Original Research Article

EVALUATION OF ACUTE AND CHRONIC TOXICITY OF TARTRAZINE (E102) ON STERIOD REPRODUCTIVE HORMONES OF ALBINO RATS

ABSTRACT

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Aim: To determine the acute and chronic effect of tartrazine on reproductive steroid hormones of albino rats.

Study design: The design involved acute and chronic study. The acute study investigated intraperitoneal and oral route of administration while the chronic study used oral route only. The rats used for the study weighed 150gm approximately. In the acute study, 48 rats (24 female and 24 male) were used for intraperitoneal treatment and were randomly selected into six groups treated with 0.0g/kg, 1.67g/kg, 3.33g/kg, 5.0g/kg, 6.67g/kg and 8.33g/kg of tartrazine. In orally treated rats, 48 rats (24 female and 24 male) were also used and were treated with 0.0g/kg, 5.0g/kg, 10.0g/kg, 15.0g/kg and 20.0g/kg of tartrazine. In the chronic study, the experiment was divided into phase 1, 2 and 3 which lasted for 30, 60 and 90 days respectively. In each phase, 80 rats were used and were divided into treatment and control groups. The treatment groups were given 7.5mg/kg of tartrazine orally on daily basis over a period of 30, 60 and 90 days while the control groups were not treated with tartrazine. **Place and Duration of Study:** The study was carried out in the Department of Medical Laboratory Science, Rivers State University, Port Harcourt, Nigeria over a period of 12 months (December, 2017 – December, 2018).

Methodology: At the end of the acute and chronic study, 5mls of whole blood specimens was collected by means of cardiac puncture into plain bottles. The specimens were spun at 4500 rpm for 10 minutes to obtain serum. The laboratory analysis of the hormonal parameters was based on Enzyme Linked Immunosorbent Assay (ELISA) Technique. Statistical analysis was performed using GraphPad Prism version 5.03 (San Diego, California, USA).

Results: In acute study, female treated rats (intraperitoneally and orally) showed significantly higher values in Progesterone (PROG) and Estradiol (E2) concentrations while male treated rats (intraperitoneally and orally) indicated significantly lower values in testosterone (TESTO) concentration compared with control rats. In chronic treatments, hormonal parameters after 30 days, 60 days and 90 days showed no significant differences in testosterone (TESTO), Progesterone (PROG) and Estradiol (E2) concentrations in tartrazine treated rats compared with their respective control rats. When the comparative analyses of treated groups after 30, 60 and 90 days using One-Way ANOVA were considered, testosterone (TESTO) concentration indicated significantly lower levels in treated male rats while Progesterone (PROG) showed significantly higher values over 30, 60 and 90 days in treated female rats. Conclusion: In the acute study, reduction in testosterone (TESTO) concentration while increase in PROG and E₂ concentrations were seen. However, in the chronic study, significant differences were not seen in testosterone (TESTO), Progesterone (PROG) and Estradiol (E2) concentrations. Finally, when the influence on duration of exposure at ADI doses (7.5mg/kg) were considered after 30, 60 and 90 days, reduction in testosterone (TESTO) and increase in Progesterone (PROG) concentrations were seen.

14 Keywords: Tartrazine, reproductive hormones, Progesterone, Testosterone, Estradiol.

16 **1. INTRODUCTION**

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Colours are important components of food and food products which gives the first impression on the psyche of the consumer [1, 2]. Though food dyes occur in natural and synthetic forms, lately synthetic food dyes are commonly used in food industries because of their availability, cost effectiveness and stability. They are mainly organic compound (Azo dyes) with the capacity to reflect light. Examples include tartrazine, erythrosine, fast green, carmoisine and so on [3, 4]. The application of dyes is also seen in textile, leather, paper, rubber, cosmetics and even in pharmaceutical industries [4].

The use of synthetic food dyes has been reported to cause renal derangements, hepatotoxicity, anaemia, leucopenia and interference with enzymes activities resulting in reduced enzymes functions when studied in rats [2, 4, 5, 6]. However, according to the Australian Government through her Department of Health [7] in a scientific review report in 2014 stated that synthetic food dyes possesses no harmful effect within the acceptable daily intake (ADI) doses.

33 A review of literature reveals that almost all synthetic food dyes originate from coal tar which 34 is toxic and carcinogenic [6]. The toxicity of synthetic dyes such as tartrazine has been 35 linked to the reductive biotransformation of the azo bond during their metabolism in the 36 intestine and liver producing reactive amines, aryl amines and free radicals [8]. These dyes 37 have also been reported to react with proteins (enzymes) covalently which leads to distortion 38 of the protein active site and configuration [5]. Though the effects of synthetic dyes on 39 reproductive parameters and organs remains controversial, very few studies have reported reproductive derangements in rats [9, 10, 11], while other studies reported no reproductive 40 41 derangements [12, 13, 14]. Reproductive parameters (Hormones) considered in this study 42 include, testosterone, estrogen and progesterone.

44 Testosterone is the main androgen hormones secreted by the mature testes [15, 16]. It is an 45 important steroid hormone that play vital role in the production and maturation of 46 spermatozoa, development, growth and differentiation of male sex organs, sexual drive and 47 secondary sexual characteristics that make sexual reproduction feasible in males [17]. In the absence of injury or toxicity of the testicular cells, there is no sharp reduction or fall in testosterone production [17]. Several chemicals and drugs have been reported to induce 48 49 50 adverse effects on the reproductive organs [17]. Sies et al. [18], Ashida et al. [19], reported 51 that tartrazine stimulates mutagenic processes and decreases cell viability. However, 52 Mehedi et al. [9], Gautam et al. [20], reported that sperm production and sperm motility were 53 decreased when xenobiobitics such as azo dyes were fed to rats.

54 55 Progesterone is one of the principal hormones secreted by the ovaries and produced mainly by the corpus luteum under the influence of Luteinizing hormone [21, 22]. The major role of 56 progesterone involves the transformation of the proliferative endometrium in the secretary 57 58 phase, which is necessary for implantation of fertilized egg [22]. Progesterone also 59 enhances the viscosity of the cervical mucus making it more viscous and less permeable and therefore play vital role in establishing pregnancy after implantation of fertilized egg(s) 60 [22]. Estrogen is produced mainly by the granulosa cells of the developing ovarian follicle in 61 the early part of the ovarian cycle and from the luteinized granulosa cells in the corpus 62 63 luteum after ovulation [21]. The principal biological active form of estrogen is the 17β-64 estradiol [21]. The major function of estrogen includes promotion of growth and development 65 of secondary sexual characteristics in the female such as growth and development of the 66 oviducts, uterus, vagina, external genitals, among others making sexual reproduction

feasible [21, 22]. According to Foster & Gray Jr., [17], exposure to xenobiotics has been 67 68 implicated in the decline of normal fertility and reproduction. Takana [12], reported that 69 773mg/kg of tartrazine dyes fed to rats in diet had no adverse effect on reproductive 70 parameters such as steroid hormones. However, Mehedi et al. [10] and Sharma et al. [23], reported in their separate studies that 2.5% of tartrazine induced significant weight reduction 71 72 of the ovaries as well as significantly lowered concentration of steroid hormones in rats.

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74 This study is particularly relevant in our society because the exposure to food dyes cuts 75 across almost everyone due to their diverse applications in the various industries and especially in the food industries (restaurants, fast food, and domestic use, among others). 76 77 Secondly, studies on toxic effect of synthetic dyes on reproductive hormones even at the 78 recommended acceptable daily intake (ADI) are still controversial, quite minimal and 79 obscure. Sreenivasa et al. [24], reported that infertility and hormonal imbalances are on the 80 increase with global record of 75 million couples suffering infertility annually of which 15% 81 are idiopathic. Therefore, this research is aimed at using acute and chronic toxicity studies to perform toxicological evaluation of tartrazine toxicity on steroid reproductive hormones using 82 83 albino rats. 84

85 2. MATERIALS AND METHODS

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2.1 Materials 88

89 Materials used in this research include Polypropylene gavage tubes (Intech Laboratory Incorporated, Plymouth Meeting, USA), Haier thermocool refrigerator (China), MPW bucket 90 centrifuge Model 351 (MPW Medical Instruments, Warsaw, Poland), Ohaus Scout-Pro 91 92 Electronic weigh balance (Ohaus Corporation, New Jersey, USA), Albino rats, Stat Fax 4200 93 Microplate Reader (awareness, USA), Tartrazine dyes (CI. 19140, CAS No 1934-21-0, MW 534,37, E102, FD& C NO 5) with serial no of FI19371 purchased in a granular form from 94 Fiorio Colori Spa, Gessete, Italy, with purity of 86.7% guaranteed by the manufacturer. 95 96 Progesterone, Estradiol and Testosterone Enzyme Linked Immunosorbent Assay (ELISA) 97 kits were purchased from BioCheck diagnostics (San Francisco, USA). Other materials used 98 include automatic pipettes and glass test-tubes. 99

100 2.2 Experimental Animals

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Male and female albino rats used for the study weighed 150gm approximately. The reason 102 for selecting male and female rats for the study was based on the fact that male predominate 103 hormone is testosterone while female predominate hormones are estradiol and 104 105 progesterone. All the rats used for the experiment were obtained by breeding. However, the parent rats used for the breeding were purchased from the University of Port Harcourt. River 106 107 State, Nigeria. The rats were fed with rat pre-mix rat feed and water ad libitum. The animals were placed in a well-ventilated rat cages with water cans and feed containers in place.

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110 2.3 Preparation of Tartrazine Food Dye

111 112 In the acute study, for intraperitoneal treatment, 250 grams of the tartrazine was weighed 113 and dissolved in a sterile container containing 1 litre of distilled water. This implies that 1.0ml of this solution contains 0.25 grams. In terms of oral treatment (acute study), 375grams of 114 115 the tartrazine dyes was also dissolved in sterile containers containing 1 litre of distilled water. 116 This implies that 1.0ml of this solution contains 0.375grams of tartrazine. Finally, in the chronic study, 1.13 grams of tartrazine was weighed and dissolved in a sterile container 117 containing 1.0 litre of distilled water. This implies that, 1.0ml of the tartrazine solution 118 119 contains 0.00113grams and which is equivalent to 7.5mg/kg when administered into a

0.15kg rat. The contents of the containers were properly mixed to ensure complete mixture
 before administration.

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2.4 Experimental Design and Administration of Food Dyes

124 125 The method of treatment in the acute studies involved both intraperitoneal and oral 126 techniques while in the chronic study, treatment was strictly orally. In the intraperitoneal 127 method, the dyes were injected into the intraperitoneal space of the rats using 2 ml and 5 ml 128 hypodermic syringes while in the oral method, the food dyes were administered using 129 orogastric tube to ensure complete delivery of the dye.

2.4.1 Acute treatment and Toxicity Study

132 133 Dose range of the tartrazine dye were determined after the obtaining the value of LD₅₀ using the arithmetic method of Karber as described by Dede et al. [25], in both oral and 134 135 intraperitoneal treated rats. The LD₅₀ was calculated to be 5.83g/kg and 11.25g/kg for intraperitoneal and orally treated rats respectively. In the intraperitoneal treatment, 48 rats 136 137 (24 male & 24 female rats) were used. The male and female rats were randomly selected 138 into six different groups separately designated as ATIP (control), BTIP, CTIP, DTIP, ETIP and FTIP and were treated with 0.0g/kg, 1.67g/kg, 3.33g/kg, 5.0g/kg, 6.67g/kg and 8.33g/kg of 139 140 tartrazine respectively. In terms of orally treated rats, 48 rats (24 males; 24 females) were 141 also used. The male and female rats were randomly selected into six different groups 142 separately. The groups were designated as A_{TO} (control), B_{TO} , C_{TO} , D_{TO} , E_{TO} and F_{TO} and were orally treated with 0.0g/kg, 2.5g/kg, 5.0g/kg, 10.0g/kg, 15.0g/kg and 20.0g/kg of tartrazine respectively. At the end of the 24 hours acute toxicity testing, blood samples were 143 144 145 collected for hormonal assay after the animals were sacrificed.

147 <u>2.4.2 Chronic Treatment and Toxicity Study</u> 148

In the chronic study, the experiment was divided into three phases depending on the 149 duration of exposure of the rats to tartrazine dves. The phase 1. 2 and 3 of the chronic 150 toxicity studies lasted for a duration of 30, 60 and 90 days respectively. Eighty (80) 151 152 experimental rats weighing approximately 0.15kg were used in each phase of the study (with 153 a total of 119 females and 116 male rats of which 5 died in the course of the experiment). In 154 each phase of the experiment, the rats were divided into two groups designated T_T (tartrazine treated group), and C (control, untreated group). Rats in each of these groups 155 were further distributed randomly into ten cages with four rats per cage, designated T_{T1}, 156 157 T_{T2...}T₁₀. In the treatment pattern, the acceptable daily intake (ADI) of 7.5mg/kg of tartrazine 158 was administered orally. The control group, were not treated with tartrazine. At the end of the 159 chronic study, the animals were anaesthesized with chloroform and blood samples collected 160 by means of cardiac puncture for hormonal investigations. 161

162 **2.5 Study Area**

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The study was carried out in the Department of Medical Laboratory Science, Rivers State University, Port Harcourt. However, samples were transported in frozen form in a thermoregulatory container to the University of Port Harcourt Teaching Hospital. However, prior to the actual assay, the serum samples were allowed to defrost at temperature. All of the hormonal parameters considered were analysed at the Chemical pathology Unit of the University Teaching Hospital.

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171 2.6 Specimen Collection, Preparation and Analysis

173 At the end of the study, the animals were anaesthetized with chloroform and 5mls of blood 174 samples was collected by means of cardiac puncture into plain bottles for hormonal assay. The specimens were spun at 4500 rpm for 10 minutes to obtain serum which was 175 176 transferred into other sets of labelled plain bottles and stored at -4°C. The laboratory analysis of the hormonal parameters was based on Enzyme Linked Immunosorbent Assay 177 (ELISA) Technique. The ELISA procedure (outlined by BioCheck Diagnostics, San Francisco, USA) for the determination of Progesterone, Estradiol and Testosterone 178 179 concentrations were based on method described by Engvall and Perlmann [26]. The 180 181 concentration of the analytes in the samples viz-a-viz the intensity of colour change in the 182 microplate wells was determined using Stat Fax 4200 microplate reader.

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184 2.7 Statistical Analysis

186 Statistical analysis was performed using GraphPad Prism version 5.03 (San Diego, 187 California, USA). Results were presented as Mean \pm Standard deviation (SD). Inferential 188 statistics using Students' statistical t-test was employed to compare values of the treated 189 rats and control rats. In addition, the One-Way ANOVA (Post Hoc: Tukey's multiple 190 comparative test) was also used to analyse the influence of treatment duration. Statistical 191 significance was set at P=.05.

193 3. RESULTS

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195 3.1 Results of Acute Treatment on Reproductive Hormonal Profile in Rats 196 Administered with Tartrazine 197

Table 1 and table 2 showed hormonal indices in Rats administered with tartrazine 198 199 intraperitoneally and orally respectively. In intraperitoneally treated rats, testosterone 200 (TESTO) in treated male rats showed a significant decrease compared to control from dose 3.33g/kg while Progesterone (PROG) and Estradiol (E2) in female rats showed significant 201 202 increases when compared to control from dose 1.67g/kg at P=.05. More so, in orally treated 203 rats, testosterone (TESTO) showed a significantly lower value in tartrazine treated male rats compared with control male rats from 2.5g/kg dosage while Progesterone (PROG) and 204 205 Estradiol (E₂) in treated female rats indicated a significantly higher value in tartrazine treated female rats compared with control female rats at \tilde{P} =.05. 206

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Table 1. Reproductive Hormonal Profile in Rats Acutely Treated with Tartrazine Intraperitoneally In

Parameters	*TESTO (ng/ml)	**PROG (ng/ml)	**E2 (ng/ml)
0.0g/kg (A _{TIP})	5.75±0.21 [°]	1.10±0.70 ^ª	11.95±2.62
1.67g/kg (В _{тіР})	3.75±1.49 ^{ac}	4.20±0.28 ^{bc}	15.0±2.83 ^{ac}
3.33g/kg (С _{тіР})	4.0±0.28	4.35±0.07	12.65±1.06 ^{ac}
5.0g/kg (D _{TIP})	3.10±0.14	^{bdfg} 7.90±0.99	26.10±11.88

Comment [V1]: Whereevr it is P=.05 replace it with P=0.05

6.67g/kg (Е _{тіР})	2.90±0.42	7.90±0.99	56.25±1.91
8.33g/kg (F _{TIP})	2.15±0.78	7.40±0.28	37.65±1.49
<i>P</i> value	0.03	0.0003	0.0008
F value	5.85	33.37	22.76
Remark	S	S	S

222 223 224 Values in each column with different superscript letter (a, b) differ significantly (P=.05) when comparing the control with other groups. Values in the same column with different superscript letter (c, d) differ significantly (P=.05) when comparing the group B_{TIP} with other groups. Values with different superscript letters (e, f) in the same column are significantly different (P=.05) when comparing group C_{TIP} with other groups. *Male rats, ** female rats. No of female Rats/group = 4 Rats, No of male Rats/group = 4 Rats.

Table 2. Reproductive Hormonal Profile in Rats Acutely Treated with Tartrazine Orally

Parameters	*TESTO (ng/ml)	**PROG (ng/ml)	**E2 (ng/ml)
0.0g/kg (A _{TO})	5.75±0.21 ^ª	1.10±0.70 ^a	11.95±2.62 ^ª
2.5g/kg (В _{то})	3.25±0.07 ^{bc}	7.05±6.45 ^{ac}	37.25±18.6
5.0g/kg (С _{то})	3.35±0.35	^{ade} 2.45±1.85	14.85±3.75
10.0g/kg (D _{то})	3.10±0.71	6.90±0.14	48.95±0.35
15.0g/kg (Е _{то})	^{bdfhi} 1.85±0.21	bcfgh 8.10±2.12	bcegh 29.25±1.20
20.0g/kg (F _{то})	bcehi 2.40±0.28	8.30±0.42	28.60±0.99
P value	0.0005	0.4101	0.020
F value	26.88	1.197	6.210
Remark	S	NS	S
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Values in each column with different superscript letter (a, b) differ significantly (P=.05) when comparing the control group (A_{TO}) with other groups. Values in each column with different superscript letter (c, d) differ significantly (P=.05) when comparing the B_{TO} with other groups. Values in each column with different superscript letter (e, f) differ significantly (P=.05) when comparing the C_{TO} with other groups. Values in the same column with same superscript letter (i) do not differ significantly (P=.05) when comparing the groups E_{TO} and 261 262 263 264 other. *Male rats **Female rats. No of female Rats/group = 4 Rats, No of male Rats/group = 4 Rats

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3.2 Results on Reproductive Hormonal Profile in Rats Chronically Treated 266 with Tartrazine Over a Period of 30 Days 267

269	The comparison of tartrazine male treated Rats and male control Rats showed no significant
270	difference in Testosterone (TESTO) (Table 3). When tartrazine female rats and female
271	control were considered, the comparison showed non-significant differences Progesterone
272	(PROG) and Estradiol (E_2) at <i>P</i> =.05 (table 3).

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274 Table 3. Reproductive Hormonal Profile in Rats Chronically Treated with Tartrazine 275 Over a Period of 30 Days 276

Parameters	Control Rats	Treated Rats	P value	T value	Remark
*TESTO (ng/ml)	4.24±2.21	3.94±1.99	0.6729	0.4259	NS
**PROG (ng/ml)	6.26±1.96	6.25±0.87	0.9950	0.0064	NS
**E2 (ng/ml)	33.30±11.84	37.97±11.07	0.7754	0.2872	NS
NS= Not Significan	t No of control r	ats: Male=18 Fem	ale=22 No	of treated ra	ts: Male = 1

NS= Not Significant. No of control
 Female=22. *Male rats, **Female rats.

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3.3 Results on Reproductive Hormonal Profile in Rats Chronically Treated with Tartrazine Over a Period of 60 Days

283 When tartrazine treated male rats were considered, no significant differences were seen in 284 Testosterone (TESTO) in tartrazine treated male rats compared with control male rats at 285 P=.05 (table 4). In addition, when tartrazine treated female rats were considered, no 286 significant differences were also seen in Progesterone (PROG) and Estradiol (E₂) of 287 tartrazine treated female rats compared with control female rats at P=.05 (table 4). 288

Table 4. Reproductive Hormonal Profile in Rats Chronically Treated with Tartrazine Over a Period of 60 Days Over a Period of 60 Days

Parameters	Control Rats	Treated Rats	P value	T value	Remark
*TESTO (ng/ml)	2.21±1.31	1.92±1.16	0.4361	0.7662	NS
**PROG (ng/ml)	16.32±11.76	13.86±6.45	0.4693	0.7321	NS
**E2 (ng/ml)	61.89±25.29	58.95±23.19	0.7266	0.3526	NS
NS= Not Significan	t. No of control ra	ats: Male=20, Fema	ale=20. No	of treated rate	s: Male = 25,
Female=15. *Male ra	ats, **Female rats.				

3.4 Results on Reproductive Hormonal Profile in Rats Chronically Treated with Tartrazine Over a Period of 90 Days

The comparison of tartrazine treated male rats and control male rats indicated no significant difference in Testosterone (TESTO) concentration of tartrazine treated male rats compared with the control male rats (table 5). When tartrazine treated female rats were considered, no significant differences were seen in Progesterone (PROG) and Estradiol (E_2) concentrations in tartrazine treated female rats compared with control female rats at *P*=.05 (table 5).

Table 5. Reproductive Hormonal <mark>Profile in</mark> Rats Chronically Treated <mark>with Tartrazine Over a Period of</mark> 90 Days

Parameters	Control Rats	Treated Rats	P value	T value	Remark
*TESTO (ng/ml)	3.14±0.98	2.58±1.21	0.1373	1.522	NS

	**PROG (ng/ml)	7.47±3.53	10.80±6.25	0.0516	2.009	NS		
	**E2 (ng/ml)	31.09±19.31	44.94±23.40	0.0514	2.012	NS		
307	NS= Not Significa	nt. No of control	rats: Male=19, F	emale=18. No	of treated ra	ats: Male = 17,		
308	Female=22. *Male r	ats, **Female rats.						
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310			oroductive Hor			hronically		
311	Treated wi	<mark>th Tartrazine</mark> o	over a Period o	f 30, 60 anc	l 90 Days			
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314	treated rats. The			0	,			
315	tartrazine treated male rats from phase 1 to phase 3. When Turkey's multiple comparison							
316	test was used, significant decreases were seen between phase 1 and phase 2 as well as							
317	phase 1 and 3. However, no significant differences were seen between phase 2 and 3. More							
318	so, Table 6 also showed hormonal parameters of female rats treated with tartrazine over a							
319	period of 30, 60 and 90 days. The ANOVA results obtained indicated a significantly higher							
320 321	value in PROG from phase 1 to phase 3. When Turkey's multiple comparison test was used,							
321 322	significant increase was seen between phase 1 and phase 2 as well as phase 1 and 3. However, no significant differences were seen between phase 2 and 3 at P =.05.							
322 323	nowever, no sign		s were seen betw	een phase z	and 5 at P=.0	<i>J</i> J.		
323 324	Table 6. Effect o	f Duration on R	oproductivo Hou	monal Profil	lo in Pote Tr	ootod with		

Tartrazine Over a Period of 30, 60 and 90 Days

Parameters	Phase 1 (Rats)	Phase 2 (Rats)	Phase 3 (Rats)	P value	F value	Remark
*TESTO (ng/ml)		1.92±1.16	2.58±1.21	0.0002	9.786	S
**PROG (ng/ml)	6.25±0.87 ^a	13.86±6.45	10.80±6.25	0.0005	8.712	S
**E2 (ng/ml)	a 37.97±11.07	^{ab} 58.95±23.19	^{ab} 44.94±23.4	0.2397	1.466	NS

Values in the same row with different superscript letter (a, b) differ significantly (*P*=.05) when comparing phase 1 with other phases. Values in the same row with same superscript letter (c) do not differ significantly (*P*=.05) when comparing phase 2 with phase 3. S = Significant. NS=Not Significant.
No of Male rats for Phase 1, 2 & 3 were:17, 25 & 17 respectively. No of Female rats for Phase 1, 2 & 3
were: 22, 15 & 22 respectively. *Male rats, **Female rats.

333 4. DISCUSSION

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334 When hormonal or reproductive hormones were considered in acute toxicity study at high 335 doses, male rats treated (intraperitoneally and orally) indicated significantly lower values in 336 testosterone (TESTO) concentration compared with control rats while female rats treated 337 (intraperitoneally and orally) showed significantly higher values in Progesterone (PROG) and 338 Estradiol (E2) concentrations. The significantly lower value in TESTO observed in tartrazine treated rats supports several findings [9, 20, 23, 27]. Mehedi et al. [9], reported that 2.5% of 339 340 tartrazine administered orally for 13 weeks in male rats induced decreased sperm count, 341 sperm abnormalities viz-a-viz reduction in testosterone concentration compared to control 342 rats. Gautem et al. [20], also reported reduction in sperm density, motility and presence of varying degree of abnormalities in the spermatozoa of rats treated with 0.2g/kg and 0.4g/kg 343 344 bodyweight of tartrazine for 30 days. More so, Sharma et al. [23], reported low levels of 345 steroid hormones including testosterone in rats treated with Kerisi powder (a dye mixture of 346 tartrazine and sunset yellow 6). Khiralla et al. [27], reported that high dose (5 times ADI) of 347 synthetic yellow dye (tartrazine) led to the reduction in testosterone concentration as well as 348 LH concentration. In a related study, Dixit & Goyal [11], reported that the use of an azo dye;

349 indigo Carmine at a dose 39mg/kg bodyweight for 6 weeks induced significant decrease in 350 the concentration of testosterone and the weight of the testes as a result vacuolation of spermatogonia of the testis, flagella distortion of the seminiferous lumen. nuclear 351 352 degeneration (pycnosis), distortion of basement membrane, distortion and loss of leydig cells in indigo Carmine treated male rats. Similarly, Ali et al. [28] reported that administration of 353 354 200mg/kg of tartrazine administered to rats induced decreased superoxide dismutase (SOD) 355 and reduced glutathione (GSH) and increased malondialdehyde (MDA) suggesting oxidative 356 stress induced by tartrazine dyes at high dose. More so, Bousssada et al. [29], reported that 357 sub-chronic treatment of tartrazine (E102) for 30 days at 300mg/kg bodyweight induced 358 altered sperm characteristics and quality accompanied with significantly lowered 359 testosterone concentrations and increased MDA levels in the testicular tissue of tartrazine treated rats. However, our finding when tartrazine was administered at high doses 360 361 contradicts the reports of [12]. Tanaka [12], reported that the administration of synthetic food 362 dyes such as tartrazine at a high dose of 773mg/kg bodyweight in rats did not induce 363 deleterious effect on reproductive hormonal parameters. The reduction in testosterone concentration observed in our study, could be as a result of disruption of the hypothalamus-364 365 pituitary-testes axis regulating testosterone production by the Leydig cells of the testes. The 366 disruption might have resulted from the oxidative insults on the testes arising from azo dye 367 metabolism which might have led to distortion or loss of spermatogenic precursors (spermatogonia) owing to pathologic alteration of the leydig (testosterone production) and 368 sertoli cells (FSH and LH production) architecture. Our present findings further support the 369 370 reports of [28, 29]. Ali et al., [28], reported that administration of 200mg/kg of tartrazine in 371 rats induced decreased superoxide dismutase (SOD) and reduced glutathione (GSH) and 372 increased malondialdehyde (MDA) suggesting oxidative stress induced by tartrazine dyes at high dose. More so, Boussada et al. [29], also reported increased MDA levels in the 373 374 testicular tissue of tartrazine treated rats indicating increased oxidative stress in the testes of 375 rats treated with 300mg/kg of tartrazine. 376

The significantly higher values seen in E_2 in our study when tartrazine was given in high 377 doses collaborates with the report of [30, 31] but contradicts the findings of [23]. Akinloye et 378 379 al.[30], reported in their work that azo dyes induced increase in E2 when azo dyes were fed 380 to rats. Zahra et al. [31], reported that the use of tartrazine and other food dyes such as sunset yellow 6 mimic estrogen in the body and thus stimulates increase in E_2 concentration 381 382 and a reduction in testosterone concentration thereby affecting libido in men. However, Sharma et al. [23], reported reduced E2 in rats treated with Kerisi powder (a dye mixture of 383 384 tartrazine and sunset yellow 6). The increase observed in E_2 in the acute study could be due 385 to xenoestrogenic attributes of tartrazine which is implicated in hormonal imbalance. More 386 so, the significant increase seen in PROG level in the acute treatment (intraperitoneal) also 387 contradicts the finding of [23]. Sharma et al. [23], also reported reduced PROG in rats 388 treated with Kerisi powder (a dye mixture of tartrazine and sunset yellow 6) as a result of 389 apoptosis of ovarian cells being induced by the synthetic dye. 390

Furthermore, when chronic treatments were considered, hormonal parameters after 30 days, 391 392 60 days and 90 days chronic treatment showed non-significant reductions in testosterone 393 (TESTO) concentration in the tartrazine treated male rats compared with their respective 394 control male rats. The non-significant difference seen in testosterone when given ADI doses 395 compared with control group is in line with the findings of [12, 13, 14]. Tanaka [12], Elhkim et 396 al. [13], EFSA [14], reported in their separate work that tartrazine did not induce any 397 deleterious effect on reproductive hormonal parameters in rats when given at ADI doses. 398 More so, Gil [32], also reported that synthetic dye such as tartrazine in a concentration of 399 1mM in a cell culture medium did not induce a decrease in testosterone concentration 400 compared to vehicle treated control. However, oxidative stress in the adrenal cortex affecting 401 steroid hormones production when exposed to high concentration of azo dyes was reported.

The non-significant decreases seen in testosterone concentration in the chronically treated rats compared with the control rats over a period of 30, 60 and 90 days could be as a result of the testes or the body system not been overwhelmed by the vehement effects of reactive oxygen species (ROS) produced by the tartrazine during metabolism.

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407 In addition, Progesterone (PROG) and Estradiol (E₂) concentrations after 30, 60 and 90 days 408 chronic treatment showed non-significant increases in tartrazine treated female rats 409 compared with their respective control female rats. The non-significant differences observed 410 in PROG and E₂ concentration concurs with the findings of [12, 13, 14] but contrast the reports of [23, 30, 31], when the effects of azo dyes on reproductive parameters were 411 evaluated. Tanaka [12], Elhkim et al. [13], EFSA [14], recorded in their separate work that 412 ADI doses of tartrazine did not cause harmful effect on reproductive hormonal parameters. 413 However, Sharma, [23], reported a fall in E2 and PROG levels due to vacuolation or 414 415 apoptosis of ovarian tissues when tartrazine mixed with sunset yellow 6 were fed to rats. In addition, Akinloye et al. [30] and Zehra et al. [31], reported in their separate studies that 416 xenoestrogenic activities of tartrazine azo food dyes induced low levels of progesterone in 417 418 plasma. The non-significant differences observed in E₂ and PROG in the chronic treatment 419 could be as a result of complete removal of reactive oxygen species by the body antioxidative mechanism when these dyes are administered at ADI doses. It is quite possible 420 that the derangements caused by synthetic food dyes are mainly due overwhelming 421 tendencies of the anti-oxidative capacity of the body system when these days are consumed 422 423 in high doses.

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425 Finally, when the comparative analyses of hormonal parameters of tartrazine treated rats were considered using One-Way ANOVA over the period of 30, 60 and 90 days, TESTO 426 427 concentration indicated significantly lower levels in tartrazine treated male rats. Significant 428 decreases were seen between 30 days and 60 days as well as between 30 days and 90 429 days. However, no significant difference was seen between 60 days and 90 days. When treated female were considered, PROG showed significantly higher values over 30, 60 and 430 90 days in tartrazine treated female rats. Significant increase was seen between 30 days 431 and 60 days as well as between 30 days and 90 days. However, no significant differences 432 433 were seen between 60 days and 90 days.

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The significantly lower value seen in TESTO over the period of 30, 60 and 90 days suggest 435 436 gradual fall in the TESTO level over time and possibly alteration in membrane structure of parenchymal cells of the leydig and sertoli cells (of the testis) disrupting the hypothalamic-437 438 pituitary-testes axis. In a related study, Helal et al. [33], reported that administration of food 439 additives such as sodium nitrate and monosodium glutamate at recommended dose induced 440 reduction in testosterone concentration. Also, Mahmoud et al. [34], reported reduction in 441 spermatogenesis when azo dye brilliant black was given to male rats orally at a dose of 0.08g/kg and 0.4g/kg over a period of 30 days. In addition, the significantly higher levels of 442 443 PROG seen in our study is contrary to the reports of [23, 32]. Sharma et al. [23], reported severe degeneration of corpus luteum of the ovaries in tartrazine treated female rats which 444 445 was attributed to reduction in plasma PROG level. More so, Gil [32], also reported no 446 significant difference in E₂ when tartrazine treated cells were compared with vehicle treated 447 control in a cell culture medium at a concentration of 1mM. The significant increase seen in 448 PROG could be related to hormonal imbalance induced by distortion of the follicular cells, 449 theca interna and externa (of the ovaries) due to persistent oxidative stress induced by azo 450 dyes. However, no significant increase was seen E_2 concentration over 30, 60 and 90 days. 451 Our finding collaborates with the report of [12, 21] but contrary to the reports of [23, 30]. 452 Tanaka [12], reported that tartrazine at a dose of 773mg/kg did not affect reproductive 453 hormonal parameters when tested in rats. More so, Meyer et al. [21], recorded that 454 administration of 0.5mg/kg and 50mg/kg of tartrazine did not induce significant change in the 455 weight of the ovaries viz-a-viz estradiol concentration. However, Akinloye et al. [30], reported 456 in their work that azo dyes such as tartrazine possesses xenoestrogenic attributes and are 457 therefore stimulates increase in E₂ which is implicated in hormonal imbalance. Sharma et al, 458 [23], further reported reduced E_2 levels in rats treated with tartrazine which they attributed to apoptosis of ovarian cells. The non-significant difference observed in E2 over the period of 459 460 30, 60 and 90 days in the treated female rats could be as a result of intact membrane structure of parenchymal (follicular and luteal) cells of the ovaries maintaining the 461 462 hypothalamic-pituitary-ovaries axis as well as poor xenoestrogenic activities of tartrazine on 463 the ovarian cells associated with the ADI doses given over time.

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465 5. CONCLUSION

466 In the acute toxicity study, reduction in TESTO concentration as well as increase in PROG 467 and E₂ was seen which suggest possible disturbance in the fertility profile or capacity when 468 these dyes are consumed in high doses. However, in chronic study, significant differences were not seen in TESTO concentration as well as increase in PROG and E2 was seen which 469 470 suggest possible disturbance in the fertility profile or capacity when these dyes are not 471 consumed in high doses. Finally, when the influence of duration of exposure at ADI doses were considered over 30, 60 and 90 days, gradual reduction in TESTO concentration and 472 473 increase in PROG concentration was seen. This implies that there could be possibility of 474 hormonal derangements when food dyes are consumed even at ADI doses on daily basis 475 over prolonged period.

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478 6. RECOMMENDATION

479 It is therefore recommended that high doses of tartrazine in foods or food products should be 480 avoided. Also, the ADI dose of tartrazine should be reconsidered by international and 481 national agencies on reducing the ADI dosage over a life time. More so, 482 marketers/consumers should be sensitized/educated/re-educated on the use of food dyes 483 and finally, governmental policies/consumer protection agency should regulate and ensure 484 appropriate labelling of food dyes and food products with additives.

486 COMPETING INTERESTS487

Authors have declared that no competing interests exist.

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491 CONSENT

Not applicable

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496 ETHICAL APPROVAL497

We hereby declare that Principles of laboratory animal care (NIH publication No. 85-23,
revised 1985) were followed, as well as specific national laws where applicable. All
experiments have been examined and approved by the Rivers State University
research/ethics committee.

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606 **ABBREVIATIONS**

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608	TESTO	=	TESTOSTERONE
609	PROG	=	PROGESTERONE
610	E ₂	=	ESTRADIOL
611	ADI	=	ACCEPTABLE DAILY INTAKE
612	PHASE 1	=	30 DAYS
613	PHASE 2	=	60 DAYS
614	PHASE 3	=	90 DAYS