

2 **Caloric Substitution of Diets with Apple Pomace was Determined to be Safe for**
3 **Renal and Bone Health Using a Growing Rat Model**
4

5 **Abstract**

6 Aims: to determine the safety of caloric substitution with 10% apple pomace substitution (g/kg)
7 to a healthy or Western diet.

8 Study design: Growing (age 22-29 days) female Sprague-Dawley rats were randomly assigned
9 (n=8 rats/group) to consume a purified standard rodent diet (AIN-93G), AIN-93G/10% g/kg
10 apple pomace (AIN/AP), Western diet, or Western/10% g/kg apple pomace (Western/AP) diets
11 for 8 weeks.

12 Results: Histological evaluation showed renal interstitial hypercellularity in rats fed AIN/AP,
13 Western, and Western/AP diets. However, there was no effects on renal expression of oxidative
14 stress and inflammatory genes or serum measures of kidney damage and function among diet
15 groups. Apple pomace is also high in calcium which can affect calcium balance. Dietary calcium
16 consumption was highest ($P < .001$) in rats consuming Western/AP. However, there was no
17 significant differences in calcium absorption and retention among diet groups. Further, there
18 was no evidence of renal calcification. There were also impact on femoral calcium and total
19 mineral content, size, and strength.

20 **Conclusions: Based on the results, apple pomace consumption was safe for renal and bone**
21 **health in a rodent model, regardless of diet quality. Future prestudies should be conducted to**
22 **further determine the efficacy and safety of apple pomace.**

23
24 **Keywords:** apple pomace, safety, minerals, Western diet, bone, kidney, sustainability
25

26 **1. Introduction**

27 Apple processing generates waste, consisting of skin, stem, seeds, and calyx,
28 collectively known as apple pomace. The environmental pollution and burden of waste disposal
29 costs to apple farmers and producers can be decreased by re-purposing apple pomace as a
30 product for human consumption. However, among popular consumed fruits, apples had the
31 highest fructose content [1]. Muir, et al. [2] reported apples to have 10.5 g of fructose/serving
32 compared to 3.2 g/serving for bananas, 6.4 g/serving for blueberries, and 2.5 g/serving for
33 oranges. Further, apple pomace contains 44.7% fructose compared to 5.8-6.0% fructose in
34 whole apple [3]. This is a health concern because fructose overconsumption has been reported
35 to contribute to renal disease and to produce deleterious effects on bone [4,5]. Apple pomace
36 contains a higher mineral content than whole apples, particularly calcium which is required for
37 bone health [1,6]. However, overconsumption of calcium can increase nephrocalcinosis and
38 reduced kidney function [7]. In turn, renal dysfunction can lead to bone loss due to mineral
39 imbalance, resulting in increased risk of osteoporosis and other bone-mineral disorders [8].

40 Diets typical of Western countries are characterized by high-fat and high-sucrose.
41 Western diet consumption has been shown to increase the risk of chronic kidney disease by
42 inducing renal steatosis, inflammation, and oxidative stress. Western diet consumption has also
43 been reported to increase risk of kidney stones due to the high sugar content [9]. Additionally,
44 consuming a Western diet can result in early onset of osteoporosis by promoting mineral
45 balance and inflammation leading to decreased bone mineral density [10,11].
46 Dietary advice suggests replacing calories in the diet with healthier food choices instead of
47 dietary supplementation with a purified isolated nutrient [12].

48 Previously, our laboratory reported caloric substitution of a Western diet with 10% g/kg
49 apple pomace attenuating features of NAFLD [13]. However, the effects of apple pomace on
50 renal and bone was not assessed in this study. To our knowledge no studies have evaluated the
51 safety of apple pomace consumption on renal and bone health. Therefore, the objectives of this

52 study were to determine the safety of apple pomace, due to its high fructose content and
53 increased calcium content, in growing rats consuming a “healthy” and Western diet. Female rats
54 were used due to their increased susceptibility to nephrocalcinosis, and growing rats because
55 kidney disease has been shown to have more severe bone effects in a pediatric population
56 [14,15]. We hypothesize apple pomace will not detriment kidney or bone health in growing
57 female rats consuming “healthy” or Western diets.

58

59 **2. Materials and Methods**

60 **2.1 Diets**

61 Locally sourced apple pomace was provided by Swilled Dog Hard Cider Company
62 (Franklin, WV). Apple pomace was freeze dried in equipment? Nutrient composition analysis of
63 apple pomace was performed by Medallion Laboratories (Minneapolis, MN). Apple pomace
64 contains 32.5% fructose compared to the published average of 5.9% fructose for whole apples.
65 Dietary calcium and phosphorus were determined by inductively coupled plasma mass
66 spectrometry (ICP) (model P400, Perkin Elmer, Shelton, CT). Freeze-dried apple pomace
67 contained 1.47 mg/g calcium and 1.97 mg/g phosphorous (**Supplementary Table 1**) compared
68 to respective published values of 0.06 mg/g and 0.11 mg/g in whole apples [13].

69 The ‘healthy’ diet was the standard purified diet American Institute of Nutrition (AIN-93G)
70 for growing rats [16] while a Western diet consisting of 45% fat and 34% sucrose was used to
71 typify the high-fat, high-sugar diet consumed by Western countries [17,18]. AIN-93G and
72 Western diet were calorically substituted with 10% g/kg freeze-dried apple pomace. AIN diets
73 were adjusted to be isocaloric (3.7-3.8 kcal/g) and Western diets were adjusted to be isocaloric
74 (4.7 kcal/g). The complete ingredient composition of experimental diets is provided as
75 **Supplementary Table 2**. Diets were stored at -20°C until fed to animals.

76

77 **2.2 Animals**

78 Weanling (age 22-29 days) female Sprague-Dawley rats (n=32) were purchased from
79 Harlan-Tekald (Indianapolis, IN). Rats were individually housed and kept in a room at constant
80 temperature of 21±2°C with a 12 h light/dark cycle throughout the study duration. Following a 7-
81 days acclimation, rats were randomly assigned (n=8 rats/group) to four dietary groups
82 consisting of: 1) AIN-93G, a standard purified rodent diet, 2) AIN-93G with 10% weight (g/kg)
83 substituted with apple pomace (AIN/AP), 3) Western diet (45% fat, 33% sucrose by kcals), or 4)
84 Western diet with 10% of weight (g/kg) substituted with apple pomace (Western/AP). Rats were
85 provided ad libitum access to their assigned diets and deionized distilled water (ddH₂O)
86 throughout the eight weeks study duration. Food intake was measured and assigned diets
87 replaced every other day while ddH₂O was replaced weekly. At the end of the study, rats were
88 fasted overnight then euthanized by carbon dioxide inhalation. The kidney was excised,
89 weighed, and then flash frozen in liquid nitrogen and stored at -80°C until analyzed. Both femurs
90 were removed, cleaned, and stored at -20°C.

91

92 2.3 Kidney histology

93 The left kidney was removed, weighed, flash frozen in liquid nitrogen, and stored at -
94 80°C until analysis. A center sagittal section was cut from each frozen tissue (n=6-8) and stored
95 in 10% neutral buffered formalin for 48 hours (fixation). After fixation, samples underwent a
96 dehydration protocol consisting of 10-15 minutes incubation in increasing ethanol
97 concentrations (50-to-100%) followed by two 20-minute incubations in xylenes. Following xylene
98 incubation, samples were incubated in molten paraffin wax for 20 minutes (infiltration) and
99 embedded into blocks. 5-7µm sections were cut and mounted on charged slides and sections
100 stained with hematoxylin and eosin. Histological evaluation included gross morphological
101 assessment which included the following: glomerular hypercellularity and matrix deposition,
102 interstitial hypercellularity, tubulointerstitial calcification, inflammation, and fibrosis. All slides
103 were analyzed using a Nikon Labophot 2 microscope (Nikon Instruments, New York, NY) at

104 magnification 10X by a trained investigator blinded to the identity of the groups. Images were
105 captured using a LCL-500-LHD digital camera with a PC Method Capture Imaging software
106 (Ludescop, Parkville, MD).

107

108 2.4 Renal RNA isolation and inflammatory gene expression

109 Total RNA was extracted from frozen kidney tissue (50 mg) using the Zymo Research
110 Direct-zol RNA Miniprep Plus Isolation Kit (Irvine, CA, catalog #R2071) according to the
111 manufacturer's instruction for total RNA isolation. Isolated RNA integrity was visualized on a
112 1.5% agarose gel and quantified by spectrophotometry (NanoDrop 100; Thermo Fisher
113 Scientific, Waltham, MA). Following DNase I treatment with TURBO DNA-free kit (Thermo
114 Fisher Scientific), total mRNA was amplified using the Superscript IV First-Strand Synthesis
115 System with oligo dT primers (Thermo Fisher Scientific).

116 Real-time quantitative polymerase chain reaction (RT-qPCR) consisted of 2.5 µl of
117 SYBR Green Master Mix (Thermo Fisher Scientific), 1 µL of cDNA (diluted 1:10), 1 µL of
118 respective forward and reverse primers and 0.5 µl of deionized distilled water for a total reaction
119 volume of 5 µl. The reactions were performed in a 7500 ABI Real-Time PCR System (Thermo
120 Fisher Scientific). The thermal profile consisted of 50°C for 2 min, 95°C for 10 min then 40
121 cycles of 95°C for 15 sec and 60°C for 1 min. A melt curve analysis was applied at the end of
122 cycling. Primers that were designed for transcription factors, nuclear factor kappa-light chain
123 enhancer of B cells (NFκB) and NADPH oxidase 4 (NOX4) and for inflammatory cytokines,
124 tumor necrosis factor-alpha (TNF-α), and interleukin-6 (IL-6) as well as for housekeeping genes,
125 β-actin and glyceraldehyde 2-phosphate dehydrogenase (GAPDH) using the Primer3 program
126 (Howard Hughes Medical Institute) and respective mRNA sequences obtained by NCBI.
127 Forward and reverse primers for genes of interest are listed in **Supplementary Table 3**:

128 2.5 Serum and urinary measures of renal function and health

129 Serum measures of kidney function included: blood urea nitrogen (BUN), creatinine, total
130 protein, calcium, phosphorous, alanine aminotransferase (ALT). Additionally, serum glucose
131 and amylase were measured. Values were determined enzymatically using a commercially
132 available Vet-16 rotor and quantified by a Hemagen Analyst automated spectrophotometer
133 (Hemagen Diagnostics Inc., Columbia, MD).

134 Serum and urine uric acid was determined by commercially available enzymatic assay
135 (Cayman Chemical). Briefly, serum and urine samples were aliquoted onto a 96-well plate and
136 incubated for 15 minutes. Reaction was initiated by adding 15 μ l of uricase and horseradish
137 peroxidase enzyme mixture, and read at an excitation of 535 nm and an emission of 590 nm
138 using a BioTek Synergy H1 microplate reader (Winooski, VT). Inter-assay coefficient of variation
139 was 32.1% for both serum and urine.

140

141 2.6 Calcium balance and retention

142 Rats were fasted overnight and euthanized by carbon dioxide inhalation. Blood was
143 collected by aorta puncture. Collected blood was centrifuged at 1,500 g for 10 min at 4°C to
144 obtain serum. Serum samples were stored at -80°C until analyzed. Serum calcium was
145 determined enzymatically using a commercially available Vet-16 rotor and quantified by a
146 Hemagen Analyst automated spectrophotometer.

147 During the initial and final weeks of the feeding study, rats were individually housed in
148 metabolic cages to collect urine and feces for 24 h. Initial and final day urine samples were
149 collected, centrifuged at 1,500 g for 10 min at 4°C, filtered through Whatman no. 1 paper, and
150 then diluted 1:10 in dd H₂O. Initial and final feces were collected and dried for 48 h, then ashed
151 in a muffle furnace (model CP18210, Thermolyne, Dubuque, IA) at 550°C for 24 h. Fecal
152 samples were then acidified in 70% nitric acid, neutralized in ddH₂O, filtered through Whatman
153 no. 1 paper, and further diluted (1:50 v/v) in ddH₂O. Ca content of feces and urine was
154 determined by ICP.

155 Measures of Ca excretion, absorption, and retention were performed according to
156 Maditz, et al. [20]. Briefly, urinary calcium excretion was calculated as urinary Ca
157 concentration/urine volume. Ca apparent absorption was calculated as $[(\text{Ca intake} - \text{fecal Ca}$
158 $\text{excretion}) / (\text{Ca intake})] \times 100$. Calcium retention was calculated as $[(\text{Ca intake} - (\text{fecal Ca}$
159 $\text{excretion} + \text{urinary Ca excretion})]$.

160

161 2.7 Femur morphometry and mineralization

162 Following CO₂ inhalation, the left and right femur were collected, and then defleshed.
163 After no bilateral differences were determined using a t-test with significance set at $P < .05$ left
164 femurs were used for all analyses. Femoral morphometry measurements of depth, width, and
165 length were determined using a Vernier caliper (Bel-Art Products, Pequannock, NJ, USA).
166 Length was measured from the medial condyle to the greater trochanter. Femurs were weighed
167 using an analytical balance (Mettler Toledo, Columbus, OH, USA).

168 Total bone mineral was determined by ashing in a muffle furnace at 600°C for 24 hours,
169 then weighed. To measure specific minerals, ash was dissolved in 2 mL of 70% nitric acid.
170 Acidified samples were filtered through Whatman no. 1 paper and diluted (1:50 v/v) to volume
171 with ddH₂O and Ca determined using ICP.

172

173 2.8 Femur biomechanical strength

174 Femoral strength indices were assessed using a TA,XT2i Texture Analyzer (Texture
175 Technologies, Scarsdale, NY, USA) fitted with a three point bending apparatus. Femora were
176 placed on supports and force applied to the midshaft marked at a position halfway between the
177 greater trochanter and the distal medial condyle. Bone was broken by lowering a centrally
178 placed blade (1 mm width) at a constant crosshead speed (0.1 mm/s). The load cell was 250 kg.
179 The load-deflection data were collected by a PC interfaced with the TA,XT2i. Sample test
180 distance was set at 10 mm with a signal collection rate of 100 points per second. Peak force,

181 ultimate stiffness, ultimate bending stress and Young's modulus were calculated according to
182 Yuan and Kitts [21].

183

184 2.9 Statistics

185 Results are expressed as mean \pm standard error of the mean (SEM). Gene expression
186 was determined as a function of mRNA abundance (A), where $A=1/(\text{gene of interest primer}$
187 $\text{efficiency} \times \Delta\text{CT (g.o.i.)} - (\text{average housekeeping primer efficiency} \times \Delta\text{CT (h.k.)})$, where the
188 product of efficiency and average of expression of β -actin was averaged with the product of
189 efficiency and average of expression of GAPDH to determine the overall expression of the two
190 housekeeping gene [17,29,22]. Gene expression data for each treatment group were log-
191 transformed prior to statistical analysis. One-way ANOVA was used to determine differences
192 among dietary groups. Post hoc multiple comparison tests were performed using Tukey's test
193 with treatment differences considered significant at $P = .05$ and a tendency at $P = .08$. All
194 statistical analyses were performed using JMP 12.2 statistical software package (SAS Institute,
195 Cary, NC).

196

197 3. Results and Discussion

198 Rats are susceptible to renal disease and diets high in fructose and high in calcium have
199 been shown to be detrimental to renal health, and high-fructose diets can detriment bone health
200 [5, 23]. In the current study, no differences were observed in body or organ weights (Table 1),
201 but histological analysis of the kidneys showed no evidence of fibrosis, glomerular
202 hypercellularity, glomerular matrix deposition, or amyloidosis.

203

204 **Table 1.** Weekly caloric and macronutrient intake, weekly body weight gain, and kidney and
205 bone weights of growing female rats consuming different diets substituted with apple pomace
206 (10% g/kg) for 8 weeks.

Measurements

Treatments¹

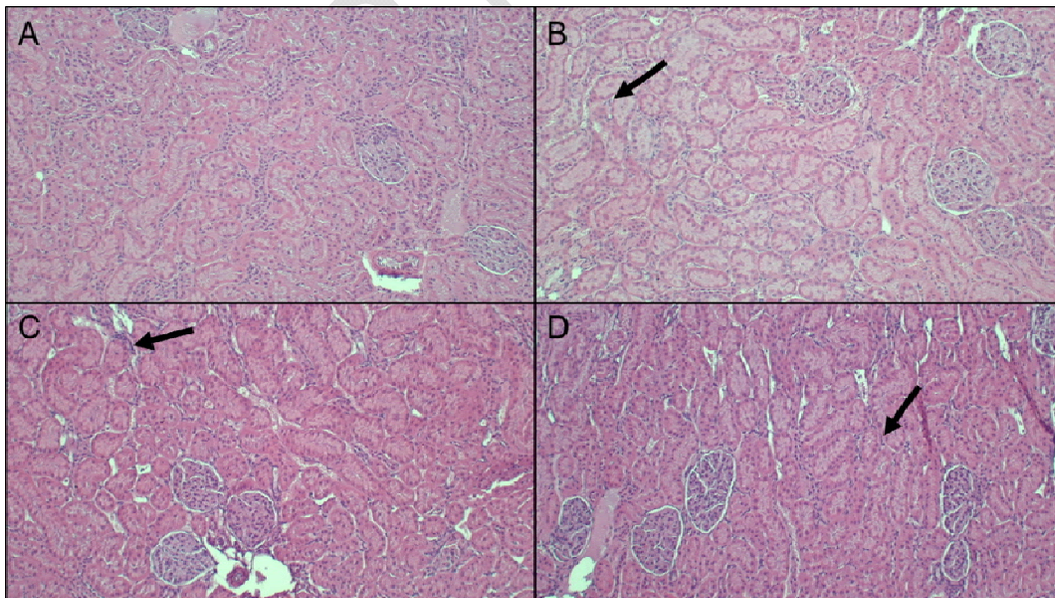
Histological changes	AIN	AIN/AP	Western	Western/AP
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	AIN	AIN/AP	Western	Western/AP	P-Value
Caloric intake (kcal/week)	368 ± 11 ^b	345 ± 8 ^b	422 ± 9 ^a	430 ± 17 ^a	<.001
Initial bwt (g)	95 ± 3	92 ± 3	95 ± 3	95 ± 3	0.80
Final bwt (g)	216 ± 4	216 ± 8	229 ± 5	234 ± 5	0.08
Average weekly bwt gain (g)	16 ± 3	16 ± 3	18 ± 3	18 ± 3	0.94
Average mineral intake (mg/d)	304.0 ± 9.3 ^b	318.8 ± 7.3 ^b	368.9 ± 7.8 ^a	374.7 ± 15.0 ^a	<.001
Right kidney weight (g)	0.69 ± 0.02	0.68 ± 0.02	0.71 ± 0.02	0.73 ± 0.02	0.28
Left kidney weight (g)	0.69 ± 0.02	0.67 ± 0.02	0.74 ± 0.03	0.74 ± 0.02	0.07
Relative right kidney weight (mg/g)	0.32 ± 0.01	0.31 ± 0.01	0.32 ± 0.01	0.31 ± 0.01	0.86
Relative left kidney weight (mg/g)	0.31 ± 0.01	0.31 ± 0.01	0.31 ± 0.01	0.32 ± 0.00	0.70
Left kidney ash (mg/g)	9.86 ± 0.56	10.07 ± 0.54	9.14 ± 1.09	10.34 ± 0.67	0.71

207 ¹Values expressed as mean ± SEM (*n* = 6–8 rats/group). Different superscript letters a and
208 b within the same row. Indicate significant difference at *P* < .05 by one-way ANOVA
209 followed by Tukey's test. Abbreviations: Bwt, body weight; CHO, carbohydrate.

210

211 However, rats consuming Western diet and diets containing apple pomace showed renal
212 interstitial hypercellularity (Figure 1), suggesting renal inflammation.



213

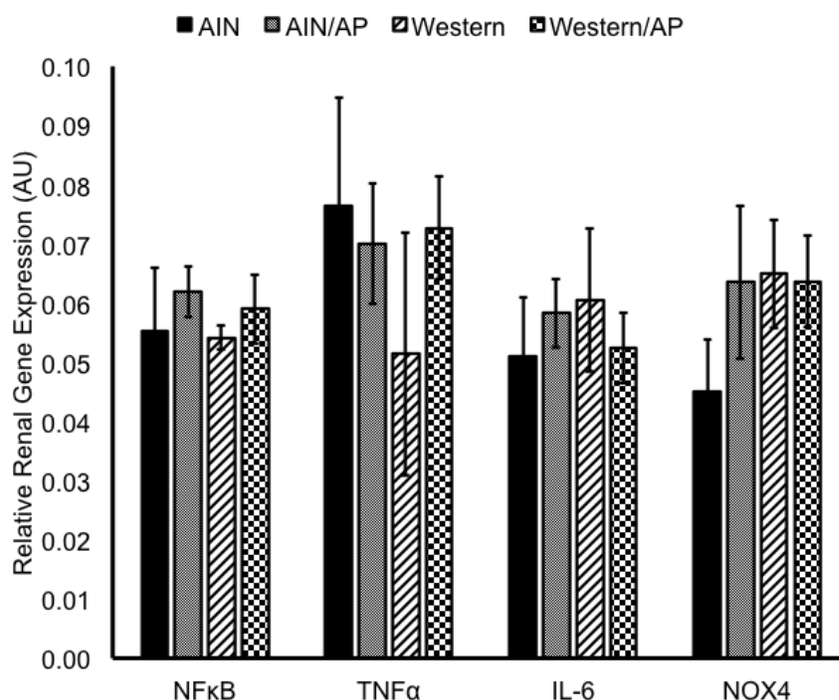
214

Inflammation	0	0	0	0
Fibrosis	0	0	0	0
Glomerular hypercellularity	0	0	0	0
Glomerular matrix deposition	0	0	0	0
Amyloidosis	0	0	0	0
Interstitial Calcification	0	0	0	0
Interstitial hypercellularity	0	2	1	2

215 **Figure 1.** Representative histological staining images of the kidney of growing female rats
 216 consuming (A) AIN, (B) AIN/AP, (C) Western, or (D) Western/AP following 8 weeks of feeding.

217

218 To further investigate, gene expression of inflammatory transcription factor, NFkB and
 219 proinflammatory cytokines, TNF- α and IL-6 as well as NOX4, a highly expressed enzyme
 220 regulating generation of reactive oxygen species, were measured in the kidneys. No significant
 221 differences were found in renal expression of any of the genes of interest among diet groups
 222 (Figure 2).



223

224 **Figure 2.** Renal expression of genes involved in inflammation and oxidative stress in rats
 225 consuming different diets substituted with 10% apple pomace (g/kg). Values expressed as
 226 mean \pm SEM (n=5-7 animals/group). Different superscript letters a and b within the same figure
 227 indicates significant difference at $P < .05$ by one-way ANOVA followed by Tukey's test.
 228 Abbreviations: AU, arbitrary units; IL-6, interleukin-6; NOX4, NADPH oxidase 4; NF κ B, nuclear
 229 factor kappa-light enhancer of B cells; TNF α , tumor necrosis factor alpha.

230
 231 Serum creatinine, BUN, ALT, and total protein also showed no significant differences among
 232 diet groups, collectively indicating absence of inflammation and oxidative stress (Table 4).

233 Increased fructose consumption and elevated uric acid may play a role in renal
 234 inflammation [24]. Elevations in uric acid levels have been shown to change the fundamental
 235 architecture of renal histology and has been implicated in acute and chronic renal failure [25].
 236 The current study results showed no significant difference in serum or urine uric acid among diet
 237 groups (Table 2).

238 **Table 2.** Effect of consumption of different diets substituted with apple pomace (10% g/kg) by
 239 growing female rats on serum and urine measurements of liver function enzymes, and uric acid
 240 following 8 weeks of feeding.

Measurements	Treatments ¹				P-Value
	AIN	AIN/AP	Western	Western/AP	
Serum Creatinine (U/L)	1.46 \pm 0.08	1.45 \pm 0.11	1.38 \pm 0.09	1.43 \pm 0.04	0.90
Serum BUN (mg/dl)	17.84 \pm 1.59	19.63 \pm 1.41	20.25 \pm 2.32	16.00 \pm 0.94	0.27
Serum ALT (U/L)	107.63 \pm 19.59	118.71 \pm 43.60	94.5 \pm 12.58	133.5 \pm 30.59	0.78
Serum Total Protein (g/dl)	3.9 \pm 0.25	4.62 \pm 0.34	4.08 \pm 0.67	4.19 \pm 0.34	0.79
Serum Phosphorous (mg/dl)	14.18 \pm 0.54	13.46 \pm 1.72	15.68 \pm 0.53	13.09 \pm 1.02	0.35
Serum Ca (mg/dl)	9.56 \pm 0.80	11.10 \pm 1.09	11.49 \pm 0.54	10.51 \pm 1.00	0.48
Serum Uric Acid (μ M)	7.24 \pm 0.31	6.27 \pm 1.61	7.19 \pm 0.86	7.57 \pm 1.25	0.86
Urine Uric Acid (μ M)	5.94 \pm 2.26	10.35 \pm 2.11	10.40 \pm 1.12	6.79 \pm 1.41	0.23

241 ¹Values expressed as mean \pm SEM (n=4-8 animals/group). Different superscript letters a and b
 242 within the same figure indicates significant difference at $P < .05$ by one-way ANOVA followed by
 243 Tukey's test. Abbreviations: ALT, alanine aminotransferase; BUN, blood urea nitrogen.

244
 245 Interstitial hypercellularity was observed in 13-29% of animals, but there were no
 246 significant differences in oxidative stress and inflammatory gene expression or serum and urine

247 measurements of renal dysfunction and injury were observed among diet groups. These results
 248 indicate renal interstitial hypercellularity was unlikely to be of biological significance. Collectively,
 249 the results indicate the fructose content of apple pomace was not a risk for renal injury and
 250 development of chronic kidney disease in either 'healthy' or Western diet.

251 In our study, Western diets were high in calcium with Western/AP diet having the highest
 252 calcium content (Table 2). Differences in calcium content in diets can have significant effects on
 253 calcium excretion, absorption, and retention [20]. Increased calcium excretion can induce
 254 nephrocalcinosis [26]. Initial urinary and fecal calcium excretion, calcium retention, and calcium
 255 absorption showed no significant differences among diet groups (Table 3). At final week, no
 256 differences were observed in rats urinary calcium excretion among all groups, but an increase
 257 ($P = .04$) in fecal calcium excretion by rats consuming a Western/AP compared to AIN was
 258 observed. This was also likely due to a combination of the high insoluble dietary fiber content in
 259 apple pomace possibly binding to calcium and the increased dietary calcium in Western/AP
 260 diets. This also explains the lack of change in apparent calcium absorption among all diet
 261 groups. No differences were observed in calcium retention among all diet groups.

262

263 **Table 3.** Calcium balance of rats fed different diets substituted with 10% (g/kg) apple pomace.

Calcium Balance	Treatments ¹				P-Value
	AIN	AIN/AP	Western	Western/AP	
Ca Intake (mg/d)	135.6 ± 4.2 ^c	140.1 ± 3.2 ^c	162.4 ± 3.5 ^b	184.9 ± 7.4 ^a	<0.001
Initial					
Urine Ca excretion (mg/dl)	0.16 ± 0.04	0.19 ± 0.04	0.17 ± 0.04	0.18 ± 0.04	0.96
Fecal Ca excretion (mg/d)	25.9 ± 3.6	22.9 ± 3.5	31.3 ± 3.7	34.7 ± 2.7	0.12
Ca retention (mg/d)	89.3 ± 9.4	94.9 ± 5.9	96.4 ± 5.8	109.8 ± 6.2	0.32
Ca absorption (%)	62.5 ± 4.6	68.0 ± 4.7	61.4 ± 4.2	63.3 ± 3.0	0.70
Final					
Urine Ca excretion (mg/ml)	0.15 ± 0.02	0.16 ± 0.04	0.16 ± 0.04	0.10 ± 0.01	0.25
Fecal Ca excretion (mg/d)	60.9 ± 2.9 ^b	79.4 ± 11.6 ^{ab}	81.2 ± 3.9 ^{ab}	99.3 ± 7.1 ^a	0.04
Ca retention (mg/d)	77.7 ± 5.3	66.7 ± 5.3	80.8 ± 5.0	78.9 ± 5.3	0.25
Ca absorption (%)	54.2 ± 4.1	41.8 ± 11.8	49.7 ± 3.2	46.3 ± 5.3	0.65

264 ¹Values expressed as mean ± SEM (n=4-8 animals/group). Different superscript letters a and b
 265 within the same figure indicates significant difference at $P < .05$ by one-way ANOVA followed by
 266 Tukey's test.

267

268 Further, renal histological evaluation showed no evidence of calcium deposition in any of the
269 diet groups, further indicating apple pomace consumption to be safe (Figure 2).

270 While Western diet (high fat and high sugar) and fructose consumption have also been
271 reported to detriment bone health, whole apples have been shown to favorably alter bone
272 health, through increased bone mineral density, decreased calcium loss, and decreased
273 inflammation due to antioxidants present in apples [27,28]. Apple pomace has been shown to
274 contain more calcium than apples [3]. Increasing dietary calcium has been shown to prevent
275 osteoporosis and to lower the risk of bone fractures [29]. Further, children with adequate
276 calcium consumption had increased bone mineral density [30]. The present study showed no
277 significant differences in femoral calcium content among diet groups. Additionally, there were no
278 significant differences in femur size or bone strength measurements including: peak force,
279 ultimate stiffness, ultimate bending stress, and Young's modulus among diet groups (Table 4).

280 **Table 4.** Femoral morphometry and strength measurements of rats fed different diets substituted with 10% (g/kg) apple pomace.

Measurement	Treatments ¹				P-value
	AIN	AIN/AP	Western	Western/AP	
Femur morphometry					
Length (mm)	29.71 ± 0.53	29.09 ± 0.78	30.52 ± 0.56	29.36 ± 0.78	0.09
Medial lateral width (mm)	2.98 ± 0.04	3.12 ± 0.12	3.06 ± 0.08	3.15 ± 0.10	0.13
Depth (mm)	2.78 ± 0.07	2.73 ± 0.12	2.60 ± 0.09	3.06 ± 0.17	0.43
Wet wt (g)	0.77 ± 0.02	0.74 ± .05	0.73 ± 0.03	0.74 ± 0.04	0.89
Dry wt (g)	0.48 ± 0.01	0.46 ± 0.03	0.45 ± 0.02	0.47 ± 0.02	0.77
Femur mineralization					
Ash (mg/g of bone)	407.92 ± 11.42	407.75 ± 9.26	399.66 ± 7.40	396.94 ± 6.46	0.80
Calcium (mg/g of bone)	37.99 ± 0.78	39.09 ± 4.41	40.09 ± 2.26	38.28 ± 2.08	0.75
Femur biomechanical strength					
Peak force (N)	1.74 ± 0.18	1.99 ± 0.25	1.55 ± 0.11	1.23 ± 0.23	0.07
Ultimate stiffness (N/S)	382.03 ± 16.28	399.49 ± 27.07	397.55 ± 38.73	347.15 ± 14.01	0.60
Ultimate bending stress (N/S)	42.32 ± 1.57	38.21 ± 2.19	40.12 ± 3.46	42.19 ± 2.59	0.48
Young's Modulus (N/mm ²)	1604.92 ± 76.18	1484.85 ± 284.92	1549.57 ± 90.13	1275.92 ± 200.17	0.75

281 ¹Values expressed as mean ± SEM (n=6-8 animals/group). Different superscript letters a and b within the same figure indicates
 282 significant difference at $P \leq .05$ by one-way ANOVA followed by Tukey's test.
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288 Another concern is rats consuming Western/AP diets had significantly increased gonadal
289 fat pad weights than rats consuming AIN diets (Table 1). Obesity and diabetes have been
290 reported to be causal factors in diet-induced kidney disease progression [4,31]. In our study,
291 despite higher adiposity in rats fed Western/AP there were no significant differences in fasting
292 serum glucose or amylase among diet groups (data now shown). Our study provides evidence
293 that high fructose and high calcium content of apple pomace was not sufficient to effect renal or
294 bone health in rats, regardless of diet. Studies on apple pomace have reported numerous health
295 benefit including decreases in body weight, as well improvements in serum lipid, insulin,
296 glucose, antioxidant status, digestion, and gut health [32-37]. Yet, few studies have evaluated
297 the safety of apple pomace consumption. Devrajan, et al. [38] fed rats unfermented or
298 fermented apple pomace for 2 weeks showed a nonsignificant increase serum BUN, but found
299 no indication of kidney damage. Additionally, histology was not used to evaluate kidney health.

300

301 **4. Conclusions**

302 Caloric substitution of a healthy or Western diet with 10% apple pomace had no impact on renal
303 or bone health in growing female rodents. Based on our results apple pomace is safe for
304 consumption, despite its high fructose content combined with a high calcium content, regardless
305 of diet quality in rodents. The study provides evidence for apple pomace, a “waste” byproduct of
306 apple processing, has a favorable nutritional profile and is safe and therefore has potential to be
307 repurposed as a sustainable food source for human consumption. However, the impact of
308 processing on apple pomace should be further evaluated for processing and undesirable
309 compounds before human clinical trials are conducted to determine the efficacy and safety of
310 apple pomace consumption.

311

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316

317 Competing Interests

318 Authors have declared that no competing interests exist.

319

320

321 **Ethics:**

322 All animal procedures were approved by the Animal Care and Use Committee at West Virginia
323 University and conducted in accordance with the guidelines of the National Research Council
324 for the Care and Use of Laboratory Animals [23]. All authors hereby declare that "Principles of
325 laboratory animal care" (NIH publication No. 85-23, revised 1985) were followed, as well as
326 specific national laws where applicable. All protocols have been examined and approved by the
327 appropriate ethics committee

328 **Consent:** NA

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445 **Supplemental Material**

446

447 **Supplementary Table 1.** Composition of locally sourced freeze-dried apple pomace.

Macronutrients (%)	
Protein	3.56
Fat	1.3
Carbohydrates	68.1
Sugars (%)	
Fructose	32.5
Glucose	9.77
Sucrose	13.9
Maltose	<0.1
Lactose	<0.1
Dietary Fiber (%)	
Insoluble Dietary Fiber	22.2
Soluble Dietary Fiber	11.0
Polyphenols (g/kg)	0.029
Minerals (mg/g)	
Total Minerals	15.5
Calcium	1.47
Phosphorous	1.97
Calories (kcal/100 g)	387

448

449

450 **Supplementary Table 2. Composition of rodent diets substituted with apple pomace**
 451 **(10% g/kg) fed to growing female rats.**

	Diet Groups *			
	AIN	AIN/AP	Western	Western/AP
Ingredients (g/kg) *				
Apple pomace	0.0	100.0	0.0	100.0
Corn Starch	397.486	392.086	63.36	57.96
Maltodextrin	132.0	132.0	60.0	60.0
Sucrose	100.0	43.9	340.0	283.9
Fructose	50	54.45	170	174.45
Total Dietary Fiber	50.0	50.0	50.0	50.0
Insoluble Fiber †	50.0	39.0	50.0	39.0
Soluble Fiber ‡	0.0	11.0	0.0	11.0
Anhydrous Milkfat	0.0	0.0	210.0	210.0
Soybean Oil	70.0	68.7	20.0	18.7
Casein	200.0	196.0	195.0	191.0
L-Cystine	3.0	3.0	3.0	3.0
Vitamin Mix	10.0	10.0	12.5	12.5
Mineral Mix	35.0	35.0	43.0	43.0
Total Minerals	22.1	24.2	26.4	28.0
Calcium	10.4	10.8	12.8	14.6
Phosphorous	7.2	7.5	7.6	7.5
Choline Bitartrate	2.5	2.5	3.1	3.1
TBHQ, antioxidant	0.014	0.014	0.04	0.04
Polyphenols	0.0015	0.0029	0.0008	0.0032
Macronutrients (% kcal)				
Protein	18.8	18.9	14.8	14.8
Fat	17.2	17.3	44.6	44.8
Carbohydrate	63.9	63.7	40.6	40.4
Calories (kcal/g)	3.8	3.7	4.7	4.7

452 * Abbreviations: AIN, the American Institute of Nutrition; AP, apple pomace; TBHQ, tert-
 453 butylhydroquinone. † Insoluble fiber is cellulose. ‡ Soluble fiber is mainly pectin [1].
 454

455 **Supplementary Table 3.** Forward and reverse primers for genes of interest in study.

Gene	NCBI Gene ID	Forward Primer	Reverse Primer
NFκB	81736	5' TTATGGGCAGGAT GGACCTA 3'	5' CCTTTCAGGGCTTT GGTTTA 3'
TNFα	24835	5' CACAAGGCTGCTG AAGATGT 3'	5' GAGGGAAGGAAGG AAGGAAG 3'
IL-6	24498	5' TGGCTAAGGACC AAGACCAT 3'	5' TTGCCGAGTAGAC CTCATAGTG 3'
NOX4	85431	5' CCTCCATCAAGCC AAGATTC 3'	5' CTCCAGCCACACA CAGACTAAC 3'
β-actin	81822	5' TTGCTGACAGGAT GCACAAG 3'	5' CAGTGAGGCCAGG ATAGAGC 3'
GAPDH	24383	5' TCAAGAAGGTGGT GAAGCAG 3'	5' CCTCAGTGTAGCC CAGGATG 3'

456

UNDER PEER REVIEW