HYDROETHANOLIC EXTRACTS OF FICUS PUMILA LINN. IS PROTECTIVE AGAINST GENATAMICIN-INDUCED KIDNEY DAMAGE IN RATS

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Original Research Article

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HYDROETHANOLIC EXTRACTS OF FICUS PUMILA LINN. IS PROTECTIVE AGAINST GENATAMICIN-INDUCED KIDNEY DAMAGE IN RATS

7 Abstract

8 Ficus pumila Linn, has been reported to be rich in phenols, hepatoprotective and antiproliferative on leukemic cancer cells. The aim of this study was to evaluate the 9 10 nephroprotective effect of hydroethanolic leaves extracts of F. puntila on gentamicin-induced kidney damage in rats. Twenty seven female Wistar albino rats were divided into 9 groups 11 (n=3). Group 1 being normal; group 2 was the gentamicin (GM) induced only (80 mg/kg b/w 12 ip for 5 days); groups 3, 4, & 5 rats were treated with gentamicin (80mg/kg b/w ip for 5 days) 13 and F. pumila extract at 100, 250, and 500mg/kg b/w orally respectively; groups 6, 7 & 8 rats 14 received the extract only (100, 250, and 500mg/kg b/w orally) respectively and group 9 being 15 gentamicin and silymarin (100 mg/kg b/w orally) for 21 days. Blood samples were taken 24 16 17 hrs after the experimented period and biochemical and haematological parameters were analyzed. GM nephrotoxicity was characterized by a significantly increased levels of serum 18 creatinine, urea, sodium, potassium and WBC, while reduced RBC, HGB, MCH and MCV 19 levels compared with normal group. Rats treated with gentamicin and the extract showed a 20 significant reduction in the levels of these markers. The results suggest that hydro-ethanolic 21 22 extract of Ficus pumila leaves protect against gentamicin-induced nephrotoxicity in female 23 Wistar albino rats.

24

25 Keywords: Nephrotoxicity, Ficus pumila Linn. Creatinine, Urea, electrolytes

26 Introduction

Nephrotoxicity is known to be one of the most common kidney problems worldwide. It occurs 27 when the body is exposed to high dosages of a drug or a toxin. Kidney damage is characterized 28 by increased levels of serum urea and creatinine and imbalance of blood electrolytes such as 29 potassium and magnesium [1]. Aminogly coside antibiotics are commonly used in the treatment 30 31 of bacterial infections. They have potent antibacterial activity against infections produced by 37 gram negative bacteria [2]. Gentamicin is an aminoglycoside antibiotic isolated from the 33 bacterium Micromonosporu purpurea. It has a hexose ring to which various amino sugars are attached by glycosidic linkages [3]. Despite its clinical benefits, it is known to be the most 34 35 nephrotoxic of all the aminogly cosides [4]. Gentamicin-induced nephrotoxicity is indicated by 36 elevated levels of plasma creatinine and urea with severe necrosis of the renal proximal convoluted tubules followed by failure of renal functions [5]. According to Al-Majed et al. [6]. 37 its nephrotoxicity is as a result of the selective accumulation of reactive oxygen species in renal 38 39 cortical areas leading to damage of membranes. 40

- 41 Some species of the Moraceae have been shown to possess significant nephroprotective
- 42 activity. They include F. religiosa latex on cisplatin [7], F. dalhousiae leaf methanolic extracts
- 43 on gentamicin and acetaminophen [8], F. carica leaf extract on gentamicin [9], F. racemosa

aqueous bark extract on gentamicin [10] and F. benghalensis latex on cisplatin [11]. Ficus 44 pumila Linn. is a creeping vine-like fig plant which also belongs to the family Moraceae. It is 45 native to South and cast China. Malaysia, Vietnam and Africa [12], F. pumila is ingested to 46 47 treat conditions such as diabetes, dizziness, skin diseases and high blood pressure [13]. The 48 hydroethanolic extract of Ficus pumila L. is a rich source of tannins, saponins, general glycosides, alkaloids, flavonoids, triterpenes, and sterols and has been demonstrated to be 49 50 hepatoprotective in animals [14,15], and it is a potent anticancer agent. The leaves of this plant have been shown to have antioxidant, antimicrobial, anti-mutagenic, anti-inflammatory and 51 analgesic activities [14, 16]. 52

53 The aim of this study was to determine the nephroprotective effect of the 50% aqueousethanolic leaves extract of *Ficus pumila* Linn, in gentamicin-induced kidney damage in female Wistar albino rats.

56

57 Materials and Methods

58 Plant collection and authentication

59 The leaves of *Ficus pumila Linn*, were collected in October, 2015 from the Republic Hall, 60 Kwame Nkrumah University of Science and Technology (KNUST) Campus. They were 61 identified based on voucher specimen deposited at the herbarium of the Department of Herbal 62 Medicine (KNUST, Kumasi, voucher number KNUST/HM1/2014/L093).

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64 Extract Preparation

The plants were washed, shade-dried for a month, and milled. 50% ethanol extraction of the plants were carried out by suspending 100 grams of the powder in 1000 ml of 50% ethanol (50: 50 ethanol, water, v/v). The leaves-solvent mixtures were allowed to stand for 24 hours at room temperature on a shaker. The extracts were filtered through cotton wool and concentrated using a rotary evaporator under reduced pressure. They were transferred into sterile bottles and freeze dried to obtain the *Ficus pumila* ethanolic leaf extract (FPE). The extract was dissolved in

- 71 distilled water at respective doses and used for the study.
- 72

73 Animal Model

74 The study was performed on twenty-seven female Wistar albino rats (150-200g). They were 75 obtained from the SMS-UG. Accra and kept at the animal holding facility at the Department of Biochemistry and Biotechnology, KNUST-Kumasi. The animals were labeled, housed in a 76 77 clean standard metal cage and had free water and standard rodent feed (Agricare, Kumasi, Ghana) ad libitum at room temperature. Food intake by animals was monitored daily. All 78 79 animal experiments were conducted in accordance with the guidelines of the Committee for 80 the Purpose of Control and Supervision of Experiment on Animals (CPCSEA, New Delhi, India) and guide for care and use of laboratory animals (Washington, US). 81

82 Experimental Drug

83 Gentamicin injection (Letap Pharmaceuticals, Ghana) at 80 mg/kg body weight was administered to the rats intraperitoneally (ip) from the 16th -20th day of treatment to induce

85 kidney damage.

86 Experimental Design

87 The rats were divided into 9 groups with 3 animals in each group. The groups were divided as

- 88 follows: Group I rats served as normal control and received I ml/kg b/w distilled water
- 89 throughout the duration of the experiment, Group II were injected with gentamicin, Group III,

90 IV and V rats were treated with gentamicin and FPE (100, 250 and 500 mg/kg body weight 91 respectively). Groups VI, VII and VIII rats were also treated with FPE only at a dose 100, 250 92 and 500 mg/kg body weight respectively. Group IX were treated with gentamicin and silymarin 93 (100 mg/kg body weight). The experiment was terminated with an overnight fast at the end of 94 21 days. The rats were sacrificed after mild ether anesthesia. Incisions were made in the 95 cervical region of the animals and blood samples were taken for biochemical and 96 haematological analysis.

97 Effect of Treatment on Body Weight

C1

98 Body weight of the rats were taken every two days and percent change in body weight 99 calculated with the following formula:

100
$$Percent Chnage in Body Weight = \frac{Weight_n - Weight_{initial}}{Weight_{initial}} \times 100$$

101 where Weight₀ is the body weight on Day 4, D8 D21 and Weight_{annal} is the body weight on 102 D0

103 Effect of Treatment on Kidney Weight

104 The kidneys of sacrificed animals were excised, washed in buffered saline and blotted with

105 paper tissue. They were weighed to obtain the absolute organ weight (AOW). The Relative

106 Organ Weight (ROW) was calculated with the following formula:

107 Relative Organ Weight (%) =
$$\frac{Absolute Organ Weight}{Body Weight at Sacrifice} \times 100$$

108 Assessment of Kidney Function

109 The blood samples were collected into clean sterile tube and left to stand for an hour and 110 centrifuged at 3000g for 15 minutes at 5°C to separate the serum for biochemical analyses 111 which included urea, creatinine, electrolytes, cholesterol, fasting blood glucose, alanine 112 aminotransferase (ALT) and total protein using the Cobas Integra Autoanalyser and kits

113 (Fortress Diagnostics, UK).

114 Haematological Analyses

Part of the blood sample was placed in EDTA tubes for haematological analyses which included red blood cell (RBC), hemoglobin (HGB), hematocrit (HCT), mean cell hemoglobin concentration (MCHC), mean cell volume (MCV) and platelets (PLT) count using the Sysmex KX21N autoanalyzer to run a full blood count in the whole blood mode.

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120 Statistical Analysis

121 Data was analysed using GraphPad Prism 5 for windows. The results were expressed as the 122 Mean ± Standard error mean (SEM). One – way Analysis of variance followed by Newman-123 Keuls multiple comparison test was used for comparison between groups (i.e. control and 124 treated groups). All statistical tests were run at a 95% confidence interval and values of P< 0. 125 05 were considered statistically significant. Percentage protection was calculated with 126 following formula based on significant indicators of nephroprotection including urea, 127 creatinine,

128

Percent Protection = 100 + Values of Toxin Contol - Values of Test sample (Values of Toxin Control - Values of Normal Control) 129

130 Results

131 Effect of treatment on body weight

Table 1 shows the effect of treatment on the body weight of the rats. There was a reduction in the body weight of rats treated with gentamicin only compared with the normal. However, the body weight of groups treated with plant extract only was almost the same as the normal but comparing the body weights of groups treated with gentamicin and plant extract at varying concentration to the gentamicin only group, a decrease was observed.

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138 Effect of treatment on relative kidney weight

- 139 Figure 1 shows the effect of the treatment of FPE on relative weight of the kidneys.
- 140 Administration of FPE and GM to the animals did not provoke any significant increase in the 141 relative kidney weights.



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143 Fig. 1: Effect of treatment on kidney weight. Each column represents a mean # SEM.

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145 Effect of treatment on some biochemical parameters

Treatments

146 Table 2 shows the biochemical data obtained for the normal and treated rats. The rats to which 147 GM only was administered showed a significant increase in the blood urea, serum creatinine, 148 total protein and fasting blood sugar levels and a decrease in ALT levels compared to the 149 normal. Those parameters however, had reduced levels in the groups that were treated with 150 FPE and GM suggesting nephroprotection, while GM significantly reduced the serum 151 potassium, sodium and chloride levels as compared to normal. The electrolyte levels were 152 however significantly increased in the treated groups.

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154 Effect of treatment on haematological parameters

- 155 Table 3 shows the effect of treatment on some hematological parameters. There were no
- 156 significant changes in the haematological parameters assayed excepted a significant increase 157 in animals treated with both GM and extract.
- 158 Percentage Protection

159 Fig. 2 shows the percent protection of extract alone and with GM on the kidney. The extract at

160 all doses protected the kidney (94-99%). With GM, only the 250 mg/kg showed a good

161 protection of 58%.

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163

164 Fig. 2: Effect of treatment on percent liver protection

165 DISCUSSION

Owing to the increasing kidney disease burden annually and the high cost of treatment, there 166 167 is the need to develop new therapies to overcome these challenges. Therefore, in this study, the nephroprotective effect of the aqueous-ethanolic leaves extract of F. pumila Linn, was 168 investigated. Administration of gentamicin (80 mg/kg b/w ip) for 5 consecutive days caused 169 marked nephrotoxicity as is evident from Table 2, showing significant increase in serum 170 creatinine (332.80 mg/dL ±12.96 mg/dL at p< 0.0001) and serum usea (261.50 mg/dL ± 26.32 171 mg/dL at p<0.0001) compared with normal serum creatinine (34.60 mg/dL ± 2.428 mg/dL) 172 173 and urea (59.12 mg/dL ± 2.43 mg/dL). The elevation of the serum creatinine is produced by 174 kidney damage, which lead to a decreasing glomerular filtration rate (GFR) and serum 175 creatinine filtration. The increase in the serum creatinine levels in the gentamicin (GM) treated 176 group is due to decreased GFR caused by the gentamicin [17]. The gentamicin nephrotoxicity was significantly protected in groups treated with gentamicin and the FPE and the 250mg + 177 178 gentamicin group reduced the urea and creatinine levels even better than the Silvmarin (test 179 drug used). The results thus indicated that FPE is effective in reducing serum creatinine and 180 urea level in gentamicin toxicity. According to Larbie et al. [14], the hydroethanolic extract of FPE had significant antioxidant activity and contains tannins, saponins, general glycosides, 181 182 alkaloids, flavonoids and triterpenes. The nephroprotective effects of FPE in GM-induced 183 nephrotoxicity may be due to flavonoids and tannins present in the extract. These findings are in accordance with those reported earlier in which Ficus carica fruit extract caused marked 184 reduction in serum urea and creatinine levels in GM-induced nephrotoxicity [18]. Serum 185 potassium, chloride and sodium were significantly reduced in groups treated with gentamicin 186 187 only compared with normal which indicated kidney damage since the kidneys are involved in osmotic and ion balance in the body, therefore an imbalance in serum electrolytes was 188 189 indicative of kidney damage [19]. The effects induced by GM were significantly prevented by

190 FPE which further buttress the fact that this plant has the potential to be used to ameliorate 191 gentamicin nephrotoxicity. Again FBG and total protein increased while ALT decreased in 192 groups treated with gentamicin only compared with normal. This can also be attributed to the 193 fact that gentamicin is known to be nephrotoxic rather than hepatotoxic.

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There was observed decreases in RBC indices (HCT, MCH, MCHC, PLT and HGB) in rats treated with GM only as compared to the normal, possibly indicating an impairment of kidneys because at normal conditions the kidneys produce enough of crythropoietin for the production of red blood cell [19]. On the other hand, the aqueous ethanolic extract of the leaves of *Ficus pumila* was able to increase the levels of these parameters upon treatment. This protection may be because the plant extract was able to increase the production of crythropoietin to enhance the production of red blood cells in the bone marrow.

Balakumar et al. [20] revealed that gentamicin in the cytosol acts on mitochondria directly and 202 indirectly to activate the intrinsic pathway of apoptosis, interrupts the respiratory chain, impairs 203 ATP production and causes oxidative stress by increasing superoxide anions and hydroxyl 204 radicals which further contribute to cell death. This means that gentamicin administration 205 206 enhances the production of free radicals indicating oxidative damage at the cellular level of the 207 renal cortex. Other manifestations of gentamicin nephrotoxicity include electrolyte imbalance and water and non-electrolyte transport in a variety of cells and tissues, the principal target 208 209 organ being the kidneys. Flavonoids, one of the phytochemical constituents of the leaves of 210 Ficus pumila Linn, has been reported to show strong antioxidant activity [14]. This may 211 account for the mechanism of the nephroprotective effect of Ficus pumila. In addition, the 212 extract was observed to restore electrolytes to near normal levels in treatment group. 213 Summarizing all these facts, it can be said that these phytoconstituents are responsible for the 214 observed biological protective effect in this study.

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216 CONCLUSION

217 In conclusion, this study gives the experimental evidence that the aqueous ethanolic extract of 218 the leaves of *Ficus pumila* Linn, was able to produce considerable protection from the 219 nephrotoxic action of gentamicin in female Wistar rats. Further studies will be required to 220 understand the mechanism of protection and also its protective effect against other nephrotoxic 221 agents.

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Table 2: Effect of FPE on biochemical parameters in gentamicin induced nephrotoxicity.

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Table 3: Effect of treatment on unne himmalishelical parameters

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