

1 **PILOT STUDY**

2

3 **Preliminary investigation of the effects of long-term dietary intake of genistein and**
4 **daidzein on hepatic histopathology and biochemistry in domestic cats (*Felis catus*)**

5

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14 **Short title:** Feline hepatology following isoflavone exposure

15 **Summary**

16 Dietary isoflavones have been hypothesised to play a role in hepatic veno-occlusive disease
17 in captive exotic felids, although empirical evidence is lacking. This study aimed to
18 investigate the effect of long-term (>1 year) dietary genistein and daidzein exposure on the
19 hepatic biochemistry and histology of domestic cats. Individual cats were assessed for hepatic
20 enzyme and bile acid production before and after the removal of isoflavones from their diet in
21 the treatment group (n=4), and at the same times in unexposed control animals (n=7). No

22 significant differences were detectable in hepatic biochemistry between treatment and control
23 groups, and all serum values were within the normal reference ranges for domestic cats.
24 Additionally, treatment animals demonstrated slightly greater areas of fibrosis surrounding
25 hepatic venules than control animals, but this difference was not statistically significant. On
26 the basis of the results presented, dietary isoflavones, at the current dose and duration of
27 exposure do not appear to modulate hepatic enzyme production or histological parameters.

28 **Keywords:** cats, feline, soya, isoflavone, liver

29 **Introduction**

30 Dietary isoflavones, such as genistein and daidzein, have previously been shown to elicit a
31 diverse array of physiological effects including endocrinological, morphological and
32 histological changes in a variety of tissues or organs in species such as rodents, pigs and
33 humans (Barnes *et al* 2000; Ford *et al* 2006; McClain *et al* 2006). Isoflavones are structurally
34 similar to oestradiol and can bind to oestrogen receptors and function as natural selective
35 oestrogen receptor modulators, although a diverse array of non-hormonal effects and tissue-
36 or species-specific effects have also been observed in both rodents and humans (Hollander
37 1997; Barnes *et al* 2000; Pike 2006).

38 The liver is also a target of oestrogen activity (Diel *et al.* 2002), and isoflavones have been
39 shown to exert a variety of effects on hepatic activities. Both genistein and daidzein have
40 been associated with hypertrophic effects in the liver (Banz *et al.*, 2004; McClain *et al.*,
41 2006b). However, studies with soy protein isolate (containing isoflavones) have produced
42 divergent results, with no effect in female rats and reduced liver weights in male rats (Peluso
43 *et al.*, 2000; Huang *et al.*, 2005; Tachibana *et al.*, 2005). Likewise, mild hepatotoxicity was
44 only demonstrated following exposure to high isoflavone doses (500mg/kg BW) with these

45 changes reversible, suggesting that normal dietary exposure (estimated to be < 10 mg/kg BW
46 for domestic cats and captive cheetahs; Bell *et al.*, 2006 and 2010) is unlikely to pose a risk to
47 hepatic health (McClain *et al.*, 2006b). Moreover, other studies have demonstrated a
48 protective role for isoflavones against various hepatic insults (Lee *et al.* 1995; Kang *et al.*
49 2001; Liu *et al.* 2002; Kuzu *et al.* 2007; Wong *et al.* 2007).

50 However, dietary isoflavones have been implicated in the onset or progression of veno-
51 occlusive disease (VOD; hepatic fibrosis) in captive cheetahs (Setchell *et al.*, 1987ab;
52 Gosselin *et al.*, 1988). This disease is responsible for significant levels of mortality in the
53 global captive cheetah population (Munson *et al.*, 2005), but the cause(s) are not yet clearly
54 defined. Hepatic architecture is modulated during VOD and histological changes include
55 hepatic congestion, haemorrhage, hepatocyte and hepatic stellate cell vacuolation, foci of
56 extra-medullary haematopoiesis (EMH; a marker for hypoxia, infection and/or precocious
57 immune response) (Törő *et al.* 2007), and increased neutrophil and macrophage cell numbers.
58 However, no controlled study has been conducted to determine the ability of isoflavones to
59 modulate hepatic parameters in any felid species. Therefore, the aim of this study was to
60 determine the potential of long term consumption of genistein and daidzein to elicit
61 detectable effects on hepatic histology or biochemical parameters (as an indication of
62 hepatocyte health and biliary secretion) in a felid species, the domestic cat.

63 **Materials and Methods**

64 Eleven female short-haired domestic cats were utilised in this study. Premature removal of
65 four cats from the treatment group (for reasons unrelated to this study) resulted in a low and
66 uneven sample size. Cats were bred and maintained at the Centre for Feline Nutrition
67 (Massey University, New Zealand). Kittens were mother-reared until six weeks of age,
68 during which time they had access to the queen's diet (a commercial diet which met the

69 AAFCO (2004) standards for gestation and lactation). Previous analysis of this diet
70 demonstrated it contained a very low (16 µg total isoflavone/g DM) isoflavone
71 concentrations (Bell *et al* 2006), and exposed the queens to a total daily isoflavone dose of
72 approximately 0.56 mg – 0.84 mg/kg BW. Intake of the maternal diet by kittens prior to
73 weaning was thought to be minimal, but accurate assessment of the intake was not possible
74 due to co-housing with the queen. However, pre-weaning exposure was identical between
75 control and isoflavone-treated animals and intake predicted to be equivalent. At six weeks of
76 age, cats were gradually weaned from the queens and separated into treatment and control
77 groups. Cats were gradually introduced to the trial diets, and the day of sole consumption of
78 the trial diet was recorded as the start of the trial for each individual cat. Cats were group-
79 housed (maximum nine per pen) in multi-level outdoor pens (approx. 5 m x 2 m x 3 m),
80 exposed to natural day/night cycles and provided with daily exercise opportunities and
81 environmental enrichment. Control and treatment groups were housed in adjacent pens. Each
82 cat was weighed weekly and body weight was recorded and tracked against the colony
83 average. Cats had consumed the trial diets for an average of 394 (\pm 25.73) days at the time of
84 blood collection.

85 Cats assigned to the treatment group (n = 4 at study completion) had been exposed to the
86 dietary isoflavones, genistein and daidzein, since weaning. Cats in the control group (n = 7)
87 consumed the same base diet as the treatment group, without the addition of isoflavones.

88 The base diet for both control and treatment groups for the duration of the trial was a moist
89 feline diet which met the requirements for growth in the domestic cat according to AAFCO
90 (2009) testing. Previous analysis of this diet demonstrated it contained no detectable
91 isoflavones (Bell *et al* 2006). The purified (99.9%) form of each isoflavone, genistein
92 (150µg/g DM) and daidzein (150µg/g DM) (LC Laboratories, MA, USA) was added to the

93 base diet of the treatment group, to provide a calculated dose of 300 µg total isoflavone/g
94 DM, which is representative of exposure through consumption of certain commercially
95 prepared feline diets (Bell *et al* 2006). Due to the small quantities of powder to be added to
96 large quantities of base diet it was necessary to use a freeze-dried inert carrier. The same
97 concentration of freeze-dried carrier was added to the control diet without the addition of
98 isoflavone powders.

99 Cats were provided water *ad libitum* during the trial, and offered enough food to provide each
100 cat with appropriate energy intake for its age (i.e. 200 kcal/kg BW/d at 10 weeks, gradually
101 reducing to 88 kcal/kg BW/d by 40 weeks; Legrand-Defretin and Munday 1993). Food intake
102 per group was accurately weighed on a daily basis. Monthly assessments were made of
103 individual food intake by separation of each cat into individual metabolism cages for 24 h,
104 during which time food intake and urinary and faecal output were recorded.

105 Sub-samples of the control and treatment diets were assayed for isoflavone content
106 intermittently throughout the trial, according to methodology described in Bell *et al* (2006).
107 Ethical approval (MUAEC Protocol No. 06/06) for this trial was obtained from the Massey
108 University Animal Ethics Committee (2006).

109 The average age at the time of initial blood collection was 428 (\pm 25.75) days. Cats were
110 fasted overnight, prior to an initial (2 ml) blood sample being collected by venipuncture of
111 the jugular vein. The cats were then offered a meal of basal diet, and a second (1 ml) blood
112 sample withdrawn by venipuncture, 2 h after ingestion of this meal. Blood was transferred
113 into vacutainers and centrifuged to collect serum. Following collection of initial blood
114 samples, the diet of the treatment group was replaced with the control diet (devoid of
115 isoflavones), while the cats in the control group continued to be maintained on the control

116 diet. Forty days following this dietary change, a second pair of blood samples were collected
117 and analysed, as described above.

118 Serum from the pre-prandial blood sample was analysed for alkaline phosphate (ALP),
119 aspartate aminotransferase (AST), alanine aminotransferase (ALT), gamma glutamyl
120 transferase (GGT) and bile acids (Gribbles Pathology Ltd., Palmerston North, NZ). Serum
121 from the post-prandial blood sample was analysed for bile acid concentration only (Gribbles
122 Pathology Ltd., Palmerston North, NZ). Serum was analysed within 48 h of collection.

123 A liver biopsy was obtained from cats that had been exposed to dietary isoflavones for
124 approximately 394 days, when these animals underwent routine gonadectomy. Briefly,
125 general anaesthesia was induced with Zoletil 100 (tiletamine and zolazepam 500 mg/ml each;
126 12 mg/kg BW, sub-cutaneously) (Virbac, Auckland, New Zealand) and maintained with
127 halothane/oxygen delivered *via* an endotracheal tube. A midline incision was made in order
128 to perform routine ovario-hysterectomy. Upon completion of this procedure, a distal liver
129 lobe was located and a wedge biopsy (0.7 – 1 g) taken from its border. One or two catgut
130 sutures were used to control haemorrhage of the liver parenchyma (Cole *et al.* 2002), before
131 routine abdominal closure. All animals received Temgesic (324 µg/ml buprenorphine
132 hydrochloride, 0.03 mg/kg BW, sub-cutaneously) (Reckitt Benckiser, Auckland, New
133 Zealand) for pain relief after surgery, and Ketofen (ketobrofen, 1 mg/kg BW *per os*) for the
134 next 48 h. Cats were maintained in individual metabolism cages for 14 days following
135 surgery, after which time sutures were removed and cats were returned to normal housing.

136 Each liver biopsy was immediately weighed and transferred to 10% neutral-buffered formalin
137 (NBF), before processing on a Leica[®] TP1050 Tissue Processor (Global Science and
138 Technology, Auckland, NZ). The samples were dehydrated through a series of graded
139 alcohols (70%, 95% and absolute ethanol, BD, Poole, UK) at ambient temperature, cleared in

140 xylene and impregnated with Paraplast[®] Tissue embedding Medium (Global Science and
141 Technology, Auckland, NZ) under pressure at 60°C. The samples were then embedded using
142 a Leica Histo Embedder (Global Science and Technology, Auckland, NZ) and 3 µm sections
143 were cut using MicroTec[®] Rotary Microtome (Global Science and Technology, Auckland,
144 NZ). The sections were floated onto a Thermo[®] Tissue Bath (Medica Pacifica, Auckland,
145 NZ) at 43°C and mounted onto Superfrost, pre-cleaned slides. Half of the slides were then
146 placed in a 60°C oven for 20 min then stained with haematoxylin and eosin (H&E) on a
147 Leica[®] Autostainer XL (Global Science and Technology, Auckland, NZ).

148 The remaining slides were placed in a 60°C oven for 20 min then dewaxed on a Leica[®]
149 Autostainer XL (Global Science and Technology, Auckland, NZ) before staining using the
150 Masson's Trichrome method. Following hydration in water, slides were left to mordant in
151 Bouin's solution (Merck, Palmerston North, NZ), overnight at room temperature. Slides were
152 then washed in tap water, stained in Celestine Blue (Merck, Palmerston North, NZ) for 10
153 min, rinsed again before staining in Mayer's Haematoxylin (Merck, Palmerston North, NZ)
154 for 10 min. Slides were rinsed again for 4 min and then stained in Beibrich Scarlet-Acid
155 Fuchsin (Merck, Palmerston North, NZ) for 2 min before further rinsing. Sections were
156 covered in 5% Phosphotungstic Acid (Merck, Palmerston North, NZ) for 15 min and then
157 rinsed prior to staining with Light Green Solution (Merck, Palmerston North, NZ). After
158 further rinsing, sections were blotted dry with filter paper and placed in 1% Glacial Acetic
159 Acid (BD, Poole, UK). Sections were then blotted dry again before dehydrating in 95%
160 ethanol, absolute ethanol, and finally clearing in xylene before mounting.

161 Liver sections were examined by a veterinary histo-pathologist (W. Roe) who provided a
162 detailed report of the sections from each cat. The histo-pathologist was blinded to the
163 treatment groups. Parameters reported were haemosiderin accumulation, intra-hepatocyte

164 vacuolation, hepatocyte degeneration, necrosis or regeneration. The presence/absence and
165 extent of histological parameters were then tabulated and averaged according to treatment
166 group.

167 The extent of fibrous tissue development around three hepatic blood venules (see Figure 1)
168 was measured using ImageJ software (version 1.38. Rasband 2007; Research Services
169 Branch, National Institute of Mental Health, MD, USA) and expressed as the percentage of
170 the total area of each blood venule.



171

172 Figure 1. Liver section from a domestic cat in the current study. Central vein surrounded by
173 subendothelial fibrosis (stained green/blue). Massons Trichrome stain

174 Changes in the serum concentrations of enzymes over time were calculated for each cat, and
175 groups were tested for significant differences in any temporal changes. Differences in
176 biochemical parameters at the first sampling time (prior to removal of isoflavones from the
177 treatment group's diet), were also tested between groups. Residual data was tested for
178 normality using the Anderson-Darling test. Differences between groups were tested for
179 significance using the Mann-Whitney test as data was found to be not normally distributed.

180 Differences between the incidence of congestion, vacuolation, extra-medullary
181 haematopoiesis (EMH) and inflammatory cells in treatment and control groups were tested
182 using Fisher's exact test. The median is reported instead of the mean (Glantz, 2005). All
183 statistical analyses were performed with Minitab software (version 15, Minitab Inc., PA,
184 USA).

185 **Results and Discussion**

186 No significant differences ($P > 0.05$) were detected in any biochemical parameters (ALP,
187 AST, ALT, GGT, fasted or fed bile acids) within the first sampling phase, prior to isoflavone
188 removal from the diet of the treatment cats (Table 1). Overall, changes in hepatic
189 biochemistry parameters were generally similar between control and treatment cats.
190 However, the results of a power analysis suggests that subtle differences were undetectable in
191 the study, and that only large differences would be considered significant with the available
192 sample size (a minimum difference of 59.6% with a power of 95%, or a minimum difference
193 of 46.1% at 80% power).

194 Changes in these parameters, as well as bile acid response to a meal within each cat (before
195 and after dietary change) did not differ between groups, or within the treatment group (i.e.
196 during isoflavone exposure compared to following removal of isoflavones from the diet; $P >$
197 0.05) (Table 2). All parameters were within normal reference ranges for domestic cats, at all
198 time points.

199 The consistent increase in ALT and GGT production between first and second sampling,
200 observed in both control and treatment groups (Table 2), may reflect altered metabolism or
201 hepatic activity as a consequence of removal of the reproductive tract.

202

203 Table 1. Median (lower quartile, upper quartile) hepatic biochemistry parameters following a
 204 394 day (± 25.73) period of dietary isoflavone exposure in the treatment group

	ALP (U/L)	AST (U/L)	ALT (U/L)	GGT (U/L)	Pre- prandial bile acids ($\mu\text{mol/L}$)	Post- prandial bile acids ($\mu\text{mol/L}$)	Bile acid response to a meal (U/L)
Control	29 (25, 31)	23 (18, 27)	43 (36, 52)	1.0 (0, 2.0)	0.5 (0.3, 0.5)	1.5 (1, 2.2)	0.9 (0.6, 1.7)
Treatment	56 (33, 82)	25 (18, 35)	45 (33, 56)	0.5 (0, 1.3)	0.4 (0.3, 0.6)	2.3 (2.0, 2.8)	1.7 (1.4, 2.3)

205 The increases observed in plasma ALT and GGT may have occurred from hepatocellular
 206 injury, hormonal action, or as a consequence of muscle injury (Roth-Johnson 2004; Webster
 207 2005), all of which are possible mechanisms in these cases. Gonadectomy would have been
 208 associated with a reduction in circulating oestrogen, whilst the muscle trauma resulting from
 209 abdominal opening during gonadectomy may have elicited the increased ALT and GGT
 210 production. However, the lack of significance between changes in the control and treatment
 211 groups ALT and GGT levels suggests that the hepatic production of these enzymes was not
 212 modulated in response to dietary isoflavones, either during exposure or following a 40-day
 213 recovery period.

214 This is in contrast to previous findings in cheetahs, in which removal of dietary isoflavones
 215 elicited a reduction in ALT three months later (Setchell *et al.* 1987a). However, this earlier
 216 study did not control for the variable nutrient composition of isoflavone-containing and

217 isoflavone-free diets and as such, the alteration in ALT cannot be apportioned solely to
 218 isoflavones.

219

220 Table 2. Median (lower quartile, upper quartile) change in hepatic biochemistry parameters
 221 after a 40 days period following the removal of isoflavones from the treatment group cats (no
 222 dietary change in the control cats).

	ALP (U/L)	AST (U/L)	ALT (U/L)	GGT (U/L)	Pre- prandial bile acids ($\mu\text{mol/L}$)	Post Prandial Bile Acids ($\mu\text{mol/L}$)	Bile acid response to a meal (U/L)
Control	13 (10, 19)	-4.0 (-12, 0.5)	38 (30, 44)	2.0 (1.5, 2.5)	0.5 (0.2, 1.6)	-0.1 (-0.6, 1.2)	0.0 (-2.7, 0.7)
Treatment	-7.5 (-18, 2.5)	6.5 (-6.3, 16)	33 (19, 48)	4.5 (2.3, 12)	0.1 (-0.1, 0.1)	0.4 (0.1, 0.7)	0.5 (0.2, 0.8)

223

224 However, the current study utilised a shorter recovery period than Setchell *et al.* (1987a) and
 225 gonadectomy occurred at the time that isoflavone exposure ceased, both of which may
 226 explain the disparate results. Likewise, it is possible that the dose used in this study was
 227 insufficient to elicit any change in these enzymes, since GGT was only slightly elevated in
 228 rats when exposed to much higher genistein doses of 500 mg/kg BW (McClain *et al.* 2006b).
 229 Additionally, the sensitivity of ALT for detection of hepatic disease is moderate at best
 230 (Jacob *et al.* 2002), and only mild elevation of ALT or AST levels were noted in a domestic
 231 cat diagnosed with hepatic VOD (Cave *et al.* 2002).

232 Rats exhibited a mild increase in plasma ALP after chronic exposure to 500 mg genistein/kg
233 BW (McClain *et al.* 2006b). In the current study, no statistically significant difference in ALP
234 concentrations were detected, although the treatment group's median ALP appeared to be
235 higher than the control group's prior to removal of the isoflavones from their diet. Likewise,
236 no significant changes in ALP were detected within individuals, but values suggest a
237 reduction in ALP in treated animals following the removal of isoflavones from their diet,
238 whilst control cats exhibited an increase in median ALP levels. Following a four week
239 recovery period, rats exhibited significant recovery in the modulation of ALP (McClain *et al.*
240 2006b), suggesting that isoflavones may have elicited a non-permanent increase in ALP
241 production in cats in this study. However, this was not statistically determined due to the low
242 sample size. In the cat, the combination of relatively low hepatic stores and the short half-life
243 mean that plasma ALP is an insensitive marker of hepatic injury (Hoffmann *et al.* 1977).
244 Thus, the absence of a significant change in serum ALP does not exclude modulation of
245 hepatic injury.

246 The lack of difference between groups during the exposure phase, and the observed increase
247 in ALT following cessation of isoflavone exposure indicates that hepatic injury was not
248 associated with isoflavone consumption. Unchanged AST concentrations provided further
249 evidence to support a lack of hepatic insult, since elevations of this enzyme, in combination
250 with increased ALT levels are generally good indicators of hepatic dysfunction in the cat
251 (Roth-Johnson 2004; Webster 2005). Rats exposed to significantly higher doses of genistein
252 also failed to exhibit any change in AST concentration (McClain *et al.* 2006b).

253 Serum bile acid concentration is used in veterinary medicine to assess hepatic clearance from
254 portal circulation and functional hepatic mass (Roth-Johnson 2004; Webster 2005). Fasted
255 bile acid concentrations may offer greater specificity for detecting hepato-biliary disease than

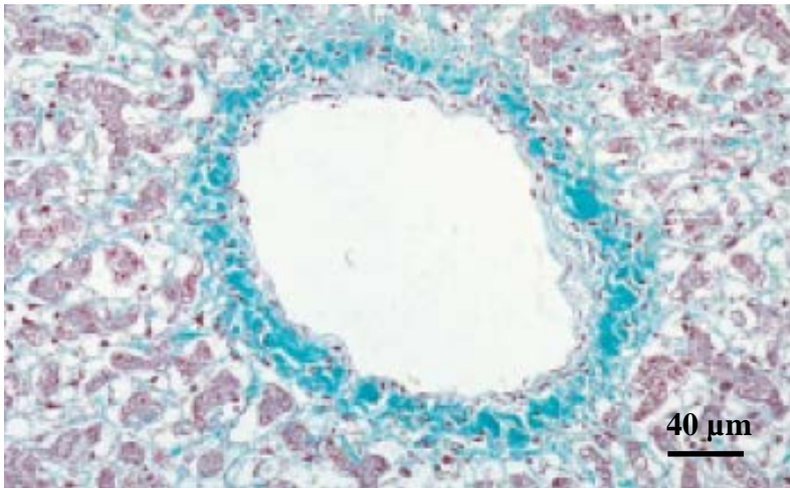
256 ALP, ALT or GGT. However, bile acid levels are poorly correlated with histological
257 findings, and may be elevated in cases of intestinal disease (Roth-Johnson 2004; Webster
258 2005). Inter-individual differences in gall bladder emptying, gastric emptying rate, intestinal
259 transit rate, ileal bile acid resorption and gut microflora are all known to affect entero-hepatic
260 recirculation of bile acids (Webster 2005). Moreover pre- and post-prandial bile acid
261 concentrations in a domestic cat known to be suffering from hepatic VOD were within the
262 normal reference range for this species (Cave *et al.* 2002). Given the variability of this
263 parameter and the small sample size in this study, it is unsurprising that no difference in bile
264 acid production was detectable between the two groups. In light of the other parameters
265 concurrently evaluated, our findings indicated no gross liver dysfunction or clinically
266 significant effects following the consumption of dietary isoflavones under the conditions of
267 this experiment.

268 Hepatic adaptation of enzyme production following chronic exposure to isoflavones may
269 have occurred in this study, and between-group differences may have been more apparent
270 following acute exposure. However, acute exposure was not measured in the current study
271 since cats were exposed to dietary isoflavones at weaning which rendered it impossible to
272 evaluate acute pre- and post-isoflavone exposure responses.

273 There were few histological abnormalities in any of the liver sections. A lack of haemosiderin
274 accumulation indicated that any observed congestion and periportal haemorrhage was a
275 recent occurrence. Intra-hepatocyte vacuolation was not significant, and any inter-individual
276 variation in vacuolation was thought to represent divergent glycogen accumulation and
277 reflective of differences in body condition and/or differences in fasting time.

278 No evidence of hepatocyte degeneration, necrosis or regeneration was observed. One
279 treatment cat had low numbers of neutrophils around some periportal areas, but these were

280 considered unlikely to be significant. The mean area of fibrous tissue surrounding hepatic
281 blood venules (Fig. 1b) in control cats represented 28.51% (\pm 2.60%; range 14.78 – 40.05) of
282 the venule area, and that of treatment cats was not significantly different ($32.84 \pm 4.18\%$;
283 range 20.72 – 51.16, $P > 0.05$). A liver section from a domestic cat suffering from hepatic
284 VOD was also measured from a published photomicrograph (Cave *et al.*, 2002) and found to
285 exhibit fibrosis covering an area equivalent to 46.75% of the venule lumen area (Figure 2).



286

287 Figure 2. Liver section from a domestic cat with diagnosed hepatic veno-occlusive disease
288 (VOD) (from Cave *et al.* 2002). Central vein surrounded by subendothelial fibrosis (stained
289 dark green/blue). Masson Trichrome stain.

290 Over half (57% or 4 of 7) of the control cat sections, but all (100%: 4 of 4) of the treatment
291 group cat sections demonstrated congestion, however this difference was not significant ($P >$
292 0.05). No significant difference in the incidence of vacuolation of hepatocytes and Ito cells
293 was found between control cat sections (86%: 6 of 7) or treatment cat sections (50%: 2 of 4)
294 ($P > 0.05$). Extra-medullary haematopoiesis was detected in 14% (1 of 7) of control cat
295 sections and 25% (1 of 4) of treatment cat sections, but no difference was detectable ($P >$
296 0.05). Neutrophils and/or macrophages were not observed in any control group sections (0 of
297 7), but were seen in 50% (2 of 4) of treatment group sections. However, this difference also

298 failed to achieve statistical significance ($P > 0.05$). A summary of the histological findings is
299 provided in Table 3.

300 The lack of histological changes detected here is in agreement with the biochemistry results.
301 Sinusoidal and haemorrhagic congestion with perivenular fibrosis are typical histological
302 signs of VOD in both cheetahs (see Figure 3) and the domestic cat (see Figure 2) (Setchell *et*
303 *al.* 1987a; Cave *et al.* 2002).

Table 3. Summary of mean hepatic parameter scores in the liver biopsies of control (N = 7) and treatment cats (N = 4) at the time of ovario-hysterectomy

	Control Group	Treatment Group
Congestion	57%	100%
Vacuolation	86%	50%
Extra-medullary haematopoiesis	14%	25%
Neutrophils/macrophage infiltration	0%	50%
Fibrous area	28.9%	32.8%

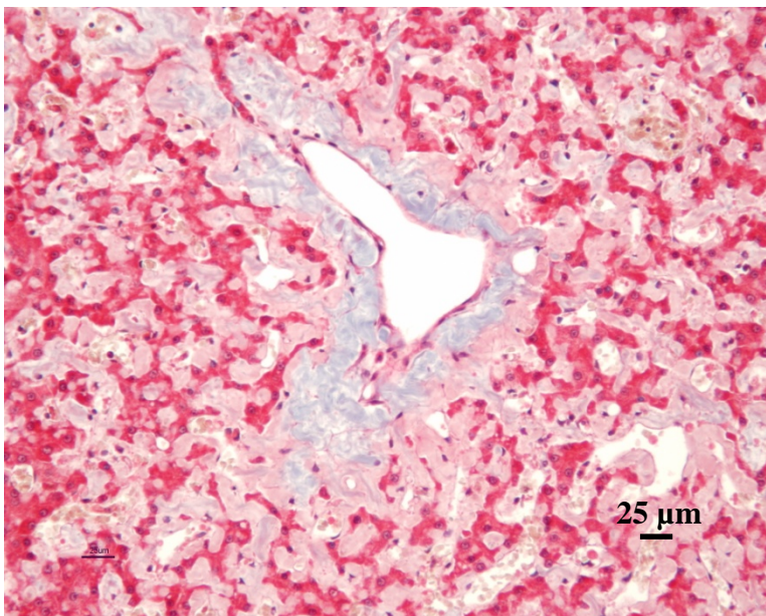
304 N.B. Data is presented as the percentage of cats with positive observation scores for each
305 parameter. The exception to this is fibrous area which is presented as the mean area of fibrous
306 tissue surrounding hepatic blood venules as a percentage of the venule area.

307

308 However, no evidence of hepatic congestion, vacuolation, EMH or inflammatory cell
309 infiltration was detected in the domestic cats following extended exposure to dietary
310 isoflavones. Although a greater proportion of cats in the treatment group, compared to the
311 control group, demonstrated hepatic congestion (100% *versus* 57%, respectively), this failed

312 to achieve statistical significance, and was most likely due to anaesthesia and surgical
313 procedures, rather than any underlying hepatic disease.

314 Similarly, no evidence of hepatic fibrosis or pathology was detected in any cat, regardless of
315 treatment group. The extent of hepatic fibrous tissue formation around blood venules did not
316 differ between groups, which suggested that these compounds are unlikely to play an
317 aetiological role in the VOD. However, hepatic fibrosis is a dynamic process, involving
318 nonspecific mechanisms which respond to inflammation and/or hepatic injury (Center, 2004).
319 Additionally, changes in liver architecture were primarily due to the deposition of
320 extracellular matrix which operates to reduce perfusion and stimulate sinusoid capillarisation
321 and collagenisation (Center 2004). The chronic nature of the appearance of these effects
322 indicates that differences between treatment and control animals in the current study may not
323 have become detectable until much later in life.



324
325 Figure 3. Liver section from a cheetah with veno-occlusive disease (Courtesy Wellington
326 Zoo, New Zealand) - central vein surrounded by subendothelial fibrosis (stained blue using
327 Masson's Trichrome stain)

328 Furthermore, unlike cheetahs in which the disease is relatively common, hepatic VOD has
329 only been reported in one domestic cat (Cave *et al.* 2002). It appears likely that the domestic
330 cat and cheetah differ in their susceptibility to VOD or the biological action of isoflavones, or
331 other environmental factors may be responsible for the incidence of VOD in captive cheetahs.
332 Budd-Chiari-like syndrome is a rare condition which has only been reported in two domestic
333 cats, and is typified by hepatic venous outflow obstruction (Cave *et al.* 2002), potentially
334 related to VOD. Elevated levels of tumour necrosis factor-alpha (TNF- α) and Interleukin-1 β
335 are observed in association with this syndrome, although it is unclear as to whether they play
336 an aetiological or responsive role (Cave *et al.* 2002). Interestingly, the ability of isoflavones
337 to inhibit TNF- α (Kang *et al.* 2005), suggests that dietary isoflavone intake is more likely to
338 reduce, rather than increase, the risk of hepatic fibrosis and VOD in domestic cats. Such a
339 protective mechanism has been postulated in other studies (Kang *et al.* 2001; Liu *et al.* 2002).
340 Although the current study was not designed to assess cellular proliferation or hepatic
341 toxicity, the lack of difference between control and treatment animals indicates that neither
342 beneficial nor detrimental effects were elicited in the liver following isoflavone exposure,
343 under the conditions of this trial.

344 **Conclusions**

345 The influence of dietary isoflavones, genistein and daidzein, on hepatic biochemistry and
346 histology in the domestic cat was investigated here for the first time. The purified aglycones
347 of genistein and daidzein, at the dose and duration of exposure utilised here, do not appear to
348 modulate hepatic enzyme production or histological parameters in the domestic cat.
349 Modulation of biochemical parameters was minor if present at all, and failed to achieve
350 statistical significance or exceed normal reference ranges for clinically healthy cats.
351 However, caution is warranted in extrapolating these findings to felids exposed to soy-

352 derived isoflavone glycosides or other phytoestrogen compounds. Although larger sample
353 sizes are needed to confirm our findings, dietary isoflavones are not considered likely to exert
354 hepatic changes with any clinical implications.

355

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360

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