophotometer (Model 2380, Perkin Elmer, Inc., Norwalk, CT, USA).

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Chemical analysis of soybean oil

Quality of oil was assessed by determining anisidine value, iodine number, per Neide value and free fatty acid according to AOAC [7]. Fatty acid composition was determined according to AOAC [7]. The derivatization was conducted following the procedure described by AOAC Official Method 991.39 (AOAC, 2012), with modification in the amcent of initial sample, the kind of extracting solvent and the temperature of heating. GC Ana 23 FAMEs from external standard or FAMEs resulted from sample derivatization were injected separately into Gas Chromatography instrument (GC). The GC analyses were perf84 med on 7890A Gas Chromatography System (Agilent Technologies, California, US) equal 5 ed with flame ionization detector and splitless injector (1 μL). Injector and detector tem 6 rature were set at 270 °C and 280 °C, respectively. The utilized column was a DB-23 (60 m ×80.25 mm, with film thickness of 0.25 μm). This column was purchased from J and W Scie 86 fic (Folsom, CA). The GC oven program was as follows: 130 °C (hold 2 min), to 170 °C at 6.5 °29 min (hold 5 min), to 215 °C at 2.75 °C/min (hold 12 min), to 230 °C at 30 °C/min (hold 30 min) Potelium and nitrogen of ultrahigh purity grade were used as carrier gases at flow rates of 11.09 and 31.24 mL/min.

The **92** emical constituents of the samples were identified using GC (Agilent Tecnologies 7890A) con **92** cted to a mass-selective detector (MSD, Agilent 7000). The flow of helium used as carrier gas **94** s retained at 1 ml/min during the run. The components were confirmed by coordinating thei **95** nass spectra and retention time with the database of National Institute of Standard and Tech **96** logy (NIST) library. The names, molecular weights and chemical structure of each of the com **97** nents of the test materials were determined.

Totalla Aflatoxin and ochratoxin standards were purchased from Sigma (St. Louis, MO, USA 9 Stock solutions of each mycotoxins were prepared by dissolving toxin in the

apparopriate solvent at concentration of 1 mg/mL. AFs in toluene/acetonitrile 99:1 and OTAOm toluene/acetic acid 99:1. Extraction and identification of aflatoxins from

Biological experiment

Eighteen rats were distributed into three groups:

Group4(1) served as negative control and fed normal diet [8] and supplied with drinking watens

Grotop6(2) served as positive control fed normal diet to which 150 ml soybean oil was addetoper kilo and supplied with drinking water.

Grotog(3) fed diet as group (2) supplied with aqueous watermelon white rind extract.

Dietand fluids were supplied ad-libitum for all groups.

At the cend of the experiment, blood samples were collected centrifuged at 4000 rpm and serum was subjected to the analysis of calcium and phosphorus.

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RESULTS AND DISCUSSION

Hemental analysis of watermelon white rind aqueous extract (Table 1) ensured the presente of iron (3.4 ppm), cupper (0.53 ppm), potassium (45.5 ppm), chromium (0.014 ppm1)16nd selenium (0.098 ppm). Aqueous rind extract was a source of mineral needed for had the maintenance as clearly demonstrated.

Dt8a in Table (2) revealed that soybean oil had anisidine value of 0.7, iodine number 131 £3,9 free fatty acid 0.43% and peroxide value of 13.62 meq O₂/Kg. Results ensured low 122 dues of free fatty acid and anisidine and best soybean oil quality.

Tementy-four compounds were detected in the GC-MS chromatogram of soybean oil. Linder acid (25.27%) was the most predominant in the tested oil, followed by montans in (15.75%), elaidic acid (9.24%), nonadecanoic acid (7%), cis-vaccenic acid (4.6824), palmitoleic acid (4.46%), 9-tetradecenol (4.42%) and cysteine (4.18%) and accdented for 59.34% of oil constituent (Table 3).

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Linderic acid is a doubly unsaturated fatty acid, known as an omega-6 fatty acid, occutating widely in plant glycosides. Linoleic acid is an essential fatty acid in human nutriden because it cannot be synthesized by humans [9].

Elai**tio** acid is the major trans fatty acid in margarine and partially hydrogenated oils **181d** also occurs in small amount in cow milk [10].

Nontale canoic acid is a long chain saturated fatty acid derived from plant sources and can be found in fats and vegetable oils [11].

Viaecenic acid, an isomer of oleic acid, is the principal ruminant *trans* fatty acid. It is produced through the biohydrogenation of linoleic acid and α -linolenic acid by mice 136 reganisms in the rumen and is found naturally in foods such as dairy and ruminant meat 37 roducts.

A38 in Table (4), fatty acid profile of soybean oil showed the existence of linoleic acid (894.28%), oleic acid (22.85%), linoleic acid (6.2%) and gadolic acid (0.21%) as unsatatoated fatty acids accounting for 83.54% of total oil content. Saturated fatty acids comparised palmitic acid (10.99%), stearic acid (4.82%), arachidic acid (0.36%) and behavior acid (0.29%) representing 16.46% of soybean oil content. These results are in accordance with Friedman and Brandon [12] who stated that soybean had low level of saturated fat and high content of linoleic acid [13].

∆45shown in Table (5), a significant difference (p<0.05) existed between negative con**tr46** (G1) and rats group fed diet with soybean oil (G2). A decrease in calcium level was **1**√2 served indicating that soybean oil affected calcium blood level.

Soybean had high phytate level [14]. Phytates can block the uptake of essential minerals as calcium, copper, iron, zinc and magnesium in intestinal tract that may contribute to mine sol deficiencies [15].

There was non-significant difference between negative control (G1) and Group 3 fed soy**b52**n oil and drunk rind extract, nor between G2 and G3.

Data 52 vealed that phosphorus blood level was not affected by any treatment and non-significant differences existed between G1 and both groups G2 and G3.

The 155 traction of plant material and isolation of biologically active compounds are essets to understand their role in disease prevention and treatment.

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CONCLUSION

Wattathelon white rind aqueous extract is a source of iron, copper, potassium, chronolium and selenium. Soybean oil decreased blood calcium level, while phosphorus was 182 ble in all treated groups. Supplementation with watermelon white rind aqueous extract calcium level to normal status as negative control.

Table (1): Elemental analysis of watermelon rind aqueous extract

| Element | Result |
|-----------------|--------|
| Copper (ppm) | 0.5 |
| Iron (ppm) | 3.4 |
| Potassium (ppm) | 45.5 |
| Chromium (ppm) | 0.014 |
| Selenium (ppm) | 0.098 |

Table (2): Chemical evaluation of soy oil

| Tested parameters | Result |
|--|--------|
| Free fatty acid (%) | 0.43 |
| Peroxide number (meq O ₂ /Kg) | 13.62 |
| Iodine number | 131.8 |
| Anisidine value | 0.7 |

Table (3): GC-MS analysis of soy oil

| RT | Compound name | Area sum (%) |
|------|---------------|--------------|
| 3.88 | Chicoric acid | 0.29 |

| 5.7 | Phytanic acid | 0.59 |
|--------|--|-------|
| 6.187 | 3,2',4',5'-Tetramethoxyflavone | 0.27 |
| 8.04 | Gardenin | 0.49 |
| 8.96 | Isovitexin | 0.59 |
| 11.7 | Lutein | 1.33 |
| 12.03 | Stevioside | 0.57 |
| 13.23 | Hexadecanoic acid, methyl ester | 2.63 |
| 13.43 | Pentadecanoic acid | 0.73 |
| 13.5 | Monensin | 15.75 |
| 13.9 | Zearalenone | 1.59 |
| 14.17 | Oleic acid | 2.83 |
| 14.35 | Cis-vaccenic acid | 4.68 |
| 14.52 | Linoleic acid, methyl ester | 25.27 |
| 14.59 | Elaidic acid | 9.24 |
| 14.66 | Cis-13-eicosenoic acid | 4.92 |
| 14.75 | Nonadecanoic acid, methyl ester | 7.0 |
| 14.93 | Linoleic acid | 4.67 |
| 15.14 | Quinine | 0.5 |
| 15.33 | 3-(3,4-dimethoxyphenyl)-4,6-dimethylcoumarin | 0.98 |
| 15.9 | Di-γ-linolenin | 1.97 |
| 16.009 | Palmitoleic acid | 4.46 |
| 16.04 | Cystine | 4.18 |
| 16.79 | 9-tetradecenal, (Z)- | 4.42 |

Table (4): Fatty acid analysis of soybean oil

| Palmitic acid C16:0 | Saturated fatty acid | 10.99% |
|------------------------|------------------------|--------|
| Stearic acid C18:0 | Saturated fatty acid | 4.82% |
| Arachidic acid C20:0 | Saturated fatty acid | 0.36% |
| Behenic acid C22:0 | Saturated fatty acid | 0.29% |
| Oleic acid C18:1n9 | Unsaturated fatty acid | 22.85% |
| Linoleic acid C18:2n6 | Unsaturated fatty acid | 54.28% |
| Linolenic acid C18:3n3 | Unsaturated fatty acid | 6.2% |
| Gadolic acid C20:1ω9 | Unsaturated fatty acid | 0.21% |

Table (5): Serum calcium and phosphorus levels in treated rat groups

| Groups Parameters | Group 1 (n=6) | Group 2 (n=6) | Group 3 (n=6) |
|--------------------|---------------|------------------|---------------|
| Calcium (mg/dl) | 13.2±0.64 | 11.3±0.48 * | 12.8±0.62 |
| Phosphorus (mg/dl) | 10.5±0.66 | 10.38±0.76 | 11.96±0.44 |

*Significant difference (p<0.05) in comparison with negative control

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