

1 ***Pgc1a* is responsible for the sex differences in hepatic *Cidec/Fsp27β***
2 **mRNA expression in hepatic steatosis of mice fed a Western diet**
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45 **Running title:** Hepatic *Cidec/Fsp27* gene expression

46

47 **Abbreviations:** *Pgc1a*, peroxisome proliferator-activated receptor gamma coactivator

48 1-alpha

49 **Abstract**

50 Hepatic fat-specific protein 27 (*Cidec/Fsp27*) mRNA levels have been
51 associated with hepatic lipid droplet extent under certain circumstances. To address its
52 hepatic expression under different dietary conditions and in both sexes, *ApoE*-deficient
53 mice were subjected to different experimental conditions for 11 weeks to test the
54 influence of cholesterol, Western diet, squalene, oleanolic acid, sex and surgical
55 castration on *Cidec/Fsp27* mRNA expression. Dietary cholesterol increased hepatic
56 *Cidec/Fsp27 β* expression, an effect that was suppressed when cholesterol was combined
57 with saturated fat as represented by Western-diet feeding. Using the latter diet, oleanolic
58 acid or squalene did not modify its expression. Females showed lower levels of hepatic
59 *Cidec/Fsp27 β* expression than males when they were fed Western diets, a result that
60 was translated into lesser amount of CIDEC/FSP27 protein in lipid droplets and
61 microsomes. This was also confirmed in *Ldlr*-deficient mice. Incubation with estradiol
62 resulted in decreased *Cidec/Fsp27 β* expression in AML12 cells. While male surgical
63 castration did not modify the expression, ovariectomized females did show increased
64 levels compared to control females. Females also showed increased expression of
65 *Pgc1a*, suppressed by ovariectomy, and the values were significantly and inversely
66 associated with those of *Cidec/Fsp27 β* . When *Pgc1a*-deficient mice were used, the sex-
67 differences on *Cidec/Fsp27 β* expression disappeared. Therefore, hepatic *Cidec/Fsp27 β*
68 expression has a complex regulation influenced by diet and sex hormonal milieu. The
69 mRNA sex differences are controlled by *Pgc1a*.

70 **Keywords:**

71 Lipids/liver, lipid droplets, animal models, gene expression, non-alcoholic fatty liver
72 disease, *Cidec/Fsp27*, *Pgc1a*, apolipoprotein E deficient mice, high-fat diet, sex.

73

74 **Introduction**

75 Fat-specific protein 27 (*FSP27*) gene encodes a protein of 27 kDa with 238
76 amino acids, belonging to the cell-death-inducing DNA fragmentation effector (CIDE)
77 family, composed of CIDEA, CIDEB, and CIDEA/CIDE-3/*FSP27*, all of which contain
78 a conserved CIDE N-domain and a unique C-terminal domain. *Cidec/Fsp27* is
79 expressed at high levels in white adipose tissue (26). By alternative splicing in HepG2,
80 *CIDE-3* gene displays two transcripts, *CIDE-3* and *CIDE-3alpha*. While *CIDE-3*
81 comprises a full-length open reading frame, *CIDE-3alpha* encodes a truncated protein
82 (29). In the liver, a third transcript, *FSP27β*, which contains 10 additional amino acids at
83 the N-terminus of the original protein and is activated through the liver-enriched
84 transcription factor cyclic-AMP-responsive-element-binding protein H (CREBH) but
85 not by peroxisome proliferator-activated receptor gamma, has been described (11, 64).
86 In this organ, *CIDEA/CIDE-3/FSP27* contributes to triglyceride accumulation both in
87 humans and pigs (28) and to the regulation of lipidation and maturation of very low-
88 density lipoproteins (63). It is localized to lipid droplets (LD) and endoplasmic
89 reticulum (56). The latter participates in the regulation of LD formation, expansion, and
90 morphology under lipid-deficient conditions (25). To promote the formation of a
91 unilocular droplet, the formation a ternary complex of AS160, the GTPase activating
92 protein for RAB8a, *FSP27* and RAB8a is required (60).

93 *Fsp27*-deficient mice show increased energy expenditure and lower levels of
94 plasma triglycerides and free fatty acids (39). Only when they are crossed with leptin-
95 deficient mice or BATless mice, or are fed them a high-fat diet, hepatic steatosis and
96 insulin resistance are observed. Therefore, *Fsp27* deficiency requires further implication
97 of genes to display hepatic insulin resistance (58, 69). In contrast, mice with adipocyte-
98 specific disruption of the *Fsp27* gene upon high-fat diet feeding are resistant to weight

99 gain and fat-storing. This results in a lipid overflow from adipose tissue that generates
100 hepatosteatosis, dyslipidemia, and systemic insulin resistance pointing out a role for this
101 adipocyte protein to prevent lipodystrophies (57). An increased expression of *Cidec* has
102 been found in a number of experimental or pathological conditions, such as in
103 endoplasmic reticulum stress (24), spontaneous mouse insulin resistance (52) and
104 hepatocellular carcinoma cells (37). Similar effects have been described in liver
105 steatosis and in obese humans (13), being the latter increase reduced by weight loss
106 (16). A homozygous human mutation of CIDEC has been reported to induce
107 lipodystrophy and insulin-resistant diabetes (40, 49). Reduced expression of hepatic
108 *Fsp27* abolished fasting-induced liver steatosis (23) and the former condition in
109 combination with a PPARalpha agonist was also found to reduce hepatic steatosis (45)
110 and even atherosclerosis (46) in *Ldlr*-deficient mice, a model of atherosclerosis and
111 hepatosteatosis (50).

112 The expression of CIDEC is controlled at both transcriptional and
113 posttranslational levels (5, 13). Different molecules seem to be involved in its
114 expression, such as CD44 (17) or osteopontin, whose absences decrease its levels (22),
115 while leptin absence displays the opposite (35). Ceramide (27) and TNF-alpha reduced
116 its expression while insulin upregulated it. In the latter response, the activity of
117 phosphatidylinositol 3-kinase was involved (21), so was the phosphatase and tensin
118 homologue, an enzyme involved in degradation of phosphorylated phosphatidylinositol
119 (51). Final effectors of these signaling pathways seem to be nuclear receptors such as
120 TAK1/TR4/NR2C2, RORalpha and PPARalpha, nuclear proteins (CAAT-enhancer-
121 binding proteins), LXR α and SREBP-1c (3, 7, 8, 18, 19, 23). Peroxisome proliferator-
122 activated receptor gamma2 (PPARgamma2) also plays a role (34). Posttranslational
123 regulation of FSP27 involves stability through the proteasomal ubiquitin-dependent

124 protein catabolic process (68), glycosylation (66) and acetylation (44).

125 A complex physiological regulation of CIDEC seems to exist, in which fasting
126 and diet composition play important roles. In this regard, during the initial stages of
127 fasting, *Fsp27* expression has been found dramatically increased by involvement of the
128 PKA-CREB-CRTC2 signaling pathway (59). The fasting effect was not present in
129 PPARalpha-deficient mice (67). However, after a long period of fasting, a decrease in
130 *Fsp27* expression was observed (59). Despite the observed hepatic steatosis after a
131 choline-deficient diet, no changes were observed for *Fsp27* (67). Nevertheless, a
132 marked induction of its expression was found in the high-fat- or methionine- and
133 choline-deficient diet-induced fatty liver, but not in alcohol-induced fatty liver. The
134 induction of *Fsp27* mRNA was independent of peroxisome proliferator-activated
135 receptor gamma (PPARgamma) levels and completely absent in the liver from
136 PPARgamma-deficient mice (2). In less extreme dietary conditions, it has been reported
137 that a high fat diet increased *Fsp27* expression through activation of PPARgamma (41).
138 In vitro, a high supply of fatty acids stimulated hepatic expression (25). Using *ApoE*-
139 deficient mice as a model of spontaneous hepatosteatosis, nature of fatty acids was
140 important to increase its expression in these mice fed a Western-type diet enriched with
141 linoleic acid isomers since only those mice receiving trans-10, cis-12-conjugated
142 linoleic acid showed this effect. Furthermore, consuming olive oil-enriched diet reduced
143 *Fsp27* expression (15). In addition, only one study has addressed the influence of sex on
144 its expression in young mice (12). Growth hormone has also found to regulate this
145 protein (53, 54). Therefore, influence of sex may be different in adult mice. Based on
146 these facts, it was hypothesized that *Fsp27* hepatic regulation might be the result of
147 complex interactions of dietary components and sex. To this end, the present work was
148 undertaken to characterize the influence of different dietary conditions and sex on

149 *Fsp27* gene expression in adult liver of several animal models.

150

152 *Animals*

153 Charles River (Charles River Laboratories, Barcelona, Spain) was the source of
154 *ApoE*-deficient mice on the C57BL/6J genetic background. Dr. Nobuyo Maeda
155 (University of North Carolina at Chapel Hill, NC, USA) generously provided these
156 mice on the C57BL/6JxOla129 genetic background. *Ldlr*-deficient mice on the
157 C57BL/6J.SJL genetic background were obtained from Dr. Vicente Andrés from CNIC,
158 (Madrid, Spain). C57BL/6J wild-type and *Pgc1a*-deficient mice were part of a colony
159 established at the IIB animal facility (Madrid) and originally derived from mice
160 provided by Dr. Bruce Spiegelman (DFCI, Boston, USA). Wistar rats were obtained
161 from Charles River (Charles River Laboratories, Barcelona, Spain).

162 For all experiments, two-month-old mice were used. Blood samples were taken
163 (after four-hour fasting) from the facial vein to determine plasma cholesterol and
164 accordingly establish groups with similar initial values. Animals, housed in sterile filter-
165 top cages, were maintained under a 12-h light/12-h dark cycle at the *CIBA, Universidad*
166 *de Zaragoza*. *Pgc1a*-deficient mice were maintained at *Autónoma Universidad de*
167 *Madrid*. Wistar rats were maintained at *Universidad de Córdoba*. Animals were handled
168 and killed observing guidelines (Directive 2010/63/UE) from the European Union for
169 care and use of laboratory animals in research. All had ad libitum access to food and
170 water and study protocols were approved by the Ethics Committees for Animal
171 Research of the Universities of Zaragoza, Madrid and Córdoba. After the diet
172 intervention, and four-hour fast, the animals were killed by suffocation with CO₂. The
173 livers were removed, weighed, frozen in liquid nitrogen, and stored at – 80 °C until
174 analysis.

175 *Experimental design*

176 Table 1 provides detailed information of all experimental designs regarding

177 characteristics of animals, type of diets, number of animals and length of intervention.
178 Since C57BL/6J mice do not express hepatic *Cidec/Fsp27* (35, 59), we decided to use
179 *ApoE*-deficient mice which showed hepatic expression of this gene influenced by some
180 dietary components (15). Using this model, we tested the effects on *Cidec/Fsp27*
181 expression of dietary cholesterol, Western diet and sex. On Western diet, the influence
182 of two modifiers of lipid droplet surface, oleanolic acid (10) and squalene (14) were
183 tested. Likewise, this diet was used to analyze sex differences and its inhibition by
184 surgical castration. A confirmation of the effects of ovariectomy on *Cidec/Fsp27*
185 expression was carried out in female Wistar rats fed a Western diet. Since sex
186 differences emerged on Western diet, this was also tested in another model of hepatic
187 steatosis, *Ldlr*-deficient mice on C57BL/6J genetic background. All previous
188 experiments were suggestive of an involvement of PGC1 α and to confirm such issue,
189 *Pgc1*-deficient mice on C57BL/6J genetic background fed a purified Western diet were
190 used to analyze the sex differences on hepatic *Cidec/Fsp27* expression. Detailed
191 compositions of purified diets are shown in supplementary Table 1.

192

193 *Isolation and quantification of hepatic RNA*

194 RNA was isolated using Tri-reagent (Ambion, Austin, TX, USA). Contaminant
195 DNA was removed using the DNA removal kit from Ambion. Absorbance at $A_{260/280}$
196 served to quantify RNA concentrations and the ratio 28S/ 18S ribosomal RNAs used to
197 estimate their quality. Changes in mRNA expression were determined by RT-qPCR.
198 cDNA synthesis was carried out using the First Strand synthesis kit (Thermo Scientific,
199 Madrid, Spain). The Sybr Green PCR Master Mix (Applied Biosystems, Foster City,
200 CA) was used to analyze gene expression by qPCR. Specific primers, designed and
201 checked as previously described (33) were purchased from Applied Biosystems.
202 Sequences are shown in supplementary Table 2. RT-qPCR reactions were performed on

203 a Step One Real Time PCR System (Applied Biosystems) following the standard
204 procedure and using equal amounts of DNA-free RNA from each animal. The relative
205 amount of all mRNAs was calculated using the comparative $2^{-\Delta\Delta Cq}$ method and
206 *Cyclophilin B (Pipb)* mRNA expression as the reference gene.

207 *Liver histology analyses*

208 Aliquots of liver, stored in neutral formaldehyde, were used and processed as
209 described (14).

210 *Hepatic homogenate and lipid extraction*

211 A piece of liver was homogenized in homogenization buffer (phosphate buffered
212 solution with protease inhibitor cocktail (Roche, Mannheim, Germany) and used to
213 assay protein concentration by the BioRad dye binding assay (BioRad, Madrid, Spain).
214 Extracted lipids according to Folch's method (9) were evaporated under N₂ stream and
215 dissolved in 100 μ L of isopropanol. Infinity kits (Thermo Scientific) were used to
216 measure total cholesterol and triglycerides.

217 *Preparation of microsomal fractions*

218 This fraction was prepared according to Osada et al. (43). Basically, 600 mg of
219 pooled hepatic tissue of each group were homogenized in 2 mL of 0.25 M sucrose
220 containing the Roche protease inhibitor cocktail at 4°C and centrifuged at 280g for 5
221 min. Supernatants were centrifuged at 1500g for 10 min followed by another
222 centrifugation at 19000g for 10 min to collect the supernatants containing cytosolic and
223 microsomal proteins. After a centrifugation at 100000g for 60 min, the obtained pellets
224 containing the microsomal fractions were resuspended in PBS containing 0.2% Triton
225 X-100 and 10% glycerol and centrifuged at 12000 rpm 10 min in order to remove
226 insoluble proteins. Protein concentrations were determined by BioRad dye binding
227 assay.

228 *Preparation of lipid droplets*

229 They were prepared following the protocol of Ontko et al. (42). Briefly, pooled
230 hepatic tissue (600 mg) of each group were homogenized in 3 ml of 65% sucrose
231 solution with protease inhibitor cocktail (Roche, Mannheim, Germany) at 4°C.
232 Discontinuous sucrose gradients were prepared as follows: 3 ml of liver homogenates in
233 65% sucrose were pipetted at the bottom of the centrifuge tubes kept in an ice bath.
234 Then 2 ml of 60% sucrose solution were slowly added, followed by 2 ml of 52%
235 sucrose, 2 ml of 44% sucrose and 2 ml of distilled water. The tubes with the gradients
236 were centrifuged at 25000g for 30 min at 4°C and the bands containing the different
237 lipid droplets were collected. They were mixed with 3 volumes of acetone and kept at -
238 80°C for 10 min and then at -20°C overnight. The tubes were centrifuged at 15000g for
239 15 min at 4°C. The pellets were washed three times, firstly with acetone: diethyl ether
240 1:1 and then twice with diethyl ether. Dry pellets were resuspended in PBS containing
241 0.2% Triton X-100 and 10% glycerol and centrifuged at 12000 rpm 10 min in order to
242 remove insoluble proteins. Protein concentrations were determined by BioRad dye
243 binding assay.

244 *Western blot*

245 20 µg of protein were loaded onto a 10% SDS-polyacrylamide gel and
246 electrophoresed for 120 min at 90V in a Bio-Rad Miniprotean cell (Hercules, CA).
247 Proteins were transferred to PVDF membranes (GE Healthcare, Madrid, Spain).
248 Membranes were blocked with PBS buffer containing 5% BSA for 1 h at room
249 temperature. The primary antibodies, diluted in PBS buffer containing 2.5% BSA and
250 1% Tween 20, were added and the membranes were incubated 2 h at room temperature
251 and then overnight at 4°C. FSP27 protein expression was evidenced by using a rabbit
252 polyclonal antibody (NB100-430 diluted 1/1,000, Novus Biologicals, Centennial,

253 Colorado, USA). Equal loadings were confirmed by using a goat polyclonal anti-HSC70
254 (TA302666 diluted 1/500, OriGene, Rockville, MD, USA). Membranes were washed
255 with PBS buffer containing 0.1% Tween 20. Conjugated goat anti-rabbit IgG (H&L)
256 DyLight 800 secondary antibody (SA5-35571, diluted 1/15,000, Thermo-scientific,
257 Waltham, MA, USA) for FSP27 detection and a donkey anti-goat IRDye 680RD (926-
258 68074, diluted 1/5,000, LI-COR Biosciences, Lincoln, NE, USA) for HSC70 detection
259 were used and incubated for 1 h at room temperature in PBS buffer containing 2.5%
260 BSA and 1% Tween 20. Images were captured using an Odyssey® Clx (LI-COR).

261 *AML12 cell culture*

262 The murine hepatocyte cell line was grown in a humidified atmosphere of 5%
263 CO₂ at 37°C in Dulbecco's modified Eagle's minimum essential medium (DMEM)
264 (ThermoFisher Scientific, Waltham, MA, USA): F12-Ham's medium (GE Healthcare
265 Life Science, South Logan, Utah) in 1:1 ratio supplemented with 10% foetal bovine
266 serum (ThermoFisher Scientific), 1:500 insulin/transferrin/selenium (Corning, Bedford,
267 MA, USA), 40 ng/ml dexamethasone (Sigma-Aldrich; Merck Millipore, Darmstadt,
268 Germany) 1% nonessential amino acids (ThermoFisher Scientific), 1% penicillin (1000
269 U/ml) (ThermoFisher Scientific), 1% streptomycin (1000 mg/ml) (ThermoFisher
270 Scientific) and 4 mM L-glutamine (ThermoFisher Scientific) in a 6 multiwell plate (in
271 triplicate). Medium was changed every two days. After one week of growth, this
272 medium was removed, and cells were washed twice with phosphate buffered saline
273 (PBS) prior to the addition of the serum-free media supplemented with 200 µm stearic
274 acid for 24 hours or 200 µM stearic acid for 24 hours and 50 nM estradiol dissolved in
275 ethanol for 6 hours. Then, media were removed and cells collected with Tri-reagent
276 solution (Ambion). RNA isolation and cDNA synthesis were performed as above
277 described.

278 *Reporter assays*

279 The genomic region -2042 bp to 0 bp at 5' side of the starting transcription site
280 of *CIDEA/FSP27β* (XM_024453700.1) from human genomic DNA was amplified by
281 PCR using direct (5'-agaaccagatcttggCAAGTGATCCACCTGCCTCG-3) and reverse
282 (5'-gatatctgcagaattGAGCAGATAACCCAACTCAGGGC -3') primers. The 2-kb PCR
283 product was cloned upstream a secreted Gaussia luciferase (GLUC) reporter gene using
284 linearized pEZX-GAO1 (Genecopeia Rockville, Maryland, USA) according to In-
285 Fusion® cloning protocol from Takara-Clontech (Cat No 638909, Kusatsu, Shiga,
286 Japan). Restriction enzymes and DNA sequencing confirmed the resulting plasmid. This
287 latter was transfected to AML12 cells alone or in combination with a plasmid
288 containing *Pgcl1a* (MN_008904) under the control of CMV promoter (MC204789,
289 Origene) using lipofectamine 3000 (ThermoFisher) following manufacturer's
290 instructions. Two days after, media were taken and secreted GLUC and alkaline
291 phosphatase, also present in pEZX-GAO1, activities were evaluated. The ratio of
292 GLUC/alkaline phosphatase was calculated.

293 *Statistical analysis*

294 The Statistical Package for Social Sciences version 15 (SPSS, Chicago, IL,
295 USA) or Prism 5 for windows software for Windows (GraphPad, S. Diego, CA, USA)
296 were used for statistical analyses. Variables, not showing normal distribution (according
297 to the Shapiro-Wilk's test), or homology of variance, were analyzed with the Mann-
298 Whitney's U test. Data are shown as medians and 10-90 percentile range of the values.
299 Correlations between variables were tested using the Spearman's correlation test. The
300 statistical significance was considered when $p < 0.05$.

301

302 **Results**

303 *Dietary fat and hepatic Cidec/Fsp27 β expression in Apoe-deficient mice.*

304 To characterize the dietary regulation of the expression of this gene in mice, the
305 supplement of dietary cholesterol to male mice was tested. Increased hepatic surface
306 occupied by lipid droplets as well as hepatic total cholesterol and triglyceride contents
307 (Fig 1a, b and c) were observed following dietary cholesterol supplementation. The
308 latter induced a significant increase in the hepatic *Cidec/Fsp27 β* expression as shown in
309 Fig 1d.

310 Hepatic cholesterol content was associated with hepatic *Cidec/Fsp27 β* expression (Fig
311 1e).

312 In a second study, the influence of a Western diet (WD), containing cholesterol
313 and palm oil as source of saturated fat, was explored in male *Apoe*-deficient mice on
314 C57BL/6J genetic background (Fig 2). Significant increased hepatic areas occupied by
315 lipid droplets (Fig 2a, b and c) as well as hepatic total cholesterol and triglyceride
316 contents were also observed in mice on the Western diet. Unexpectedly, a significant
317 decrease of hepatic *Cidec/Fsp27 β* expression was found (Fig 2d). These expression
318 changes were inversely associated with hepatic cholesterol (Fig 2e).

319 To further explore this dissociation between hepatic *Cidec/Fsp27 β* and Western
320 diet, its expression was tested in two dietary components that had been shown to
321 influence dietary droplets without altering lipid content (oleanolic acid) or viceversa
322 (squalene). In the first experiment and as expected, male mice receiving an oleanolic
323 acid-enriched WD showed an increase in the hepatic area occupied by lipid droplets
324 (Supplementary Fig 1a, b and c) without changes in hepatic cholesterol and triglyceride
325 contents (Supplementary Fig 1c). In these conditions, no significant change was
326 observed for hepatic *Cidec/Fsp27 β* expression (Supplementary Fig 1d). In the second

327 experiment, the effect of a squalene-enriched WD was explored, again in males. No
328 significant changes were noted for the percentage of hepatic surface occupied by lipid
329 droplets despite the decreased liver cholesterol and triglyceride contents
330 (Supplementary Fig 2a, b and c). Nor was there any significant change in the hepatic
331 *Cidec/Fsp27 β* expression by squalene (Supplementary Fig 2d). Overall, these nutritional
332 experiments emphasize that hepatic *Cidec/Fsp27 β* expression possesses a fine
333 nutritional regulation at transcriptional level in *Apoe*-deficient mice, where cholesterol
334 increased its levels and saturated fat reverted this finding, being the latter not influenced
335 by minor dietary components, such as oleanolic acid or squalene, despite the changes in
336 hepatic lipids.

337

338 *Hepatic Cidec/Fsp27 β expression is influenced by sex hormones in ApoE-deficient mice*
339 *and in vitro.*

340 The influence of sex on *Cidec/Fsp27 β* expression was explored in *ApoE*-
341 deficient mice on a chow diet of low fat content. As shown in Supplementary Fig 3,
342 panels a,b,c, females showed lower surface occupied by lipid droplets despite a
343 significant increase in hepatic cholesterol content and no changes in triglycerides. In
344 this experiment, no significant changes were observed in hepatic *Cidec/Fsp27 β* between
345 sexes. In a second experiment, the differences between sexes were explored when both
346 groups received a WD. As shown in Fig 3a, b and c, no significant change was observed
347 in the percentage of liver surface occupied by lipid droplets. However, the levels of
348 hepatic total cholesterol and triglycerides were significantly lower in females than in
349 males. In this experimental approach, females showed significantly decreased hepatic
350 *Fsp27 β* expression (Fig 3d). The latter was significantly associated with hepatic
351 triglyceride contents (Fig 3e). This mRNA decrease was translated in decreased

352 amounts of CIDEC/FSP27 protein in lipid droplets and microsomes (Fig 3f and g).
353 These data indicate that sex is playing an important role in hepatic *Cidec/Fsp27 β*
354 expression in the presence of WD and these changes are reflected in a lesser amount of
355 CIDEC in hepatic lipid droplets of female livers.

356 The involvement of hormonal changes on sex-differences was characterized in
357 *ApoE*-deficient mice of both sexes that underwent surgical removal of gonads and were
358 fed a purified Western diet. As shown in Supplementary Fig 4, no significant change in
359 *Cidec/Fsp27 β* expression was observed in orchietomized males; nor was there any
360 significant change in hepatic total cholesterol, or in hepatic triglycerides. However,
361 there was a significant increase in the liver surface occupied by lipid droplets in
362 orchietomized males. In contrast, ovariectomy resulted in significant increases in
363 hepatic cholesterol, triglycerides, and in the surface occupied by lipid droplets (Fig 4c).
364 Ovariectomized females showed a significant increase in *Cidec/Fsp27 β* expression
365 compared to control females (Fig 4d). A positive significant association was also found
366 between hepatic *Cidec/Fsp27 β* values and those of hepatic triglycerides (Fig 4e). The
367 increase in mRNA expression was translated into increased contents of CIDEC proteins
368 in lipid droplets and microsomes (Fig 4f and g). These results indicate that ovarian
369 hormones are responsible for the decreased hepatic *Cidec/Fsp27 β* expression observed
370 in females consuming WD. In fact, incubation of stearic-stimulated hepatic AML12
371 cells with estradiol resulted in a significant decrease in *Cidec/Fsp27 β* expression
372 (Supplementary Fig 5a).

373 *Pgc1a* is involved in hepatic *Cidec/Fsp27 β* expression sex differences in vivo.

374 PKA and PPAR have been described in the regulation of hepatic *Cidec/Fsp27*
375 expression (3, 7, 8, 18, 19, 23). To verify whether or not those agents were involved in
376 the observed sex-dependent responses, hepatic *Prka2* expression was determined and no

377 significant changes were observed (data not shown). Regarding PPARgamma, the
378 hepatic expression of its regulator, *Pgc1a*, was significantly increased in females
379 compared to males consuming the Western diet (Fig 5a) and an inverse significant
380 relationship was found between *Cidec/Fsp27β* expression and that of *Pgc1a* in both
381 sexes (Fig 5b). While orchietomy had no effect on *Pgc1a* expression (Fig 5c),
382 ovariectomy induced a significant decrease in its expression in *Apoe*-deficient females
383 (Fig 5d). Likewise, ovariectomized female rats also showed a trend to increase
384 *Cidec/Fsp27β* expression (Supplementary Fig 6e) and decreased hepatic *Pgc1a*
385 expression (Supplementary Fig 6f). Both effects were even more pronounced in rats
386 neonatally androgenized by testosterone administration and then ovariectomized. In this
387 model, hepatic fat, cholesterol and TG contents followed a similar pattern
388 (Supplementary Fig 6d) and *Cidec/Fsp27β* expression was associated with hepatic TG
389 content (data not shown). The sex differences in *Cidec/Fsp27β* expression were
390 observed in *Ldlr*-deficient fed on WD as well (Supplementary Fig 7d). Concomitantly, a
391 significant increase in *Pgc1a* expression was observed in these female mice
392 (Supplementary Fig 7e). Decreased *Cidec/Fsp27β* gene expression in females was
393 translated into lower amounts of CIDEC/FSP27 protein in lipid droplets and
394 microsomes (Supplementary Fig 7f and g). Overall, these findings are suggestive of an
395 inverse association between *Cidec/Fsp27β* and *Pgc1a* expressions as a general response,
396 independent of absence of APOE. These mRNA changes are reflected in CIDEC/FSP27
397 protein present in lipid droplets from female livers.

398 According to this association, it was hypothesized that *Pgc1a* would reduce the
399 transcriptional activity of a reporter gene under the control of CIDEC promoter. This
400 was the case, as shown in Supplemental Fig 5b. The opposite hypothesis would be
401 that sex-differences in hepatic *Cidec/Fsp27* would be abolished in the absence of

402 *Pgc1a*. To test this, *Pgc1a*- deficient mice from both sexes were fed WD. In this model,
403 female mice increased hepatic fat area, total cholesterol and TG contents (Fig 6a, b, c).
404 As shown in Fig 6d, no differences were observed in hepatic *Cidec/Fsp27 β* expression
405 between sexes when using homozygous *Pgc1a*-deficient mice. However, the sex
406 differences at the CIDEC protein levels in lipid droplets and microsomes remained in
407 absence of PGC1a (Fig 6e and f). These results suggest that absence of PGC1A
408 abolishes the sex-induced mRNA changes of hepatic *Fsp27 β* expression in response to
409 WD, being the transcription factor a transcriptional repressor. However, the sex-induced
410 differences in CIDEC present in lipid droplets and microsomes are independent of
411 PGC1A.

412

413

414 **Discussion**

415 The present work explores the putative hepatic *Cidec/Fsp27 β* transcriptional
416 changes induced by dietary components and sex. Using *Apoe*-deficient mouse as a
417 model of hepatic steatosis, dietary cholesterol increased hepatic *Cidec/Fsp27 β* , which
418 was repressed when combined with saturated fat. The latter was not influenced by
419 dietary minor components such as oleanolic acid or squalene administered at
420 pharmacological doses. Moreover, our study revealed a previously unnoticed sex
421 regulation dependent on the prevailing diet, being the female sex a negative regulator.
422 An effect observed in two models of genetic hepatic steatosis (*Apoe*- and *Ldlr*-deficient
423 mice) and reflected in CIDEC/FSP27 content of lipid droplets. Using ovariectomized
424 females, it was shown that ovarian hormones are crucial for the observed decrease in
425 *Cidec/Fsp27 β* expression noted in *Apoe*-deficient mice. This effect was also observed in
426 Wistar female rats. An increased expression of *Pgc1a* inversely associated with that of
427 *Cidec/Fsp27* and the lack of such effect after ovariectomy in *Apoe*-deficient mice allow
428 us to infer that ovarian hormones are executing their action through *Pgc1a*. This was
429 confirmed in mice lacking *Pgc1a* where the sex differences on hepatic *Cidec/Fsp27 β*
430 were erased providing further *in vivo* support for this role. However, the sex differences
431 at the CIDEC/FSP27 content of lipid droplets and microsomes are independent of
432 PGC1a.

433 As shown in Supplemental Figure 8, four set of primers were used to study
434 hepatic *Cidec/Fsp27* m RNA expression in mice. With the exception of primers, named
435 α , corresponding to exon 1, which showed no expression in the liver (data not shown),
436 the remaining three sets gave concordant results in all experimental conditions. None of
437 the selected primers amplified the truncated form. Thus, the observed changes
438 corresponded to *Fsp27 β* , a recently described isoform of the protein regulated by
439 CREBH (64).

440 The present work has explored the influence of two main components of
441 Western diet, cholesterol and saturated fat. Using the first dietary component, an
442 increase in the *Cidec/Fsp27 β* expression was noted. Using information from ENCODE,
443 it was observed that both SREBP1 and 2 bind to this gene (6). Recently, the
444 involvement of SREBP-1c has been proved (7). Surprisingly, the combination of
445 cholesterol and saturated diet decreased hepatic *Cidec/Fsp27 β* expression. In this
446 regard, variable effects of high fat diets have been described depending on the length of
447 fat administration (36). While a short-term administration (3 weeks) increased the
448 expression, a long-term administration of 12 weeks had the opposite effect. In this
449 sense, our study lasted 11 weeks and would be in agreement with the latter finding.
450 Similar results were observed in *ApoE*-deficient mouse males receiving an olive oil-
451 enriched diet (15). Likewise, a decreased expression was found in a postprandial
452 regimen after a virgin olive oil bolus in male Wistar rats and this decrease was inversely
453 associated with hepatic triglyceride and cholesterol contents (32). In the latter case, the
454 hepatic mRNA changes occurred just 4 hours after fat intake. In fasting rats, a rapid
455 increase was equally observed four hours after its start (59). Elevations of *Cidec/Fsp27*
456 mRNA expression by high fat diets required additional dietary deficiencies such those
457 of methionine and choline (Table 2) or under certain metabolic derangements such as
458 those posed by Db mice, PPAR- α -deficient mice (Table 2). In a previous study,
459 using *ApoE*-deficient mice with C57BL/6JxOla129 genetic background and fed Western
460 diets with different conjugated linoleic acid (CLA) isomers, we observed high hepatic
461 *Cidec/Fsp27* mRNA expression in those mice receiving the trans-10,cis-12 CLA isomer
462 and the levels were associated with the hepatic surface occupied by lipid droplets. In
463 contrast, when the cis-9, trans-11 CLA isomer was provided resulted in decreased
464 *Cidec/Fsp27* mRNA expression (15). Overall, regimen of administration and nutritional

465 components are critical modulators of hepatic *Cidec/Fsp27* expression and this mRNA
466 undergoes a rapid metabolic variation in few hours.

467 In the present study, the intake of oleanolic acid, a pentacyclic triterpene, and
468 squalene, a lineal triterpene, had no effect on *Cidec/Fsp27 β* expression despite the
469 changes induced in lipid droplet area (10). Similar finding was reported by the
470 administration of a dietary supplement of *Boswellia serrata*, an extract rich in particular
471 derivatives of boswellic acid, also a pentacyclic triterpene-based compound (20). As
472 triterpenes tend to accumulate in the liver altering distribution of triglycerides in lipid
473 droplets (30, 31), it could be hypothesized that those lipid droplets would not need
474 changes in *Cidec/Fsp27 β* expression or these are not executed at the mRNA level.

475 In a previous study, we observed that hepatic *Cidec/Fsp27* gene expression was
476 significantly associated with hepatic surface occupied by lipid droplets in *ApoE*-
477 deficient mice fed different conjugated linoleic acid isomers, in *Cbs*-deficient mice and
478 in olive oil-fed *ApoE*-deficient mice (15). This was not the case in the present study.
479 Notably, *Cidec/Fsp27 β* expression was associated with hepatic triglyceride (Figures 3
480 and 4) or cholesterol contents (Figures 1 and 2). The genetic background and the diet
481 composition are main differences between the previous and the current study. The
482 former one used Ola129xC57BL/6J mixed genetic background mice while the present
483 study has been carried out using C57BL/6J mice. Due to both strains do have important
484 differences in hepatic fat content (55), the experimental setting may have influenced the
485 outcome. The second aspect is the use of AIN-93 purified diet (48) in the present study
486 compared to commercial ones in the previous one. This choice was forced by the high
487 variability noted in our lab among control mice for years in atherosclerotic lesions when
488 using commercial chows and the failure of obtaining the same batch throughout years.
489 Indeed, source of protein has also been shown to induce changes in *Cidec/Fsp27*

490 expression (61, 65). By and large, dietary components are an important source of
491 variation (50), and our current study, in well-defined conditions of mouse strains and
492 purified diets, adds further evidence supporting this contention.

493 A striking result observed in this work was the decreased hepatic *Cidec/Fsp27 β*
494 expression in female mice consuming WD in *Apoe*- and in *Ldlr*-deficient mice. As
495 consequence of this decrease, the amount of CIDEC/FSP27 protein in lipid droplets was
496 decreased in females. This fact points to a sex-difference in hepatic regulation of lipid
497 droplet enlargement considering the role of CIDEC/FSP27 in this process. An effect
498 that was abolished when ovariectomy was performed in *Apoe*-deficient mice and Wistar
499 rats. Interestingly, female mice lacking steroid receptor coactivator-2 showed increased
500 hepatic expression of this gene (Table 2). Steroid receptor coactivator-2 promotes the
501 transcriptional activation of estrogen receptor in some tissues (62). These results are
502 indicating a negative regulation of the gene by the influence of female hormones. This
503 could be executed through *Pgc1a* as the significant inverse association noted between
504 *Pgc1a* and *Cidec/Fsp27 β* suggests. Further evidences to this suggestion come from the
505 binding of PGC1a to this gene as evidenced by CHIP assays reported by the ENCODE
506 consortium (6). Indeed, estradiol action has been found to be modulated by *Pgc1a* (4)
507 and *Pgc1a* decreased *CIDEC* promoter activity. When we carried out ovariectomy, the
508 decrease in *Cidec/Fsp27 β* expression was lost. Deficiency of *Pgc1a* as the case of the
509 experiment carried out in *Pgc1a*-deficient mice is also supporting the role of *Pgc1a* in
510 the *in vivo* sex differences but only at the mRNA levels. This would be in line with a
511 *Pgc1a*- independent control of downstream processes thereby *Cidec/Fsp27 β* mRNA is
512 translated into FSP27 β protein either at pre- or posttranslational stages. At least in
513 adipocytes, FSP27 protein levels are controlled by the catabolic intervention of
514 proteasomal ubiquitin-dependent proteins. In fact, isoproterenol increases FSP27 levels

515 through a delayed degradation rate mediated by decreased ubiquitination (47), while
516 AMPK activation promoted its degradation (68). Whether or not such mechanisms exist
517 for FSP27 β in the liver are interesting aspects to carry out future experiments.

518 In conclusion, the present report evidences two axes of hepatic *Cidec/Fsp27 β*
519 regulation defined by diet and sex. Regarding the first one, cholesterol and the nature of
520 fatty acids are a key component. On the other hand, the fact that the female decrease in
521 hepatic gene expression was not observed in ovariectomized mice strongly suggests that
522 ovarian hormones are involved in the control of hepatic *Cidec/Fsp27 β* mRNA
523 expression and this is modulated by *Pgc1a*. However, the sex-differences at the
524 CIDEC/FSP27 protein levels observed in lipid droplets and microsomes are
525 independent of PGC1a.

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788 Table 1. Summary of experimental conditions

| Experiment | Genetic background | Diet | Sex | Groups and sample size | Influence |
|-----------------------------------|--------------------|--|---|---|----------------------------------|
| <i>ApoE</i>-deficient mice | | | | | |
| 1 | C57BL/6J x OLA 129 | Commercial chow (B & K Universal Ltd, Humberside, UK) w/wo 0.1% cholesterol for 10 weeks (1) | Males | Control (n=7) Cholesterol (n=7) | Cholesterol |
| 2 | C57BL/6J | Purified chow and Western diets for 11 weeks | Males | Chow (n=13) Western (n=9) | Western diet |
| 3 | C57BL/6J | Purified Western w/wo 0.01% oleanolic acid (OA) (Extrasynthese, Genay, France) for 11 weeks (10) | Males | Western (n=8) Western + OA (n=9) | Oleanolic acid |
| 4 | C57BL/6J | Purified Western w/wo 1% squalene (Sigma, Madrid, Spain) for 10 weeks (14) | Males | Western (n=9) Western + Squalene (n=10) | Squalene |
| 5 | C57BL/6J | Purified chow for 11 weeks | Both sexes | Males (n=13) Females (n=13) | Sex in chow diet |
| 6 | C57BL/6J | Purified Western for 11 weeks | Both sexes | Males (n=9) Females (n=10) | Sex in Western diet |
| 7 | C57BL/6J | Purified Western for 11 weeks | Orchiectomized and non-orchiectomized males | Control (n=9) Orchiectomized on postnatal day 30 (n=9) | Testicular contribution in males |
| 8 | C57BL/6J | Purified Western for 11 weeks | Ovariectomized and non-ovariectomized females | Control (n=9) Ovariectomized on postnatal day 30 (n=9) | Ovarian contribution in females |
| <i>Ldlr</i>-deficient mice | C57BL/6J.S JL | Purified Western for 11 weeks | Both sexes | Males (n=17) Females (n=18) | Sex in Western diet |

| | | | | | |
|------------------------------------|----------|--|---|---|--|
| <i>Pgc1a</i>-deficient mice | C57BL/6J | Purified Western for 11 weeks | Both sexes | Males (n=8) Females (n=8) | Sex in Western diet |
| Rats | Wistar | Purified Western for 100 post-weaning days(38) | Ovariectomized and non-ovariectomized females | Control (n=6) Ovariectomized (n=6) Ovariectomized + a single injection of 1250 µg of testosterone propionate on postnatal day 1 (n=6) | Ovarian contribution and neonatal androgenization in females |

789 w/wo, with or without
790

791 Table 2. Changes in hepatic *Cidec/Fsp27* expression according to Genome Expressed
 792 Omnibus data bank and Array express.

| Experimental condition | Type of change | Signal log ₂ ratio | Accession number |
|---|------------------------------|-------------------------------|------------------|
| Caspase 1 deficient mice | Increased | 0.3 | GDS4922 |
| Glycerol kinase knockout | Increased | 1.9 | GDS1555 |
| NADH-cytochrome P450 reductase deletion effect on liver | Increased | 1.3 | GDS1093 |
| Stearoyl-CoA desaturase 1-deficient mutants on a very low-fat, high-carbohydrate diet | Increased | 2.2 | GDS1517 |
| Steroid receptor coactivator-2-deficient female mice | Increased | 0.8 | GDS4785 |
| Thioredoxin reductase 1-null liver | Increased | 1.1 | GDS4928 |
| Fasting | Increased | 0.5 | GDS4918 |
| Fasting and LPS in male BL6/SV129 mice | Increased | 5.5 | GDS4546 |
| Alcoholic hepatitis | Increased | 0.7 | GDS4389 |
| Sebacic acid supplemented diet effect on db/db liver | Increased | 0.7 | GDS3807 |
| Ketogenic diet effect on the liver | Increased | 5.7 | GDS2738 |
| High-fat high-calorie diet effect on liver | Increased | 1.3 | GDS2413 |
| Liver response to a high fat diet deficient in methionine and choline | Increased | 5.3 | GDS4883 |
| Liver response to a high fat diet: time course | Increased 12 h | 2.4 | GDS4783 |
| Western diet induced changes in liver | Increased | 4.9 | GDS279 |
| Perfluorooctanoic acid effect on livers lacking PPAR-alpha | Increased | 7.1 | GDS3407 |
| Peroxisome proliferator-activated receptor subtype activation effect | Increased by PPAR γ 2 | 0.6 | GDS1373 |

| | | | |
|---|---------------------|------------------------|----------|
| on liver cell | | | |
| Female receiving dexamethasone | Increased | 1.7 | GDS5036 |
| Hepatocyte nuclear factor 4 alpha depletion on hepatocellular carcinoma cell line | Decreased | -0.6 | GDS4798 |
| Transcriptional coactivator PGC-1beta hypomorphic mutation effect on the liver | Decreased | -1.3 | GDS3197 |
| ROR α -deficient staggerer mice fed high fat diet | Decreased | -5.7 | GSE23736 |
| SIRT3 deficient liver response to a high fat diet | Decreased | -0.1 | GDS4817 |
| GPR120-deficient liver response to a high fat diet | Decreased | -1.3 | GDS4830 |
| TAK1/TR4-deficient mice | Decreased | -18 | GSE21903 |
| Conditional GBA1 deletion model of Type 1 Gaucher disease | Decreased | -1.4 | GDS4162 |
| Atherogenic diet effect on the liver: time course | Decreased long term | -4.8 | GDS2292 |
| Streptozotocin induced type 1 diabetes | Decreased | -0.3 | GDS4845 |
| Adrenalectomized liver at light and dark periods of the circadian cycle | Variable | Dark 1.0 Light -1.4 | GDS1870 |
| Sex specific transcription in somatic and reproductive tissues | Decreased | -0.6 | GDS565 |

793 <http://www.ncbi.nlm.nih.gov/gds/>
794 <https://www.ebi.ac.uk/arrayexpress/>.
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797 **Fig. 1. Effect of dietary cholesterol on hepatic steatosis and *Cidec/Fsp27β***
798 **expression in male *ApoE*-deficient mice.** Representative liver micrographs at x600
799 magnification from *ApoE*-deficient mice consuming the chow (a) and cholesterol-
800 enriched (b) diets. Morphometric evaluation of surface of liver section occupied by fat,
801 total cholesterol and triglyceride contents (c). Hepatic *Cidec/Fsp27β* expressions
802 determined by RT-qPCR normalized to *Cyclophilin B* (d). Data are medians and 10-90
803 percentile range for control (n=7) and cholesterol (n=7) groups. Statistical analyses
804 were done according to Mann-Whitney's U test. ^a, P< 0.05 vs control. Association
805 between hepatic cholesterol content and *Cidec/Fsp27β* expression (e). Open squares
806 correspond to control and striped squares to cholesterol-fed mice. Spearman's
807 correlation is shown.

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812 **Fig. 2. Effect of Western diet on hepatic steatosis and *Cidec/Fsp27β* expression in**
813 **male *ApoE*-deficient mice.** Representative liver micrographs at x600 magnification
814 from *ApoE*-deficient mice consuming the chow (a) and Western (b) diets. Morphometric
815 evaluation of surface of hepatocyte occupied by fat and hepatic total cholesterol and
816 triglyceride contents (c). Analysis of hepatic *Cidec/Fsp27β* expression determined by
817 RT-qPCR normalized to *Cyclophilin B* (d). Data are medians and 10-90 percentile range
818 for chow (n=13) and Western (n=9) groups. Statistical analyses were done according to
819 Mann-Whitney's U test. ^a, P< 0.05 vs chow. Association between hepatic cholesterol
820 content and *Cidec/Fsp27β* expression (e). Spearman's correlation is shown. Open
821 squares correspond to control and striped squares to Western-fed mice.

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824 **Fig. 3. Effect of sex on hepatic steatosis, *Cidec/Fsp27β* expression and**
825 **CIDEC/FSP27 content in lipid droplets and microsomes in *ApoE*-deficient mice fed**
826 **on a Western diet.** Representative liver micrographs at x600 magnification from male
827 (a) and female (b) *ApoE*-deficient mice consuming Western diets. Morphometric
828 evaluation of surface of hepatocyte occupied by fat and hepatic total cholesterol and
829 triglyceride contents (c). Analysis of hepatic *Cidec/Fsp27β* expression was determined
830 by RT-qPCR normalized to *Cyclophilin B* (d). Data are medians and 10-90 percentile
831 range for male (n=9) and female (n=10) groups. Relationship between hepatic
832 triglyceride content and *Cidec/Fsp27β* gene expression (e). Open squares correspond to
833 males and striped squares to females. Correlations were calculated according to
834 Spearman's test. FSP27 protein levels normalized to HSC70 in lipid droplets (f) and
835 microsomes (g), inserts show representative Western blots. Statistical analyses were
836 done according to Mann-Whitney's U test. ^a, P< 0.05 vs male.

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843 Fig. 4. **Effect of ovariectomy on hepatic steatosis, *Cidec/Fsp27* expression and**
844 **CIDEC/FSP27 content in lipid droplets and microsomes in female *ApoE*-deficient**
845 **mice fed on a Western diet.** Representative liver micrographs at x600 magnification
846 from mock (a) and surgically castrated (b) female *ApoE*-deficient mice consuming
847 Western diets. Morphometric evaluation of surface of hepatocyte occupied by fat and
848 hepatic total cholesterol and triglyceride contents (c). Analysis of hepatic *Cidec/Fsp27*
849 expression determined by RT-qPCR normalized to *Cyclophilin B* (d). Data are medians
850 and 10-90 percentile range for control (n=9) and castrated (n=9) groups. Relationship
851 between hepatic triglyceride content and *Cidec/Fsp27* gene expression (e). Open
852 squares correspond to control and striped squares to ovariectomized females.
853 Correlations were calculated according to Spearman's test. FSP27 protein levels
854 normalized to HSC70 in lipid droplets (f) and microsomes (g), inserts show
855 representative Western blots. Statistical analyses were done according to Mann-
856 Whitney's U test. ^a, P< 0.05 vs control.

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859 **Fig. 5. Effect of sex and castration on hepatic *Pparg1a/Pgcl1a* expression in *Apoe-***
860 **deficient mice fed on a Western diet.** Influence of sex on hepatic *Pgcl1a* expression in
861 *Apoe*-deficient mice (a). Relationship between *Cidec/Fsp27 β* and *Pgcl1a* gene
862 expression levels (b). Open squares correspond to males and striped squares to females.
863 Correlations were calculated according to Spearman's test. Effect of orchiectomy on
864 hepatic *Pgcl1a* expression in male *Apoe*-deficient mice (c). Effect of ovariectomy on
865 *Pgcl1a* expression in female *Apoe*-deficient mice (d). Analysis of hepatic *Pgcl1a*
866 expression was determined by RT-qPCR normalized to *Cyclophilin B*. Data are medians
867 and 10-90 percentile range for each group. Statistical analyses were done according to
868 Mann-Whitney's U test. ^a, P< 0.05 vs control.

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874 **Fig. 6. Effect of sex on hepatic steatosis, *Cidec/Fsp27* expression and**
875 **CIDEC/FSP27 content in lipid droplets and microsomes in *Pgc1a*-deficient mice**
876 **fed on a Western diet.** Representative liver micrographs at x400 magnification from
877 male (a) and female (b) *Pgc1a*-deficient mice consuming Western diets. Morphometric
878 evaluation of surface of hepatocytes occupied by fat and hepatic total cholesterol and
879 triglyceride contents (c). Analysis of hepatic *Cidec/Fsp27* expression was determined
880 by RT-qPCR normalized to *Cyclophilin B* (d). Data are medians and 10-90 percentile
881 range for male (n=8) and female (n=8) groups. FSP27 protein levels normalized to
882 HSC70 in lipid droplets (e) and microsomes (f), inserts show representative Western
883 blots. Statistical analyses were done according to Mann-Whitney's U test. ^a, P< 0.05 vs
884 male.

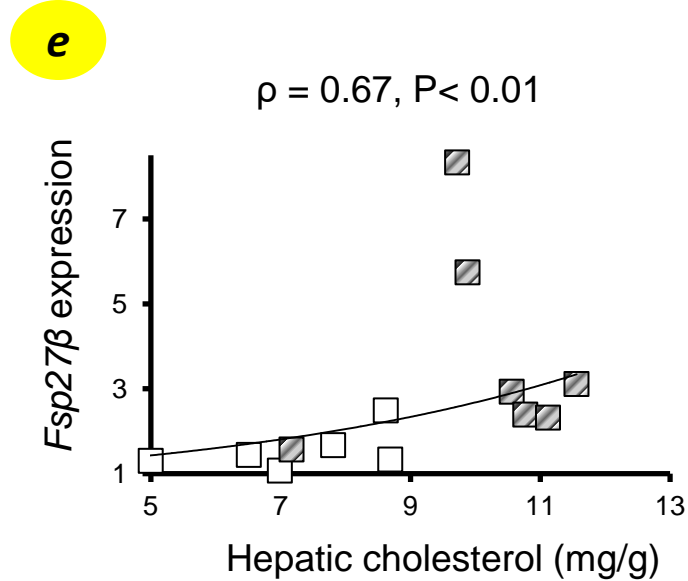
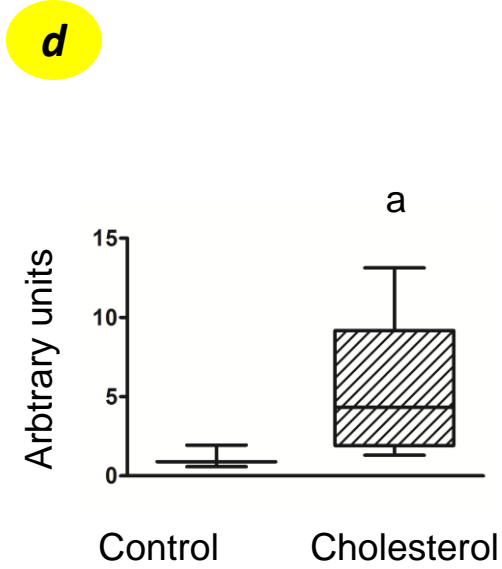
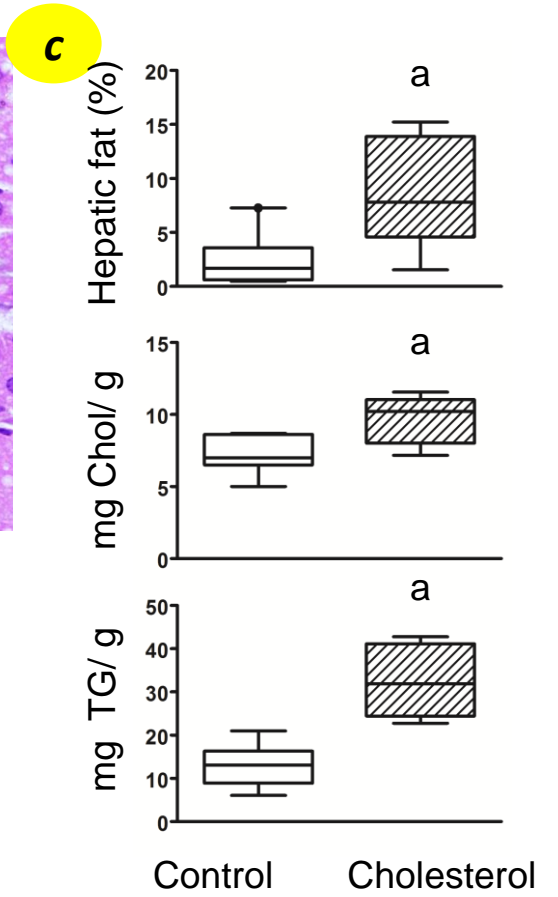
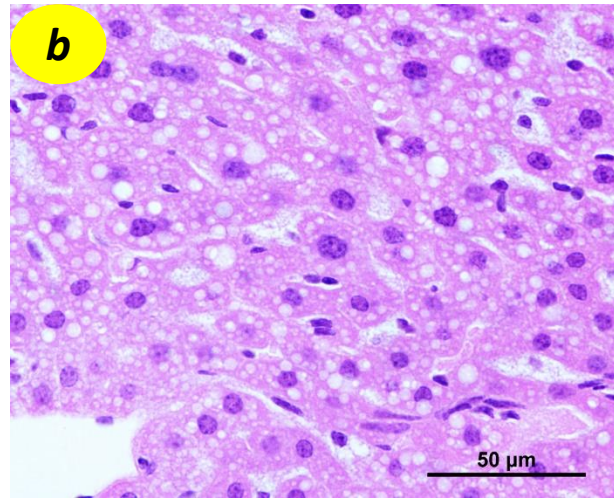
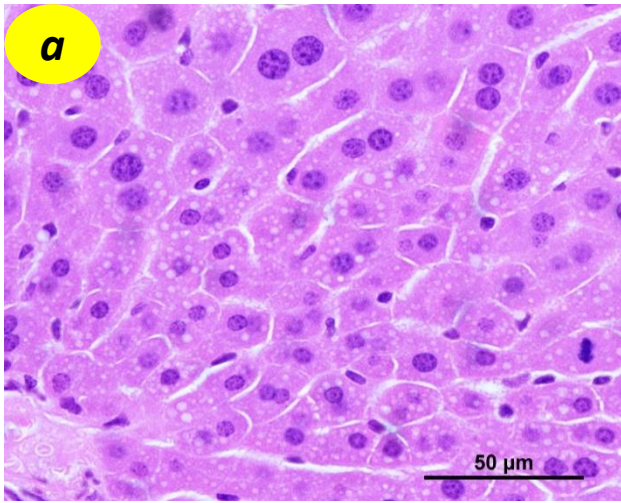
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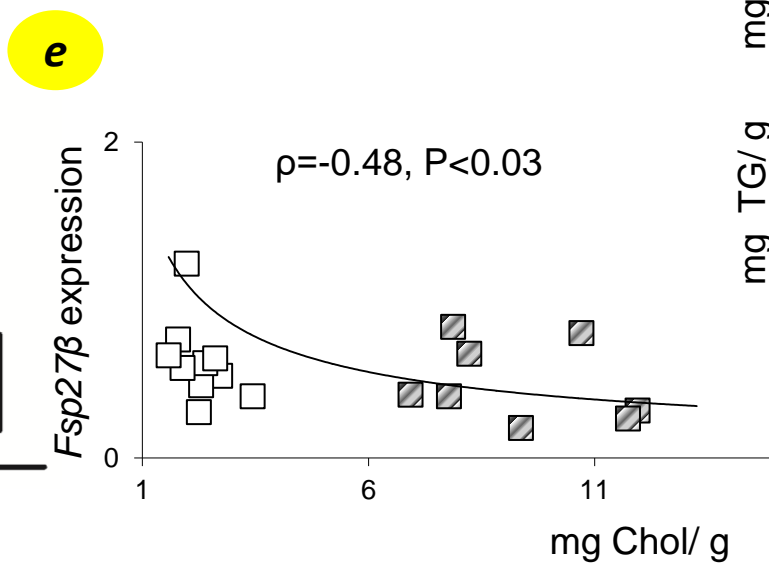
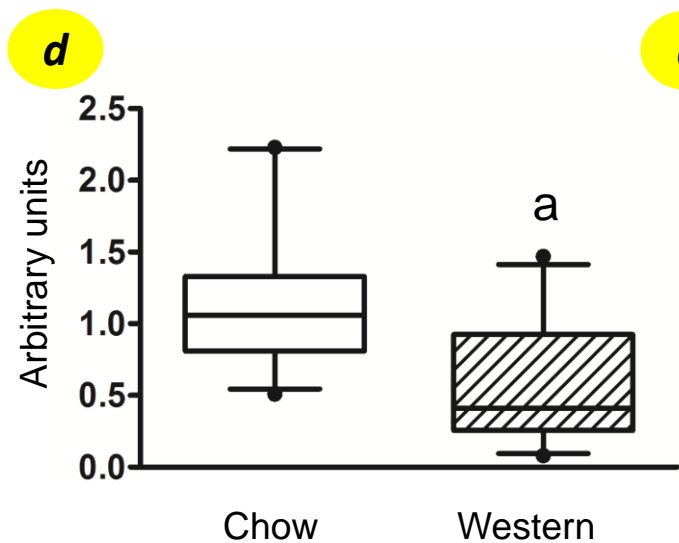
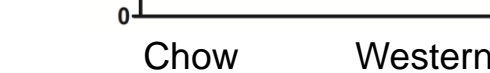
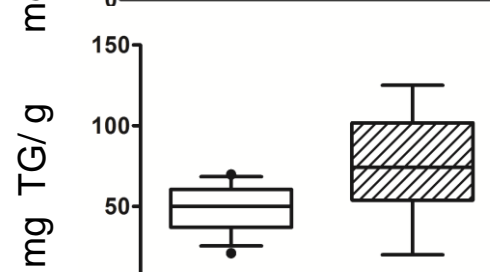
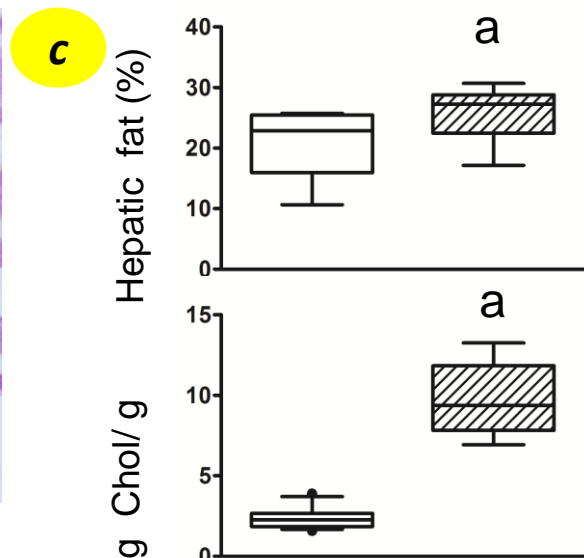
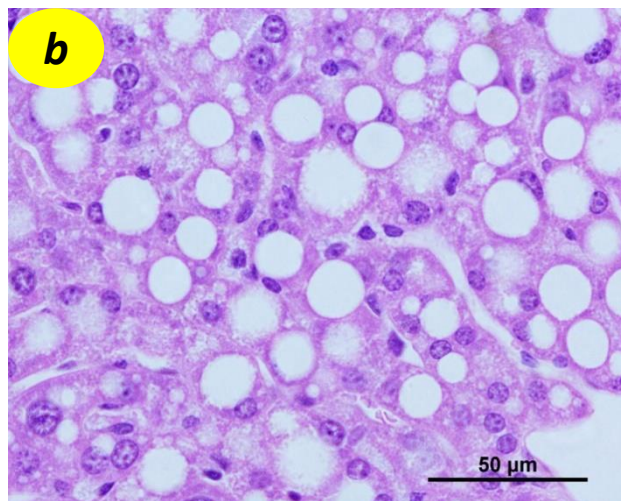
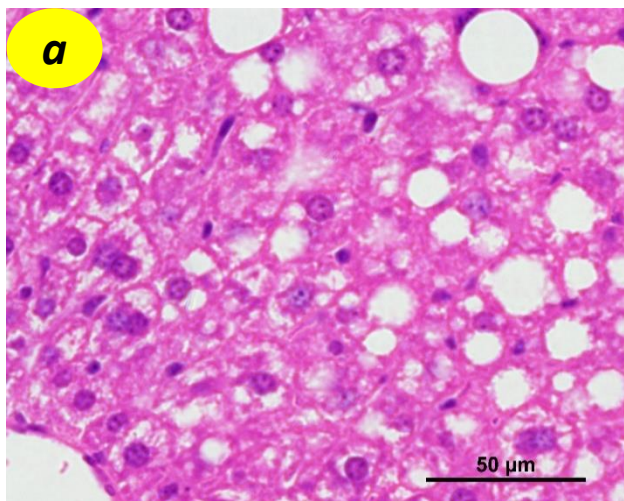
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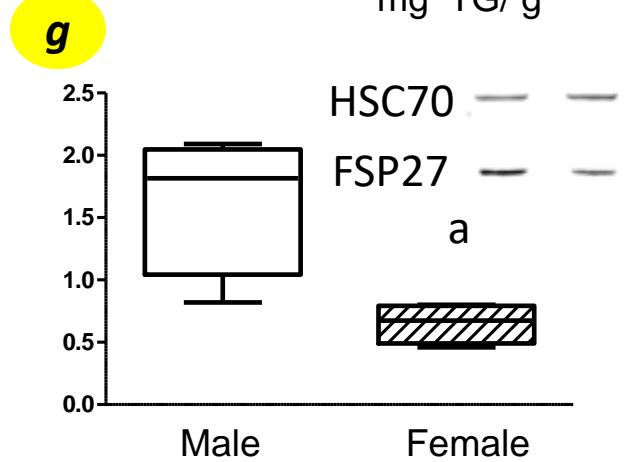
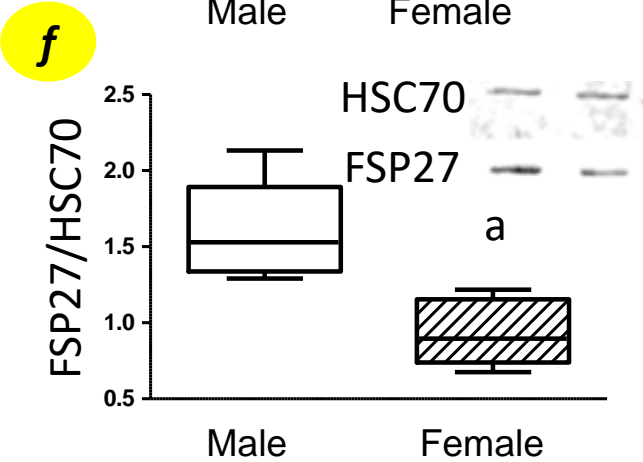
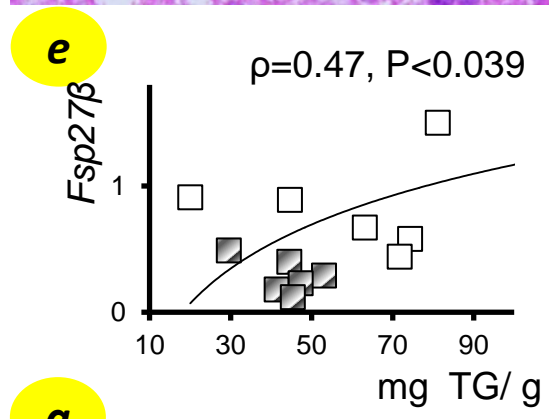
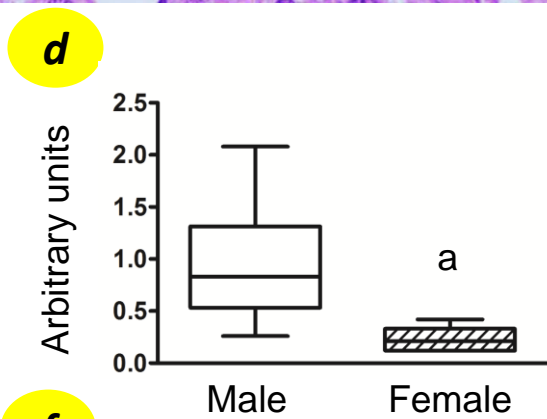
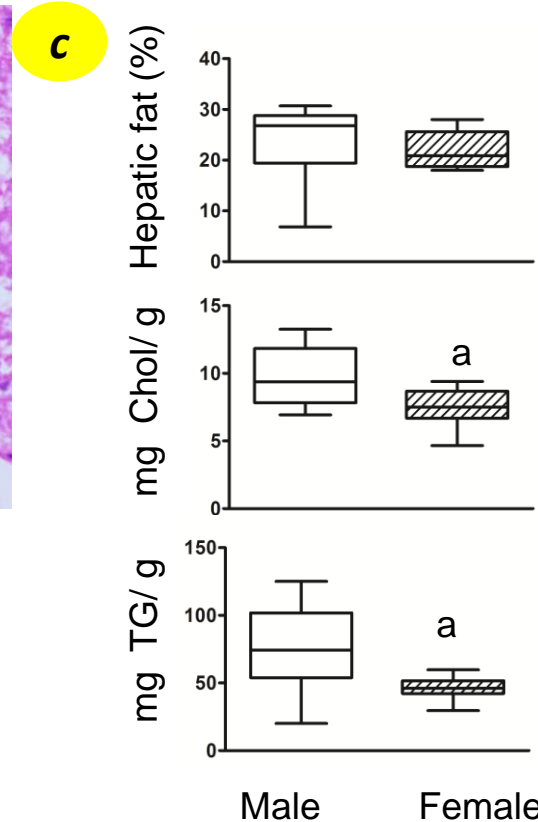
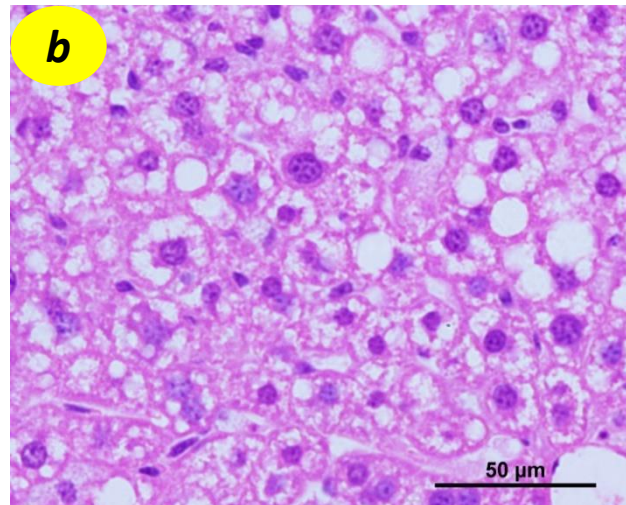
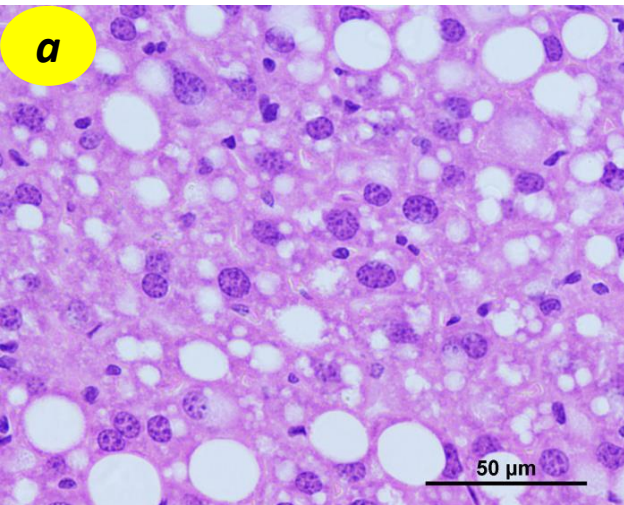
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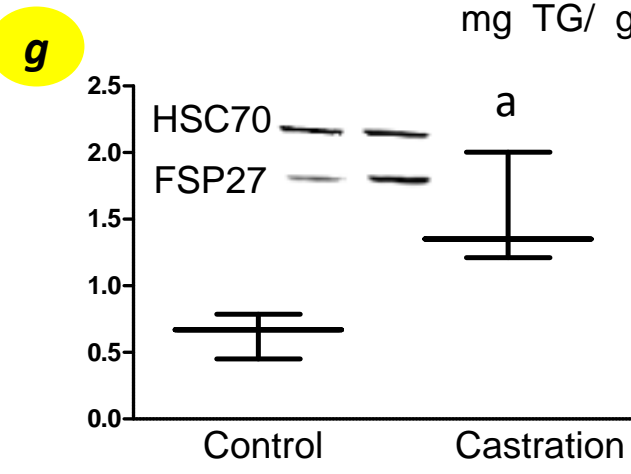
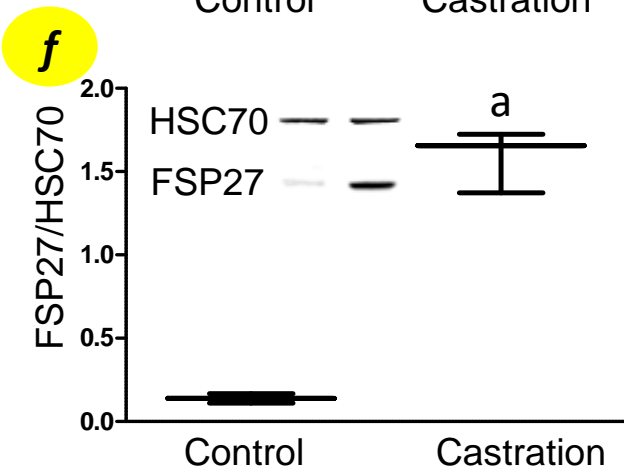
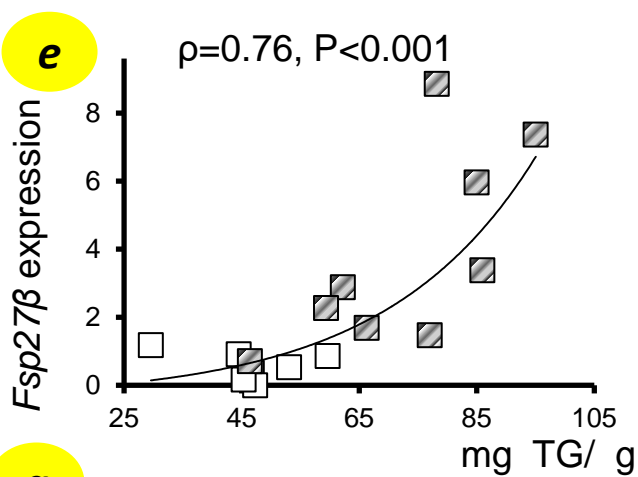
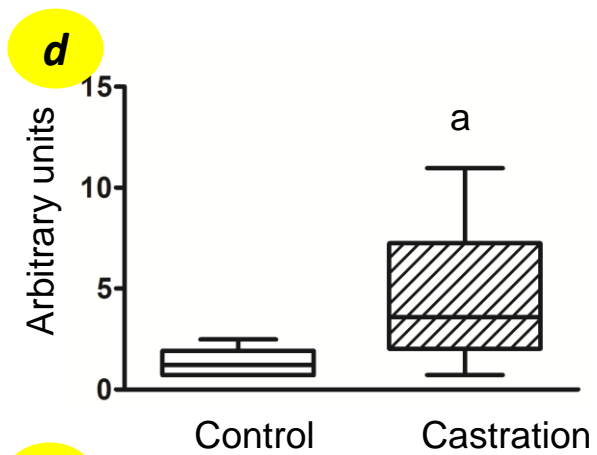
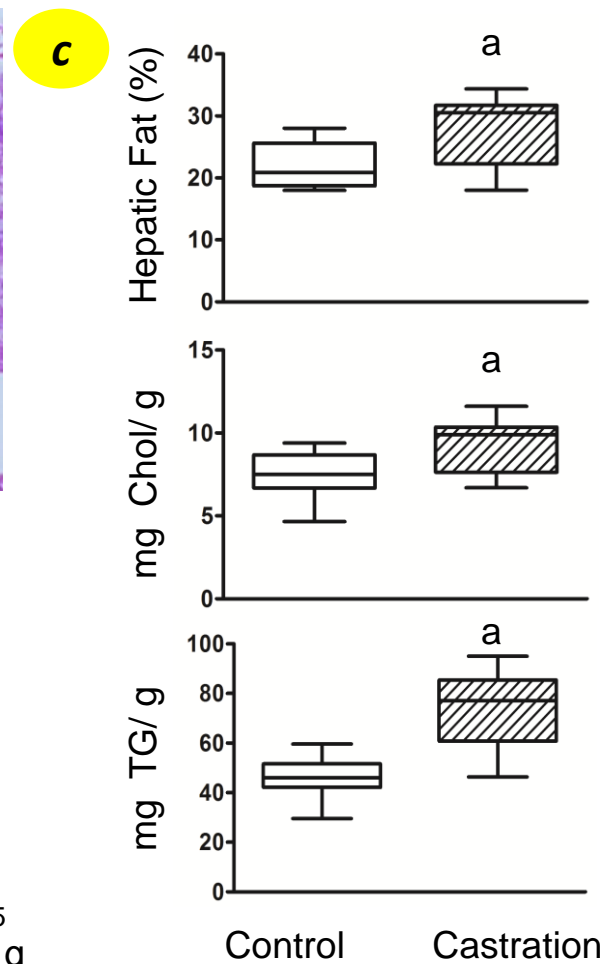
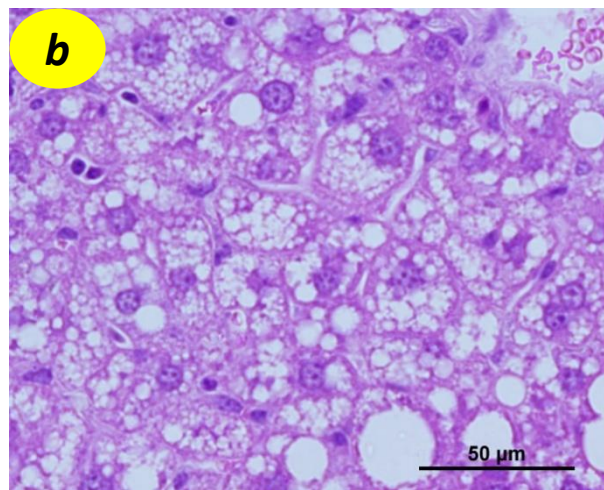
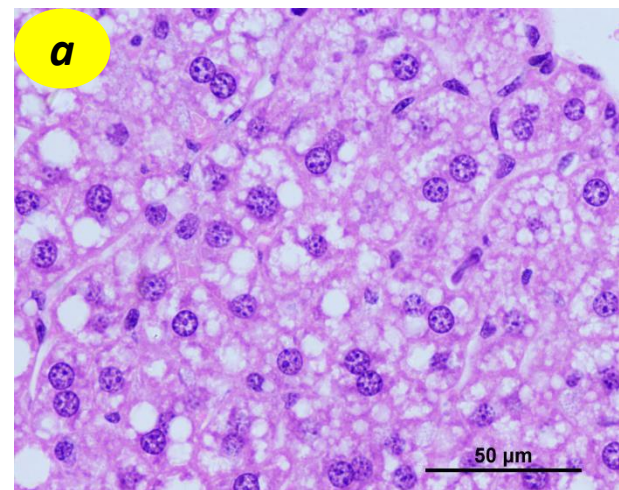
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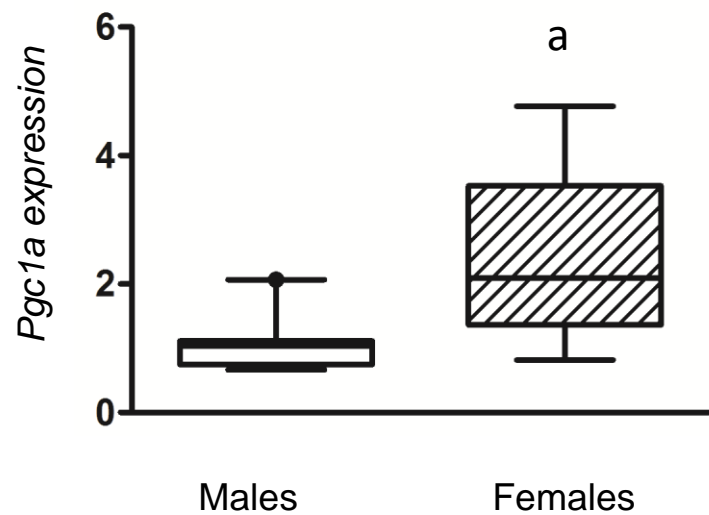
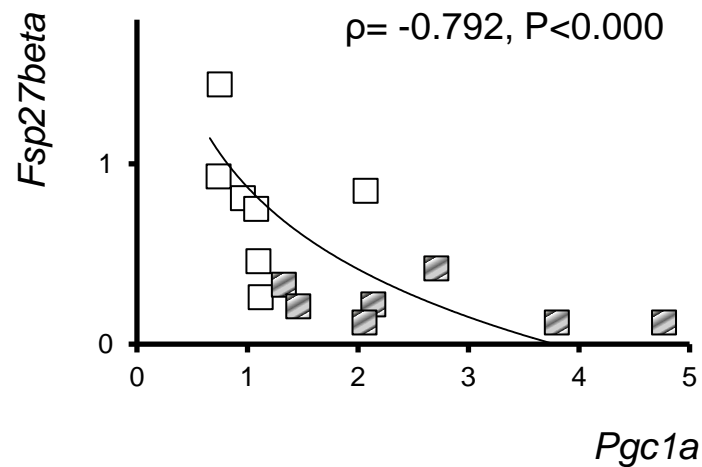
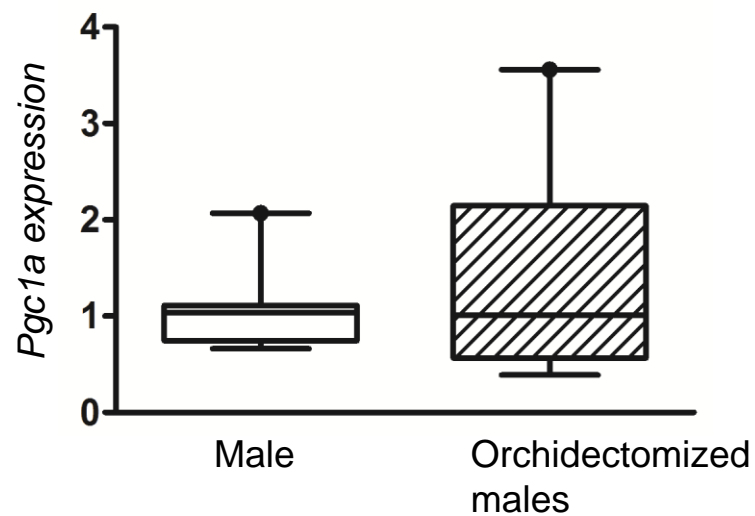
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