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Adrenoceptor-related decrease in serum triglycerides is independent of PPAR α activation

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31 **Key words:** Triglycerides; PPAR α ; Adrenergic receptors; TRLs;
32 hypertriglyceridemia; LDL

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34 **Short title:** Adrenergic regulation of serum triglycerides

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40 **Abbreviations:**

41 AADAC, arylacetamide deacetylase; ACADM, acyl-CoA dehydrogenase; ACOX, acyl-CoA
42 oxidase; ACOT, acyl-CoA thioesterase; AKT, protein kinase B; ALT, alanine
43 aminotransferase; AST, aspartate aminotransferase; ApoE, apolipoprotein E; AR, adrenergic
44 receptor; ATGL, adipose triglyceride lipase; BBAT, bile acid CoA; CREB, cAMP-response
45 element-binding protein; cAMP, cyclic AMP; CD36, cluster of differentiation 36;
46 CES3/TGH, carboxylesterase 3; DGAT, diacylglycerol O-acyltransferase; EIA, Elisa; FFA,
47 free fatty acids; FoxO1, forkhead box protein O1; HNF4 α , hepatocyte nuclear factor 4 α ;
48 HDL, high density lipoprotein; HSL, hormone sensitive lipase; ISOP, isoprenaline; MTTP,
49 microsomal triglyceride transfer protein; NEFA, non-esterified fatty acids; Nr4A, nuclear
50 receptor 4 α ; LPL, lipoprotein lipase; LDL-r, low density lipoprotein receptor; VLDL, very
51 low density lipoprotein; PCSK9, proprotein convertase subtilisin/kexin type 9; PH,
52 phenylephrine; PI3k, phosphatidylinositol 3-kinase; PKA, protein kinase A; PPAR α ,
53 peroxisome proliferator-activated receptor- α ; PCR, polymerase chain reaction; RIA,
54 radioimmunoassay; TG, triglycerides; TRLs, triglyceride-rich lipoproteins; W.A.T., white
55 adipose tissue.

56 **ABSTRACT**

57 Adrenoceptor (AR)-linked pathways belong to the major components of the stress
58 response system and are associated with the pathophysiology of diseases within the
59 spectrum of metabolic syndrome. In this study, the role of adrenoceptor stimulation in
60 serum triglyceride (TG) regulation in mice was investigated. For this purpose, α_1 -ARs
61 were activated with phenylephrine (PH) and $\beta_{1/2}$ -ARs with isoprenaline (ISOP). Both
62 AR-agonists markedly reduced serum TG levels independently of PPAR α activation.

63 These drugs also significantly activated the hormone sensitive lipase in the white
64 adipose tissue indicating increased mobilization of TGs in this tissue. In addition, PH
65 and ISOP up-regulated Lpl, Nr4A, Dgat1, Mttp, Aadac and Cd36 genes, critical in TG
66 regulation, whereas the observed decrease in serum TG levels was independent of the
67 hepatic very low-density lipoprotein (VLDL)-TG secretion. Interestingly, PH and
68 ISOP also inactivated the hepatic insulin/PI3k/AKT/FoxO1 signaling pathway,
69 holding a critical role in the regulation of genes involved in TG synthesis. Taken
70 together, the findings of the present study indicate that stimulation of α_1 - and $\beta_{1/2}$ -ARs
71 markedly reduced serum TG steady state levels as a result of alterations in TG
72 synthesis, uptake, transport, hydrolysis, metabolism and clearance, an effect induced
73 by PPAR α independent mechanisms.

81 Introduction

83 Hypertriglyceridemia is a major pathological feature of metabolic syndrome, which is
84 associated with accumulation of triglyceride-rich lipoproteins (TRLs) in circulation.
85 Patients with elevated serum TRLs are at high risk for cardiovascular and renal
86 disease, as well as for steatohepatitis and other disorders. To this date, treatment with
87 fibrates is the most effective pharmacological approach in clinical practice for the
88 reduction of serum TG levels. Fibrates are used either as monotherapy or in
89 combination with statins and other hypolipidemic drugs [1, 2].

90 Fibrates are ligands for the peroxisome proliferator-activated receptor α
91 (PPAR α), which is activated by psychophysiological stress via stimulation of AR-
92 linked pathways and glucocorticoids [3, 4]. PPAR α acts as a cellular “lipostat” that
93 transduces alterations in cellular lipid levels to the transcriptional regulation of
94 various target genes, which are critical for the fate of fatty acids [5-7]. In particular,
95 activation of PPAR α up-regulates a broad array of genes encoding enzymes that are

96 involved in fatty acid uptake, transport, as well as in mitochondrial and peroxisomal
97 fatty acid β -oxidation and microsomal fatty acid ω -oxidation. In addition, several
98 apolipoproteins are regulated by PPAR α including apolipoproteins (Apo) AI, AII and
99 CIII, a fact that indicates the central role of PPAR α in the extracellular transport and
100 metabolism of TG-rich lipoproteins in blood [3]. These PPAR α -mediated changes in
101 gene expression result in reduced serum TG-rich lipoproteins and increased high
102 density lipoprotein (HDL) levels [8, 9], although the exact mechanisms that link TG
103 and HDL levels are currently poorly defined.

104 The apparent causative relationship between serum TRL levels and a wide
105 range of human pathologies has triggered the development of several biological drugs
106 targeting TRL metabolism, such as Volanesorsen, [10], Evinacumab [11-13] and
107 IONIS-ANGPTL3-LRx [14]. Nonetheless, the effective prevention of
108 hypertriglyceridemia requires a deeper understanding of the biochemical mechanisms
109 involved and more precisely, the triggers leading to excess TRL accumulation in
110 serum.

111 The role of stress in the regulation of lipid homeostasis is well documented. In
112 particular, chronic stress deregulates lipid and carbohydrate homeostasis and is
113 considered as a causative factor of several pathologies related to the metabolic
114 syndrome, such as visceral obesity, insulin resistance, dyslipidemia, dyscoagulation
115 and hypertension [15-23]. It has been also reported that humans with low sympathetic
116 nervous system (SNS) activity, reduced beta-adrenergic sensitivity and lipid
117 mobilizing efficacy of catecholamines display lowered energy expenditure and are at
118 high risk to develop obesity compared to physiological subjects. Therefore, adrenergic
119 receptors, major components of the SNS, have been considered as putative therapeutic
120 targets against obesity [24]. Accumulating evidence also suggests that short-term
121 exposure to stress has a beneficial effect on TG regulation. Specifically, subacute
122 exposure to repeated restraint stress markedly reduces serum TG steady-state levels,
123 predominantly via adrenergic receptor (AR)-linked pathways. In particular, α_1 - and
124 $\beta_{1/2}$ -ARs appear to hold major roles in this regulation, as blockade of these receptors
125 prior to stress completely inhibited the suppressive effect of stress on serum TG levels
126 [4]. Activation of the hormone sensitive lipase (HSL) in the white adipose tissue
127 (W.A.T.) by stress or epinephrine, a major effector of the stress response, is
128 potentially responsible, at least in part, for this suppressive effect. In addition, stress
129 via stimulation of α_1 - and $\beta_{1/2}$ -ARs up-regulated several genes in the W.A.T., which

130 are critical in the synthesis and metabolism of TG depots, such as the diacylglycerol
131 acyltransferase (*Dgat*)1 and 2, lipoprotein lipase (*Lpl*), adipose triglyceride
132 lipase/patatin-like phospholipase domain containing 2 (*Atgl/Pnpla2*), arylacetamide
133 deacetylase (*Aadac*), microsomal triglyceride transfer protein (*Mttp*) and the orphan
134 nuclear receptor (*Nr4A*) [2, 25-32].

135 To better understand the molecular mediators of the stress-related
136 hypertriglyceridemia, in this study we investigated the involvement of α_1 - and $\beta_{1/2}$ -
137 ARs in the regulation of serum TRL homeostasis. For this purpose, pharmacological
138 manipulations of α_1 - and $\beta_{1/2}$ -AR-linked pathways by phenylephrine (PH) and
139 isoprenaline (ISOP) respectively, were used. The data revealed a strong suppressive
140 effect of the $\beta_{1/2}$ -AR agonist and less of the α_1 -AR agonist on serum TG steady-state
141 levels, independent of PPAR α activation, shedding light in novel-signaling pathways
142 triggered by the adrenergic system with significant roles in TRL homeostasis.

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145

146 **Results**

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149 **Alterations in serum lipid levels, post-prandial triglyceride kinetics, and hepatic** 150 **VLDL triglyceride secretion**

151

152 Pharmacological stimulation of α_1 -ARs markedly reduced serum TG, free fatty acid
153 (FFA) and total cholesterol levels in wild-type mice (Fig. 1A-1C). Beta-AR
154 stimulation decreased only serum TG and FFA concentration, whereas it had no effect
155 on total cholesterol levels (Fig. 1A-1C). Interestingly, stimulation of ARs also
156 suppressed serum TG steady state levels in *Ppara*-null mice thus indicating a PPAR α -
157 independent mechanism in TG regulation by PH and ISOP (Fig. 1A). In an effort to
158 provide a mechanistic interpretation of the observed reduction in serum TG levels
159 following stimulation of ARs with AR-agonists, the hepatic VLDL-TG secretion
160 kinetics was determined in control and AR-agonist-treated *Ppara*-null mice.
161 Stimulation of β_1/β_2 -ARs with ISOP resulted in a significant increase in the rate of
162 hepatic VLDL-TG secretion in treated mice when compared to controls (Fig. 2A),

163 while PH (α_1 -AR agonist) did not have any significant effect (Fig. 2A), suggesting
164 that mobilization and secretion of hepatic TG into VLDL particles could not account
165 for the observed reduction of serum TG levels following PH treatment (Fig. 1A &
166 2A). Similarly, no significant changes in serum AST, ALT and body weight levels
167 were observed following the above mentioned drug treatments (Tables 1 and 2).

168

169 **In vivo assessment of the role of AR-related pathways in TG regulation**

170

171 To further elucidate the mechanisms underlying the strong suppressive effect of AR-
172 linked pathways on serum TG steady state levels, the expression of various genes
173 encoding factors involved in TG synthesis, metabolism and clearance were
174 determined by qPCR and western blot analysis. Both PH and ISOP increased hepatic
175 *Lpl* mRNA expression (Fig. 3A). *Nr4A* mRNA expression was also increased by PH
176 and ISOP in the liver (Fig. 3A). In contrast, the expression of *Atgl/Pnpla2*, *Hsl* and
177 *Aadac* mRNAs were suppressed by PH in this tissue (Fig. 3A and 3B). *Dgat2* mRNA
178 transcripts were also increased in the liver of PH- and ISOP-treated mice (Fig. 3B),
179 whereas *Dgat1* mRNA expression was not affected (Fig. 3A). Moreover, *Mttp*,
180 carboxylesterase 3 (*Ces3/tgh*) and cluster of differentiation 36 (*Cd36*) mRNAs were
181 increased to the same extent in the liver of both, PH- and ISOP-treated mice compared
182 to placebo treated animals (Fig. 3B). No effect was observed on hepatic low-density
183 lipoprotein receptor (*Ldl-r*) mRNA expression (Fig. 3C). Similarly, APOE protein
184 expression was not affected by either AR-agonists (Fig. 3C).

185 It is of interest to note that the AR-agonists, PH and ISOP, also up-regulated
186 Nr4A in the W.A.T., which may in turn trigger the up-regulation of *Lpl* (Fig. 4A). *Hsl*
187 and *Atgl/Pnpla2* mRNA and protein were not affected by either PH or ISOP in this
188 tissue (Fig. 4A and 4D). PH though, induced HSL phosphorylation at Ser563 in the
189 W.A.T. compared to controls (Fig. 4D), whereas ISOP increased HSL
190 phosphorylation at Ser660 in this tissue (Fig. 4D). Notably, total perilipin and
191 specifically, PLIN5 protein levels were not modified by the AR-agonists in the
192 W.A.T. (Fig. 4D). *Dgat1* mRNA expression was increased only by ISOP in the
193 W.A.T. (Fig. 4A), whereas *Mttp* and *Cd36* were up-regulated by both drugs (Fig. 4B).
194 *Ces3/tgh* mRNA expression was not affected (Fig. 4B). Interestingly, TG content in
195 the W.A.T. was lower in PH- and ISOP-treated mice compared to controls (Fig. 1C).

196 In contrast, TG levels were higher in the livers of ISOP-treated mice compared to
197 controls and PH-treated animals (Fig. 2C).

198 In order to determine the role of AR-agonists in lipid β -oxidation, the effect of
199 PH and ISOP on mRNA encoding ACADM, the rate-limiting enzyme of this reaction,
200 was assessed in the liver and W.A.T. using qPCR analysis. Only stimulation of α_1 -
201 ARs with PH markedly increased *Acadm* mRNA expression in the liver compared to
202 controls; ISOP had no effect (Fig. 3C). In contrast, PH repressed *Acadm* mRNA
203 expression in the W.A.T. (Fig. 4C).

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208 **In vivo and in vitro assessment of the AR-induced alterations in Hnf4 α** 209 **regulation**

210

211 Stimulation of α_1 - or $\beta_{1/2}$ -ARs with PH or ISOP, respectively, markedly increased
212 *Hnf4a* mRNA and HNF4 α protein levels (Fig. 5A). The AR-induced *Hnf4a*
213 expression triggered the up-regulation of the Hnf4 α target genes, *Cyp8b1* and bile
214 acid CoA: amino acid N-acyltransferase (*Baat*) (Fig. 5B). Further investigation
215 revealed that the drug-induced up-regulating effect on hepatic Hnf4 α is due to a direct
216 effect of the drug on hepatocyte α_1 - or β_1 -ARs, respectively. Treatment of primary
217 hepatocytes with either PH or ISOP markedly induced hepatocyte *Hnf4a* mRNA
218 expression (Fig. 5C). This up-regulating effect on Hnf4 α was blocked by pre-
219 treatment of the cells with the PKA inhibitor, H89, and the phosphatase- and ATPase
220 inhibitor, NaOV (Fig. 5C). The ISOP-induced Hnf4 α up-regulation was also
221 prevented mainly, by the phosphatase- and ATPase inhibitor, NaOV and to a lesser
222 extent by H89 (Fig. 5C).

223

224 **In vivo assessment of the role of AR-linked pathways in PI3k/AKT/FoxO1 and** 225 **cAMP/PKA activation**

226

227 In order to further investigate the mechanism underlying the reduction in serum TG
228 levels following PH or ISOP treatment, total cellular proteins were analyzed by
229 Western blot. When compared to controls, both, PH and ISOP reduced AKT and

230 consequently, FoxO1 phosphorylation in the liver (Fig. 6), whereas they increased
231 CREB phosphorylation (Fig. 6), indicating inactivation of the PI3k/AKT/FoxO1 and
232 activation of the AR/cAMP/PKA/CREB signalling pathway.

233 Discussion

234

235

236 Accumulated experience over the past many decades of basic and clinical research
237 established unequivocally a major role of LDL-C in the development and progression
238 of atherosclerosis. However, the aggressive LDL-C lowering in patients following the
239 current medications is still associated with substantial residual cardiovascular risk,
240 and strongly suggests that the benefit from LDL-C lowering strategies has reached a
241 plateau [33]. Identifying and targeting alternative processes that are highly associated
242 with atherogenesis may provide new ways to complement existing therapies and
243 augment their benefit against the development of diseases, thus further reducing the
244 residual cardiovascular risk, which is associated with the current pharmacotherapy
245 [33].

246 The apparent causative relationship between high TRL serum levels and
247 atherosclerosis led to the development of several investigational drugs currently in
248 clinical trials, that target TRL metabolism [33]. Volanesorsen, an apolipoprotein C3
249 (Apo C3) antisense oligonucleotide, targets selectively Apo C3 mRNA and blocks
250 protein synthesis, due to the enhanced ribonuclease H1-mediated degradation of Apo
251 C3 mRNA [10]. Another experimental drug is Evinacumab, an angiopoietin-like
252 protein 3 (ANGPTL3) monoclonal antibody that blocks ANGPTL3, a protein known
253 to increase plasma TRL and TG levels [11, 34, 35]. Another similar ANGPTL3-
254 targeting drug is the IONIS-ANGPTL3-LRx, an ANGPTL3 antisense oligonucleotide.
255 Despite these developments, the molecular triggers that are associated with the
256 disease development and eventually, with the elevated plasma TRL accumulation
257 remain largely unexplored. There is a strong evidence that both, central and peripheral
258 nervous systems, may be involved in this regulation [4, 36-38].

259 Pharmacological stimulation of α_1 - or $\beta_{1/2}$ -AR linked pathways markedly
260 reduce the steady state levels of serum TG in mice. These data are in line with studies
261 reporting that subacute repeated restraint stress via mainly AR-related pathways
262 reduces serum TG levels [4]. Investigation of the potential mechanisms involved in

263 the AR-agonist-induced serum TG reduction indicated that PPAR α activation is not a
264 part of this mechanism, because treatment of *Ppara*-null mice with either PH or ISOP
265 triggered a reduction in serum TG levels that was comparable with that observed in
266 PPAR α expressing mice. Therefore, fibrates may not be an effective therapy for the
267 stress-related hypertriglyceridemia. The contribution of other molecular factors, such
268 as APO C3 and ANGPTL3, needs to be investigated.

269 The significant reduction of serum FFA observed following treatment with PH
270 and ISOP may also suggest that AR-agonists potentially increase energy requirements
271 in the treated mice. This hypothesis is supported by previous studies reporting that
272 agents, which stimulate adrenergic neurons increase energy expenditure, lipolysis and
273 fat oxidation [24]. Free fatty acids derived from TG β -oxidation are a major source of
274 energy. It is plausible that the rapid reduction in serum TG levels observed in mice
275 following treatment with AR-agonists represents an immediate uptake of plasma
276 TRLs by energy craving tissues in treated mice. Circulating TRLs serve as an
277 immediate source of FFA. However, since circulating TRLs represent a limited
278 supply, HSL activity needs to be stimulated in order to mobilize additional
279 intracellular deposits of TGs for sustained energy production in the W.A.T..

280 In addition, given the complex and multifactorial regulation of TG
281 homeostasis, it is possible that AR-stimulation by PH and ISOP influences numerous
282 and diverse processes responsible for the observed reduction in serum TG levels. For
283 example, PH and ISOP treatment could affect dietary lipid absorption, their packaging
284 into chylomicrons, the processing of these chylomicrons in plasma via lipoprotein
285 lipase and their subsequent clearance from the circulation by the LDL-r, the tissue
286 deposition and mobilization of these TGs once they reach the respective tissues, their
287 combustion via β -oxidation of fatty acids, and their shuttling between VLDL/LDL
288 and HDL via CETP [25, 26, 30]. The precise effects of AR-stimulation on these
289 mechanisms need further investigation.

290 The present data indicated that stimulation of α_1 - or $\beta_{1/2}$ -ARs resulted in the
291 up-regulation of several genes holding determinant roles in the fate of TGs [12, 13,
292 27, 28, 39, 40]. In particular, AR-agonists stimulated the hepatic expression of genes
293 encoding factors involved in TG metabolism and clearance, including *Lpl*, *Nr4A*,
294 *Mttp*, *Dgat2*, *Ces3/Tgh* and *Cd36* [27, 28]. These genes, with the exception of
295 *Ces3/Tgh* and *Dgat2*, were also increased in the W.A.T.. *Dgat1* was up-regulated only
296 by ISOP and *Aadac* only by PH in the W.A.T.. It is of interest also to note that both

297 AR-agonists activated HSL in the W.A.T. and reduced TG concentration in this
298 tissue, indicating an increased TG hydrolysis rate [12, 13, 27, 28, 39-41]. Treatment
299 of mice with either PH or ISOP promotes a considerable decrease in serum TG levels.
300 Notably, ISOP results in a more significant effect. Real-time PCR analysis indicated
301 that both agonists induce *Lpl* expression, indicating that the suppressive effect of
302 ISOP on serum TG levels could be mainly due to increased *Lpl* expression; the
303 nuclear receptor NR4A may have triggered the ISOP-induced *Lpl* up-regulation,
304 while the effect of PH on TG is mediated by downstream to LPL events involved in
305 the clearance of TG rich lipoproteins, such as enhanced holoparticle uptake by the
306 LDL-r [26]. ISOP treatment significantly increased hepatic VLDL-TG secretion,
307 while PH had no effect, suggesting that mobilization and secretion of hepatic TGs is
308 not a factor in the observed decrease of serum TG levels shown in PH-treated mice.

309 Although the above findings are strongly indicative for the role of AR-agonists
310 in the regulation of factors determining the fate of TGs in the body, future studies
311 should be designed to pinpoint the AR-agonist-induced alterations in the afore-
312 mentioned proteins, focusing mainly in alterations at enzyme activity levels. It is well-
313 established that HSL activity is highly regulated by adrenergic stimulation followed
314 by PKA and AMPK activation [42]. In cases where catecholamines are
315 physiologically elevated in humans (i.e. during physical exercise), the level of HSL
316 phosphorylation at Ser563 and Ser660 (PKA regulatory sites) is increased in both,
317 skeletal muscles and adipose tissue. This induced phosphorylation results in HSL
318 activation. FFA coming from the enzymatic lipolysis of W.A.T triglycerides enter to
319 hepatocytes where they are converted into triglycerides that will be eventually
320 incorporated into nascent VLDL particles [43]. Epinephrine, an α/β -AR agonist, is
321 known to induce phosphorylation of HSL at Ser563 and Ser660 to the same extent.
322 Our present data indicate that PH and ISOP differ from epinephrine in that they
323 selectively promote phosphorylation of either sites. Specifically, although PH induced
324 HSL phosphorylation at Ser563 in the W.A.T. compared to controls (Fig. 4D), ISOP
325 increased HSL phosphorylation at Ser660 in this tissue (Fig. 4D). Given that
326 phosphorylation at both residues is required for a significant induction of the HSL
327 activity, and based on the VLDL-TG secretion data (Fig. 2), we hypothesize that the
328 PH-induced Ser563 phosphorylation may be a weaker inducer of HSL activity, thus
329 resulting to less FFAs available for VLDL production compared to those following
330 the ISOP-induced Ser660 phosphorylation (Fig. 7). In support of our hypothesis is the

331 report that epinephrine (α/β -AR-agonist), also activates the AMP-activated kinase
332 (AMPK), which is considered to block the PKA-dependent activation of HSL in
333 adipocytes, when HSL phosphorylation occurs at Ser563, while it is preserved when it
334 occurs at Ser660. Our data are in line with previous findings indicating the lesser
335 significance of α_1 -ARs in the HSL-dependent lipolysis in adipocytes compared to that
336 of β -ARs [42, 44]. Clearly, additional enzymatic studies are essential to verify this
337 hypothesis.

338 The increased fatty acid β -oxidation in the hepatic mitochondria also
339 profoundly contributes to the α_1 -AR-induced decline in serum TG steady state levels,
340 as PH led to an up-regulation of hepatic *Acadm* that encodes the rate limiting enzyme
341 in this metabolic pathway [12]. In the W.A.T., fatty acid β -oxidation does not appear
342 to participate in PH- and ISOP-induced decline of serum TG levels as both AR-
343 agonists had no effect on *Ppara* and *Ppar γ* expression, whereas PH repressed *Acadm*
344 in this tissue (Fig. 4C).

345 Notably, PH and ISOP significantly up-regulated hepatic *Hnf4 α* that holds
346 determinant roles in a regulatory network required for the maintenance of the
347 hepatocyte phenotype and the regulation of several metabolic genes involved in lipid
348 homeostasis. It is of interest also to note that *Hnf4 α* is acting in a coordinating fashion
349 with the transcription regulators, *Ppara* and *Ppar γ* , on their downstream target genes
350 encoding factors important in fatty acid metabolism [9, 12, 45]. In both cases of AR-
351 agonists, the *Hnf4 α* induced expression appears to be mediated by activation of
352 several phosphatase- and ATPase-linked signaling pathways, as pre-treatment of
353 hepatocytes with the inhibitor of these enzymes, NaOV, drastically prevented the up-
354 regulating effect of PH and ISOP on *Hnf4 α* . The involvement of the α_1/β -
355 AR/cAMP/PKA signaling pathway in this induction is also indicated by the fact that
356 the PKA inhibitor, H89, restricted the drug-induced effect on *Hnf4 α* .

357 It is well documented that the insulin/PI3k/AKT/FoxO1 signaling pathway
358 regulates several lipogenic genes involved in TG synthesis [46, 47]. Inactivation of
359 this signaling pathway was detected following stimulation of α_1 -ARs with PH or $\beta_{1/2}$ -
360 ARs with ISOP, suggesting that this effect may be responsible, at least in part, for the
361 strong reduction of serum TG steady state levels that are observed following treatment
362 with these AR-agonists.

363

364 **Conclusion**

365

366 The present data indicate that stimulation of α_1 - or $\beta_{1/2}$ -AR- can efficiently reduce
367 serum TRL levels via stimulation of TG hydrolysis, transport, metabolism and
368 clearance, as well as inhibition of hepatic TG synthesis. Given that stress-response
369 includes adrenoceptor stimulation, our data further support that the stress-induced
370 changes in serum TG levels are mediated by α_1 - and $\beta_{1/2}$ -ARs [4] in a PPAR α -
371 independent fashion, further supporting that PPAR α activators, such as fibrates, may
372 not be effective in the treatment of stress-related hypertriglyceridemia. Additional
373 research may identify these PPAR α -independent triggers providing alternative
374 pharmacological targets for new pharmacological entities that may complement
375 current therapies.

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384 **Materials and methods**

385

386 **Animals**

387

388 Adult male *Ppara*-null mice [48, 49], 7-8 weeks old, grown on the 129/SV
389 background and strain-matched wild-type littermate controls, raised at NIH Animal
390 Center, were used in this study. All mice followed a NIH-31 rodent chow based diet
391 (Zeigler, Gardners, PA) and had an *ad libitum* continuous access to drinking water.
392 Five mice per cage were housed under a standard 12-h light, 12-h dark cycle and all
393 mice were monitored daily in order to detect outward signs of distress or adverse
394 health effects. All studies involving experimental animals were carried out in
395 accordance with Institute of Laboratory Animal Resources guidelines and were
396 approved by the National Cancer Institute Animal Care and Use Committee.

397

398 **Drugs and treatment**

399

400 Phenylephrine hydrochloride (Sigma-Aldrich; 2mg/kg i.p.; PH) and isoprenaline
401 hydrochloride (Sigma-Aldrich; 2mg/kg, i.p.; ISOP), were dissolved in normal saline
402 and administered twice to three times a day and for four consecutive days (dosing
403 regimen: Total 2-3-3-2=10 injections), in order to stimulate α_1 -ARs and $\beta_{1/2}$ -ARs,
404 respectively. The selection of the dosing schedule of adrenergic receptor agonists was
405 based on the literature to achieve sufficient stimulation of the adrenergic receptors
406 [50]. The controls received normal saline and mice were not fasted during treatment.
407 Two hrs after the last drug treatment (3-4p.m.), mice were killed by carbon dioxide
408 asphyxiation and trunk blood was collected in BD Microtainer Serum Separator
409 Tubes (Becton, Dickinson and Company, USA) for biochemical and hormonal
410 analyses. Liver and white adipose tissue (W.A.T.) samples were dissected for total
411 RNA, cellular and nuclear protein extraction and were kept along with serum samples
412 at -80 °C until assayed. Each treatment group included five to six animals and the
413 findings were confirmed by three different experiments.

414

415 **Quantitative real-time PCR**

416

417 Total RNA was isolated from the liver and W.A.T. using the Trizol reagent
418 (Invitrogen, Carlsbad, CA) following the manufacturer's protocol. The concentration
419 of total RNA was determined spectrophotometrically. Quantitative real-time PCR
420 (qPCR) was performed with cDNA generated from 1 μ g total RNA using the
421 SuperScript III reverse transcriptase kit (Invitrogen). The gene-specific primers were
422 designed for qPCR using the Primer Express software (Applied Biosystems, Foster
423 City, CA). The sequences for the forward and reverse primers used are shown in
424 Table 3. For the real-time reactions the SYBR Green PCR master mix (Applied
425 Biosystems, Warrington, UK) was used. These reactions were carried out using the
426 ABI PRISM 7900 HT sequence detection system (Applied Biosystems). The relative
427 mRNA expression levels were normalized to β -actin mRNA and the absolute levels
428 were determined using the comparative threshold cycle method.

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434 **Western blot analysis**

435

436 Nuclear extracts of liver samples were used for the immunoblot analysis of PPAR α
437 and hepatocyte nuclear factor 4 α (HNF4 α) protein expression. The NE-PER nuclear
438 extraction kit (Pierce, Rockford, IL) was used for the preparation of these extracts.

439 The phosphorylation of protein kinase B (Akt) and forkhead box protein O1 (FoxO1)
440 was assessed in total cellular proteins, while the phosphorylation of cAMP-response
441 element-binding protein (CREB) was analyzed in nuclear proteins. Drug-induced
442 alterations at hepatic ApoE protein levels were assessed in total cellular proteins.

443 Alterations in the phosphorylation of HSL, in total ATGL, perilipin 5 (PLIN5) and
444 total perilipin apoprotein levels were assessed in total cellular proteins extracted from
445 the W.A.T.. The BCA protein assay (Pierce, Rockford, IL) was used for the

446 determination of protein concentrations. Proteins were subjected to sodium dodecyl
447 sulfate-polyacrylamide gel electrophoresis and immunoblotting using the following
448 antibodies: goat polyclonal HNF4 α IgG (Santa Cruz Biotechnology), rabbit

449 polyclonal phospho-AKT IgG (Ser473; Santa Cruz Biotechnology), rabbit polyclonal
450 phospho-FOXO1 (Ser256; Santa Cruz Biotechnology), rabbit polyclonal phospho-
451 CREB-1 IgG (Ser133; Santa Cruz Biotechnology) and rabbit monoclonal anti-mouse

452 ApoE IgG (Meridian USA). In addition, the rabbit polyclonal phospho-HSL (Ser563,
453 Ser565, Ser660) and total HSL and Perilipin IgGs were also used (Lipolysis
454 Activation Antibody Sampler kit, Cell Signaling). The goat polyclonal PLIN5 and

455 ATGL IgGs, as well as the rabbit polyclonal AKT, FOXO1 and CREB IgGs (Santa
456 Cruz Biotechnology) were also used. As loading control the immunoblotting with

457 mouse β -actin, Histone-H3 and GAPDH antibodies (Santa Cruz Biotechnology) was
458 used. The anti-rabbit, anti-goat or anti-mouse IgG horseradish peroxidase conjugated

459 antibodies (Cell Signaling Technology) were used as secondary antibodies and the
460 proteins were detected using an enhanced chemiluminescence detection kit (Thermo

461 Scientific-Pierce, Rockford, IL). All western blot images were submitted to
462 quantitation using the Image Processing and Analysis in Java soft ware (Image J).

463

464 **Preparation of hepatocyte cultures**

465

466 For the *in vitro* experiment, hepatocytes were prepared following a modified method
467 based on a previous report [51]. In brief, for the isolation of parenchymal hepatocytes
468 the *in situ* perfusion of the murine liver was used. The isolated hepatocytes were
469 suspended in Williams' Medium E supplemented with L-glutamine, penicillin and
470 streptomycin and then, they were plated at a density of $0.80-1.0 \times 10^6$ cells in 60 mm
471 diameter collagen type I coated dish (BIOCOAT, Cell Environment, Becton
472 Dickinson Labware, UK). The trypan blue dye exclusion was used to check the
473 viability of isolated cells. Only primary hepatocytes with viability higher than 85%
474 just before plating were cultured at 37°C for 24 h under an atmosphere of humidified
475 5% CO_2 in order to allow the cells to adhere to the dish. Time and dose response
476 experiments started 24 hours later. The cells were cultured in the presence of either
477 AR-agonists, PH or ISOP, at different doses (1-100 μM) and for a period of time
478 ranging from 4-36 hours. Here are presented only data from the incubation of primary
479 hepatocytes with the AR-agonists at a concentration of $25\mu\text{M}$ for 24-h, as they clearly
480 indicate the direct effect of PH and ISOP on *Hnf4a* expression.

481

482

483 **Determination of post-prandial triglyceride kinetics following oral** 484 **administration of olive oil**

485

486 In order to compare the effect of AR-agonist treatment on the post-prandial
487 triglyceride kinetics, groups of 6-8 mice were used. The determination of the post-
488 prandial triglyceride kinetics was performed as previously described [52, 53]. Values
489 are expressed in $\text{mg/dl} \pm$ standard error of the mean.

490

491 **Rate of hepatic very low density (VLDL) triglyceride production in mice treated** 492 **with phenylephrine or isoprenaline**

493

494 In order to assess the effects of AR-agonist treatment on hepatic VLDL triglyceride
495 secretion, 6-8 mice per treatment group were used. Briefly, treated mice were injected
496 intraperitoneally with Triton-WR1339 at a dose of 500 mg/kg b.w, using a 15%
497 solution (w/v) in 0.9% NaCl. Triton-WR 1339 inhibits completely VLDL catabolism,

498 as previously described [29, 31]. Serum samples were collected 90 min following the
499 injection with Triton WR 1339, in order to minimize the influence of handling stress
500 on the tested mice. As a baseline control, serum samples were collected 1 min
501 following the injection with the detergent. Then, serum TG levels were determined
502 again at 90 min post-injection and linear graphs of TG concentration vs time were
503 generated. The rate of VLDL-triglyceride secretion (expressed in mg/dl/min) was
504 calculated from the slope of the linear graphs for each individual mouse. The slopes
505 were grouped together and plotted in a bar-graph as mean \pm standard error of the
506 mean. Statistical analysis was performed using the Student t-test.

507 **Determination of total hepatic cholesterol and triglyceride content**

508

509 Tissue triglyceride determination was performed following the method previously
510 described by Karavia et al [53]. Results are expressed as milligram (mg) of
511 triglycerides per gram of tissue \pm standard error of the mean.

512

513 **Hormonal and biochemical determinations**

514

515 Serum total cholesterol levels were measured using the Cholesterol EIA kit (Wako
516 Diagnostics, Richmond, VA) and the levels of serum non-esterified fatty acids were
517 determined using the NEFA C, EIA kit (Wako Chemicals GmbH, Neuss, Germany).

518 Serum triglyceride levels were analysed using the GPO-Trinder Kit (Sigma). In brief,
519 the serum sample (10 μ l) was diluted in 40 μ l Phosphate buffered saline (PBS), and the
520 dilute sample (7,5 μ l) was analyzed for triglycerides, following the manufacturer's
521 instructions. Triglyceride concentrations were determined spectrophotometrically at
522 540 nm as previously described [54].

523 Serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST) [17]
524 levels were determined using the Discrete Pak ALT and AST Reagents kits
525 (Catachem Inc, Bridgeport, CT).

526

527 **Statistical analysis**

528

529 The data of the present study are presented as the mean \pm SE and were analysed using
530 the one-way analysis of variance (ANOVA) that was followed by multiple
531 comparisons with Bonferonni's and Tuckey's least honest significant difference
532 methods. The significance level for all analyses was set at probability of less than
533 0.05.

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	<i>Wild type</i>			<i>Ppara null</i>		
	Control	PH	ISOP	Control	PH	ISOP
<i>ALT</i>	9.1±0.9	11.3±4.9	7.1±2.0	10.1±2.3	11.4±2.7	8.3±3.4
<i>AST</i>	25.2±4.6	29.5±11.2	15.5±2.0	24.6±5.3	28.2±8.9	18.9±2.7

Table 1. AR-induced alterations in serum ALT and AST concentration

Adrenergic receptor (AR); Alanine aminotransferase (ALT), aspartate aminotransferase (AST), phenylephrine (PH), isoprenaline (ISOP), (wild type, n=20; *Ppara*-null, n=15).

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Treatment	1 st day	4 th day
Control	25.1±1.0	26.3±0.8
Phenylephrine	27.9±0.4	28.6±0.5
Isoprenaline	24.0±0.6	25.2±0.5

Table 2.
Alterations
in the body
weight
following
adrenergic

receptor agonist treatment.

Body weight values are expressed in g. Phenylephrine, α_1 -adrenergic receptor (AR) agonist; Isoprenaline, $\beta_{1/2}$ -AR agonist.

830 **Table 3**

831 The list of 5' to 3' oligonucleotide sequences used as forward and reverse primers

832			
833	PPAR α	CAGTGGGGAGAGAGGACAGA	AGTTCGGGAACAAGACGTTG
834			
835	PPAR γ	CACAAGAGCTGACCCAATGGT	AATAATAAGGTGGAGATGCAGGTTCT
836			
837	HNF4 α	CGGAGCCCCTGCAAAGT	ACTATCCAGTCTCACAGCCCATTC
838			
839	Cyp8b1	ACGCTTCCTCTATCGCCTGAA	GTG CCTCAGACGCAGAGGAT
840			
841	BAAT	ACAGGCCTGGCCCCCTTCA	CCCATGGGGTGGACCCCAT
842			
843	ACADM	AGCTCTAGACGAAGCCACGA	GCGAGCAGAAATGAACTCC
844			
845	HSL	CCTCCAAGCAGGGCAAAGA	GCGTAAATCCATGCTGTGTGA
846			
847	ATGL/PNPLA2	CCACTCACATCTACGGAGCC	TAATGTTGGCACCTGCTTCA
848			
849	AADAC	ACCGCTTCCAGATGCTATTG	TGATTCCCAAAGTTCACCA
850			
851	MTTP	CGTGGTGAAAGGGCTTATTC	TCGCGATACCACAGAATGAA
852			
853	DGAT1	GACGGCTACTGGGATCTGA	TCACCACACACCAATTCAGG
854			
855	DGAT2	CGCAGCGAAAACAAGAATAA	GAAGATGTCTTGGAGGGCTG
856			
857	LPL	TTGGCTCCAGAGTTTGACC	TGTGTCTTCAGGGTTCCTTAG
858			
859	CES3/TGH	TGGTATTTGGTGTCCCATCA	GCTTGGGCGATACTCAAACCT
860			
861	CD36	GCGACATGATTAATGGCACA	CCTGCAAATGTCAGAGGAAA
862			
863	NR4A	ATTGAGCTTGAATACAGGGCA	GCTAGAAGGACTGCGGAGC
864			
865	LDL-r	GGGAACATTTGGGGTCTGT	AGTCTTCTGCTGCAACTCCG
866			
867	β -actin	TATTGGCAACGAGCGGTTCC	GGCATAGAGGTCTTTACGGATGTC

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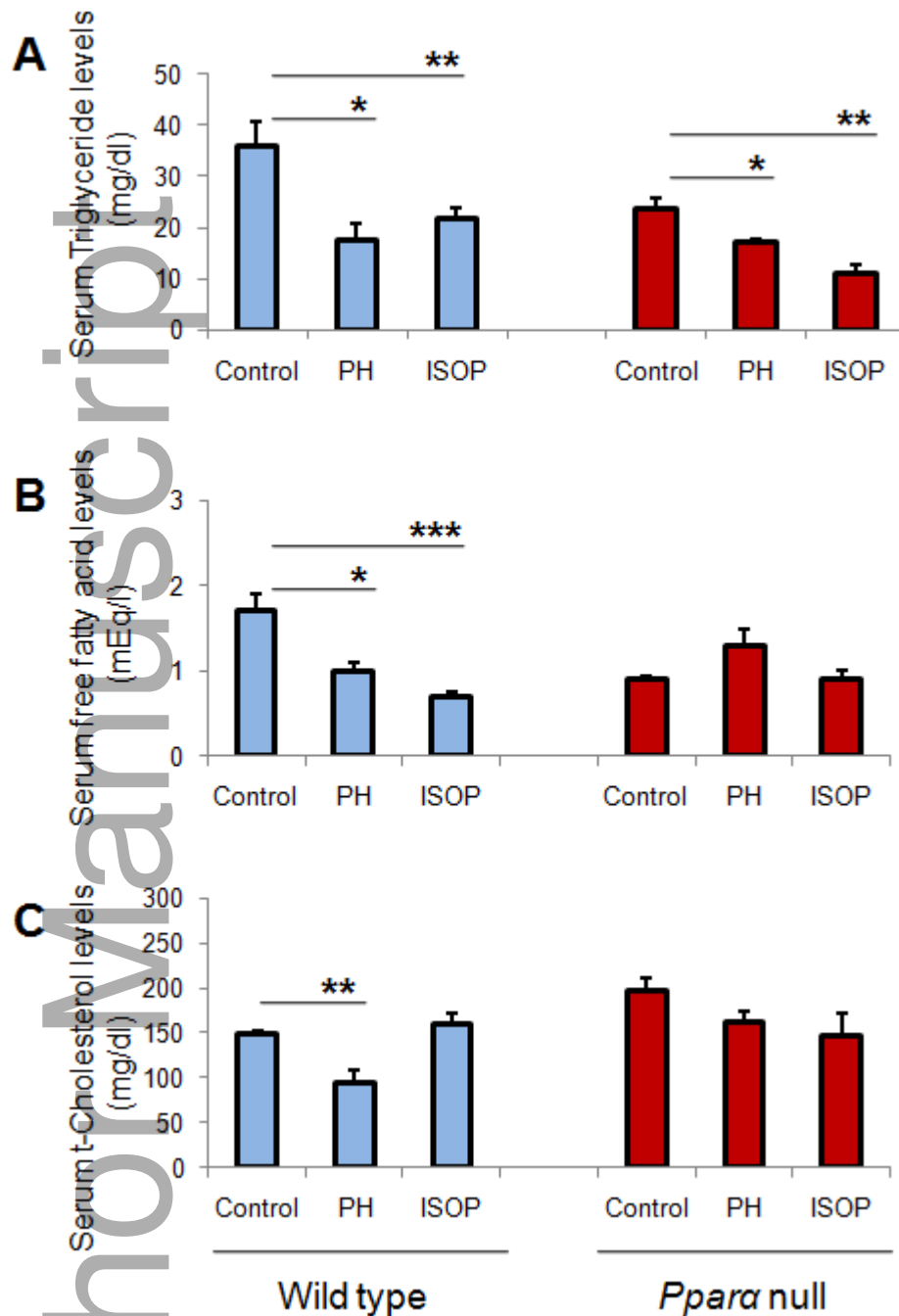
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880 **Legends**

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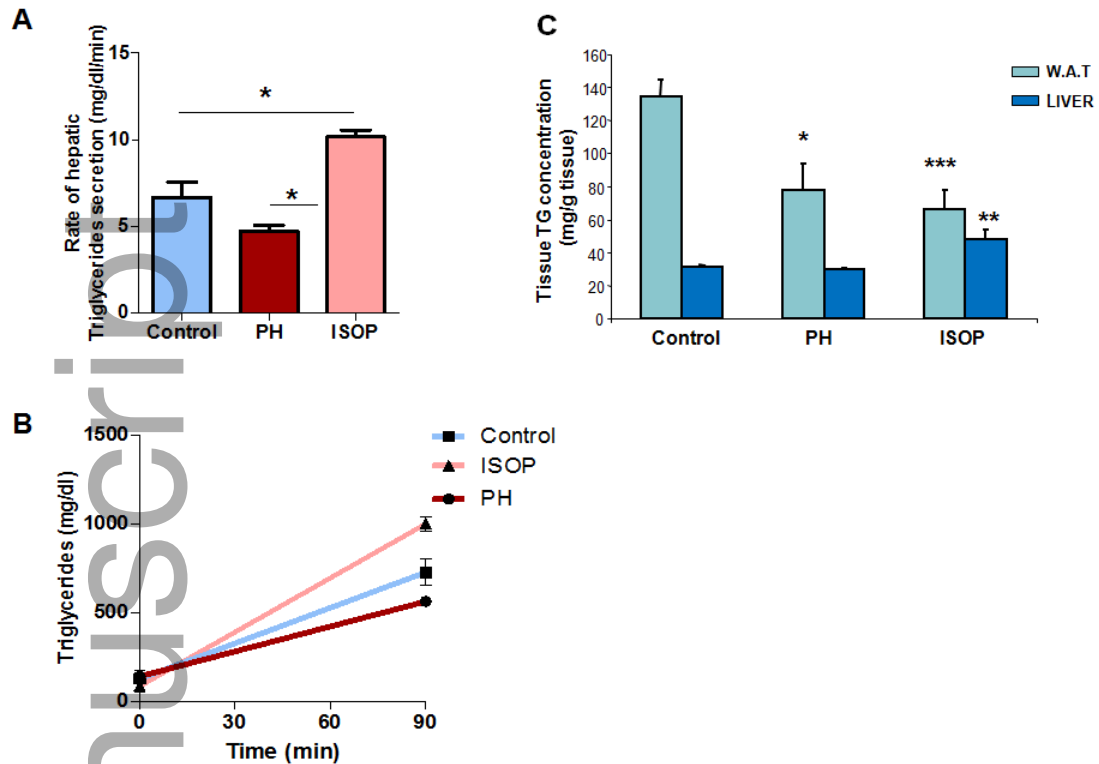
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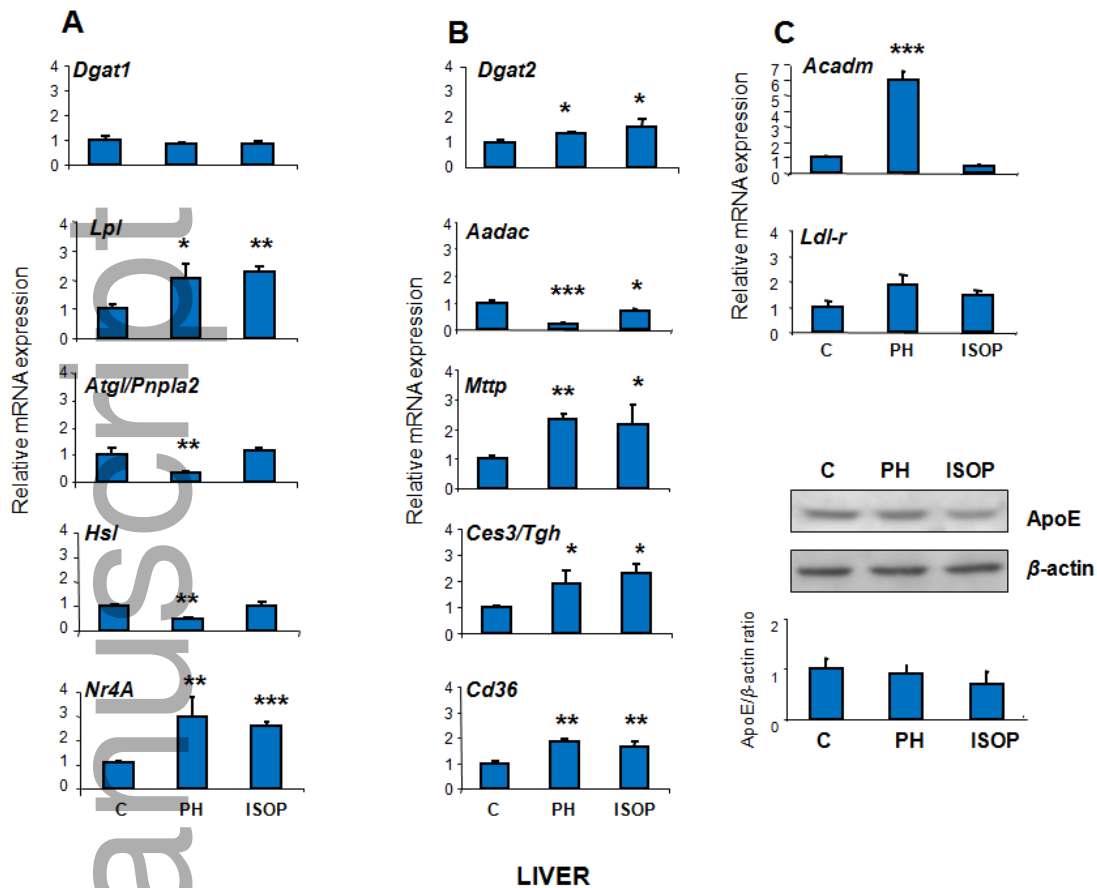
884 **Fig. 1.** Effects of PH and ISOP treatment on serum lipid markers. PH: phenylephrine
 885 (α_1 -AR agonist); ISOP: isoprenaline (β_1/β_2 -AR agonist). Values are expressed as
 886 mean \pm SEM, n: 5 per treatment group and comparisons took place between controls
 887 and drug-treated mice; * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$.



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890 **Fig. 2.** Effects of PH and ISOP treatment on kinetic parameters of serum triglyceride
 891 metabolism. Panel A shows the rate of hepatic VLDL triglyceride secretion of the PH-
 892 or ISOP-treated mice and controls and Panel B represents the kinetics of post-prandial
 893 triglyceride clearance in PH- and ISOP-treated mice. Panel C shows TG concentration
 894 in the liver and white adipose tissue (W.A.T.) of mice following treatment with either
 895 PH: phenylephrine (α_1 -AR agonist), ISOP: isoprenaline ($\beta_{1/2}$ -AR agonist) or normal
 896 saline (controls). Values are expressed as mean \pm SEM, n: 5 per treatment group and
 897 comparisons took place between controls and drug-treated mice. Group differences
 898 were calculated by one-way ANOVA, followed by Bonferonni's test. All experiments
 899 were performed as described in Materials and Methods. * $P < 0.05$, ** $P < 0.01$, *** P
 900 < 0.001 .



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902 **Fig. 3.** Adrenergic receptor-mediated effect on hepatic factors regulating TG serum
 903 levels.

904 (A) Effect of AR-agonists on genes involved in TG synthesis and lipolysis in the liver.

905 (B) Effect of AR-agonists on genes involved in TG metabolism and clearance in the

906 liver. (C) Effect of AR-agonists on factors important in lipid β -oxidation, the
 907 clearance of triglyceride rich lipoproteins and the transport of free fatty acids.

908 Comparisons were between controls and drug-treated mice. *Dgat1*: diacyl glycerol

909 acyltransferase, *Dgat2*: diacyl glycerol acyltransferase 2, *Lpl*: lipoprotein lipase, *Hsl*:

910 hormone sensitive lipase, *Atgl/Pnpla2*: adipose triglyceride lipase/patatin-like
 911 phospholipase domain containing 2, *Nr4A*: orphan nuclear receptor NR4A, *Aadac*:

912 arylacetamide deacetylase, *Cd36*: cluster of differentiation 36 or fatty acid transporter,

913 *Ces3/tgh*: carboxylesterase 3, *Mttp*: microsomal triglyceride transfer, *Acadm*: acyl-

914 CoA dehydrogenase, *Ldl-r*: low density lipoprotein receptor, ApoE: apolipoprotein E.

915 In the western blot, three samples per treatment were loaded in three different blots.

916 C: Control, phenylephrine (α_1 -AR-agonist, PH), isoprenaline ($\beta_{1/2}$ -AR agonist, ISOP).

917 Values are expressed as mean \pm SEM, n:5-6 mice per treatment group. Group

918 differences were calculated by one-way ANOVA, followed by Bonferonni's test. **P*
919 < 0.05, ***P* < 0.01, ****P* < 0.001.

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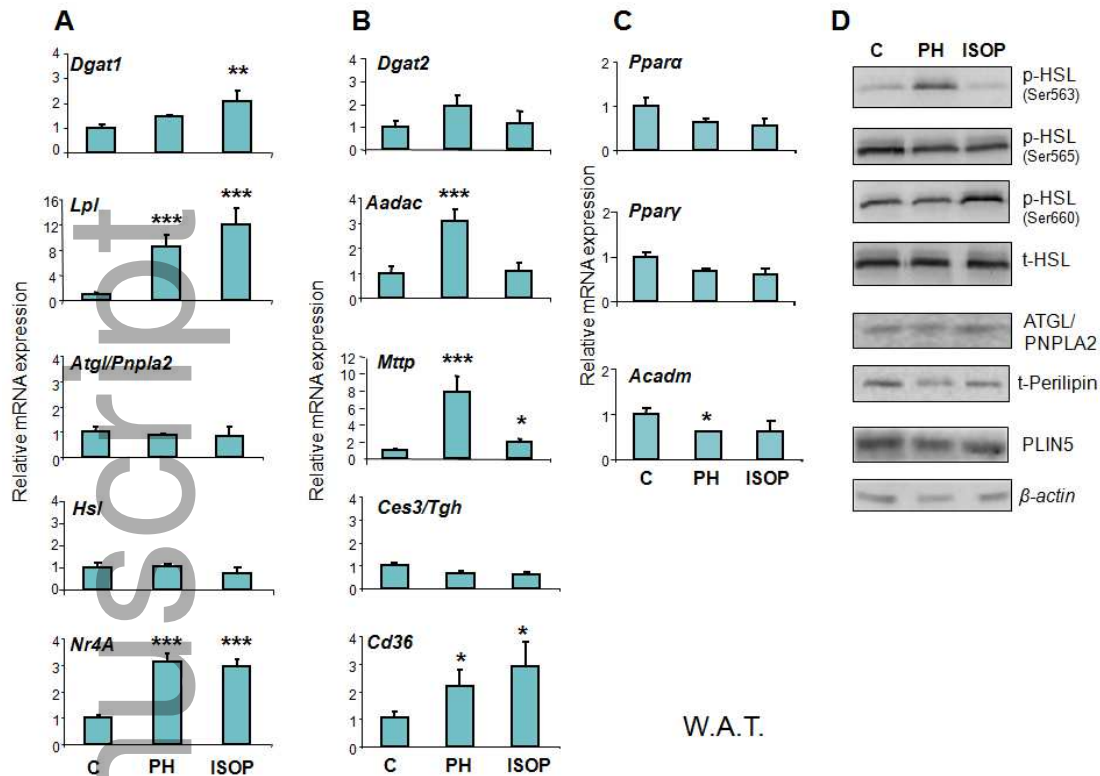
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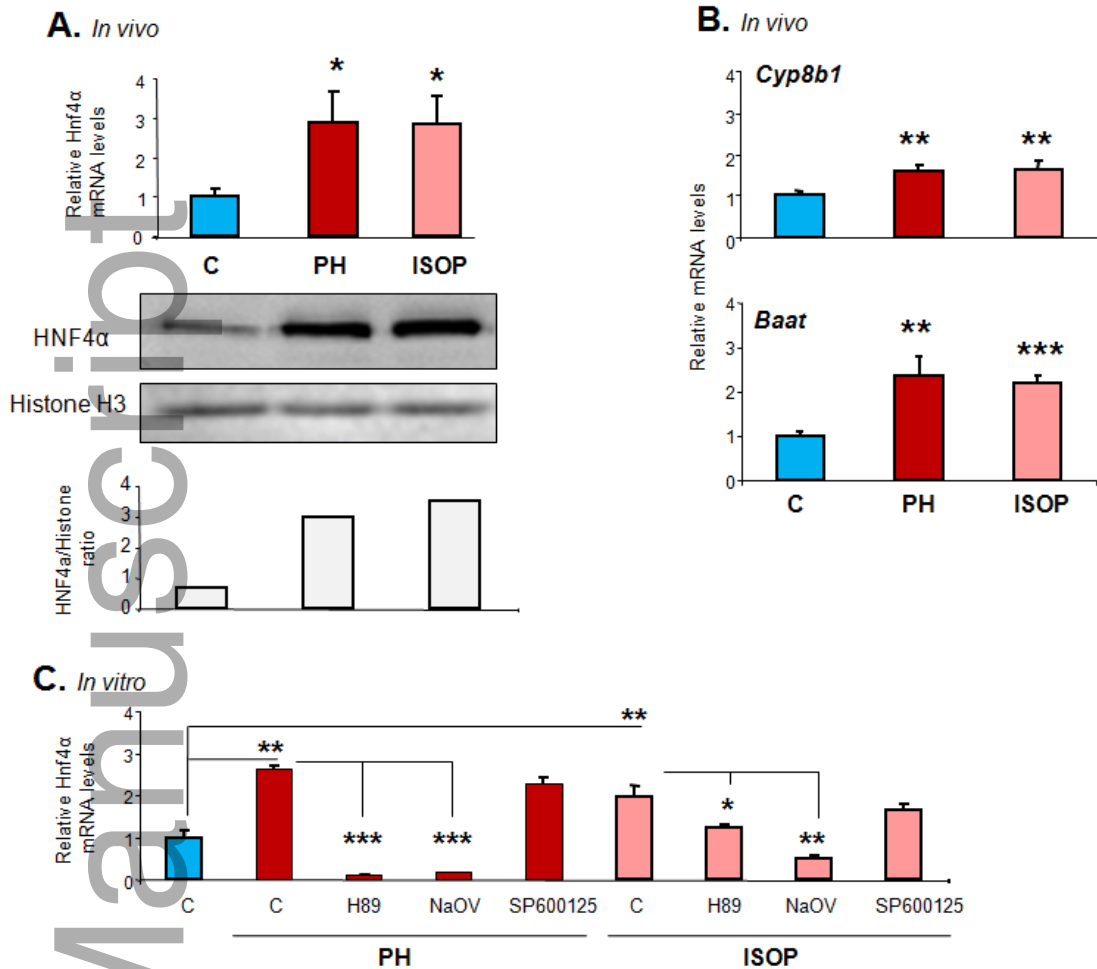
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941 **Fig. 4.** Adrenergic receptor-mediated effect on various factors expressed in the
 942 W.A.T. regulating TG serum levels (A) Effect of phenylephrine (α_1 -AR agonist, PH)
 943 and isoprenaline ($\beta_{1/2}$ -AR agonist, ISOP) on the expression of genes involved in TG
 944 synthesis and lipolysis. (B) Effect of PH and ISOP on the expression of genes
 945 involved in TG metabolism and clearance. (C) Effect of PH and ISOP on factors
 946 important in lipid β -oxidation, the clearance of triglyceride rich lipoproteins and the
 947 transport of free fatty acids. Comparisons were between controls and AR-agonist-
 948 exposed mice; *Dgat1*: diacyl glycerol acyltransferase 1 (acyl coenzyme A (CoA),
 949 *Dgat2*: diacyl glycerol acyltransferase 2, *Lpl*: lipoprotein lipase, *Hsl*: hormone
 950 sensitive lipase, *Atgl/Pnpla2*: adipose triglyceride lipase/patatin-like phospholipase
 951 domain containing 2, *Nr4a*: orphan nuclear receptor, *Aadam*: arylacetamide
 952 deacetylase, *Cd36*: cluster of differentiation 36 or fatty acid transporter, *Ces3/tgh*:
 953 carboxylesterase 3, *Mtp*: microsomal triglyceride transfer, PLIN5: perilipin 5, AR:
 954 adrenergic receptor, C: Control, W.A.T.: white adipose tissue. Values are expressed as
 955 mean \pm SEM, n:5-6 mice per treatment group; Group differences were calculated by
 956 one-way ANOVA, followed by Bonferonni's test. * $P < 0.05$, ** $P < 0.01$, *** $P <$
 957 0.001. Lanes in western blots correspond to one sample per treatment and represent
 958 one sample of three separate samples tested in different blots.

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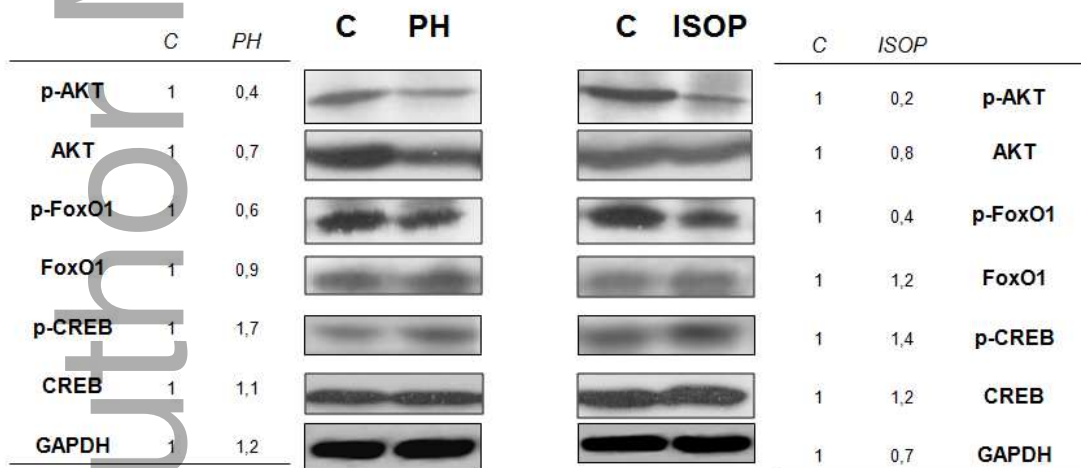


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980 **Fig. 5.** The effect of adrenergic receptor-related pathways on *Hnf4a* expression. (A)
 981 Following treatment with AR-agonists, hepatic *Hnf4a* mRNA levels were analysed in
 982 wild-type mice by qPCR. HNF4 α protein was determined in liver nuclear fractions by
 983 western blot analysis. Histone H3 served as a loading control. In the bar graph the
 984 quantified data from the western blot image are shown presented as the ratio of
 985 HNF4 α /Histone H3. (B) *Cyp8b1* and *Baat* mRNA levels were analyzed in livers of
 986 wild-type mice by qPCR following treatment with AR-agonists. (C) *Hnf4a* mRNA
 987 levels were determined by qPCR following treatment of primary hepatocyte cultures
 988 with AR-agonists for 24 hours. Primary hepatocytes were also treated with AR-
 989 agonists in combination with either the JNK inhibitor, SP600125, the PKA inhibitor,
 990 H89, or the phosphatase- and ATPase inhibitor, NaOV (concentration of the inhibitors
 991 in the medium: 10 μ M and duration of incubation: 24 hrs). Values were normalized to
 992 β -actin and are expressed as mean \pm SEM (n=8-10). In the *in vivo* experiment
 993 comparisons were between controls and drug-treated mice. In the *in vitro* experiment
 994 comparisons were between DMSO and drug-treated hepatocytes, (n=3-4). AR:

995 adrenergic receptor, C: control (DMSO treated primary hepatocytes), PH:
 996 Phenylephrine (α_1 -AR agonist), ISOP: Isoprenaline ($\beta_{1/2}$ -AR agonist). Group
 997 differences were calculated by one-way ANOVA, followed by Bonferonni's test. **P*
 998 < 0.025, ***P* < 0.01, ****P* < 0.001.

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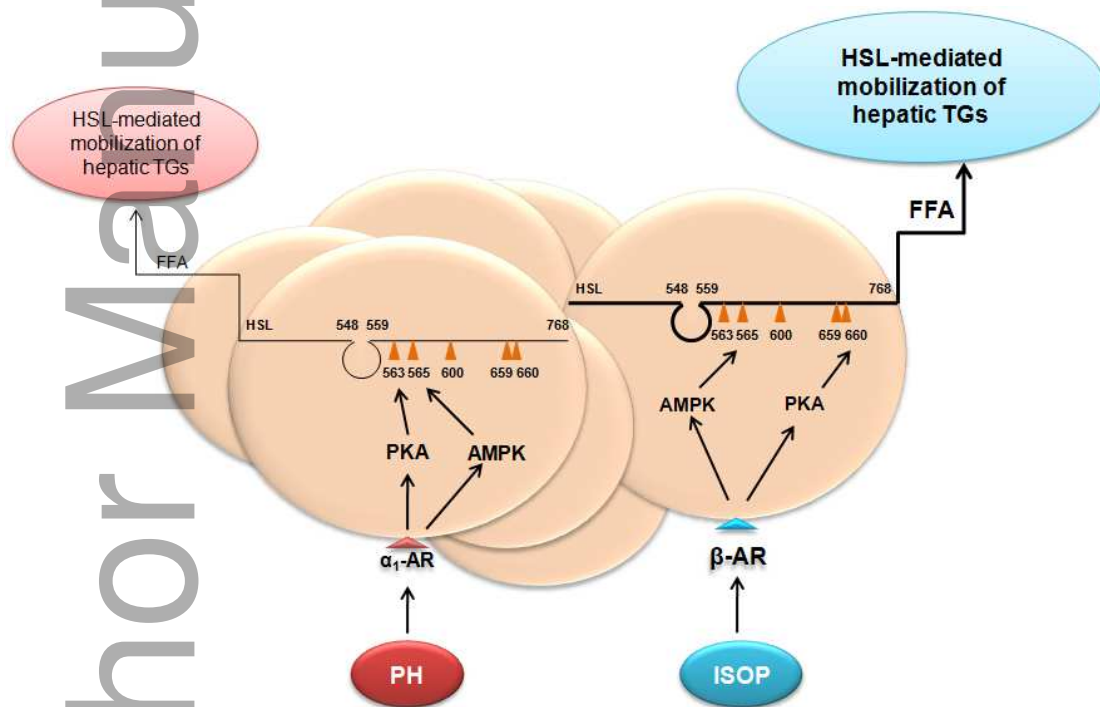


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1015 **Fig. 6.** The role of adrenergic receptors in the activation of insulin/PI3k/AKT/FoxO1
 1016 and AR/cAMP/PKA/CREB signaling pathways. Total and phosphorylated AKT and
 1017 FoxO1 expression levels were examined in hepatic total cellular proteins using
 1018 Western blot analysis. CREB phosphorylation was assessed in hepatic total cellular

1019 proteins. C: control, PH: phenylephrine (α_1 -AR agonist), ISOP: isoprenaline ($\beta_{1/2}$ -AR
 1020 agonist). The numbers next to the lanes represent the relative protein expression that is
 1021 defined as the ratio between the drug-treated and control expression, which is set at 1.

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1035 **Fig. 7.** Hypothetical model summarizing the impact of α_1 - and β -AR agonists on HSL
 1036 phosphorylation in the white adipose tissue and the subsequent hepatic TG
 1037 mobilization. The present data indicated that exposure to phenylephrine (PH, α_1 -AR
 1038 agonist) activated the cAMP-PK (PKA) resulting in HSL phosphorylation at Ser563,
 1039 whereas activation of PKA, the induced by isoprenaline (ISOP, $\beta_{1/2}$ -AR agonist) led to

1040 phosphorylation of HSL at Ser660. Both, PH and ISOP, also activated the AMP-
1041 activated kinase (AMPK), which is considered to block the PKA-dependent activation
1042 of HSL in adipocytes, when HSL phosphorylation occurs at Ser563, while it is
1043 preserved when it occurs at Ser660. The current data confirm the lesser significance
1044 of α_1 -ARs in the HSL-dependent lipolysis in adipocytes compared to that of β -ARs
1045 [42, 44]. FFA: free fatty acids.

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1049 **Acknowledgments**

1050

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1057

1058 **Author contribution**

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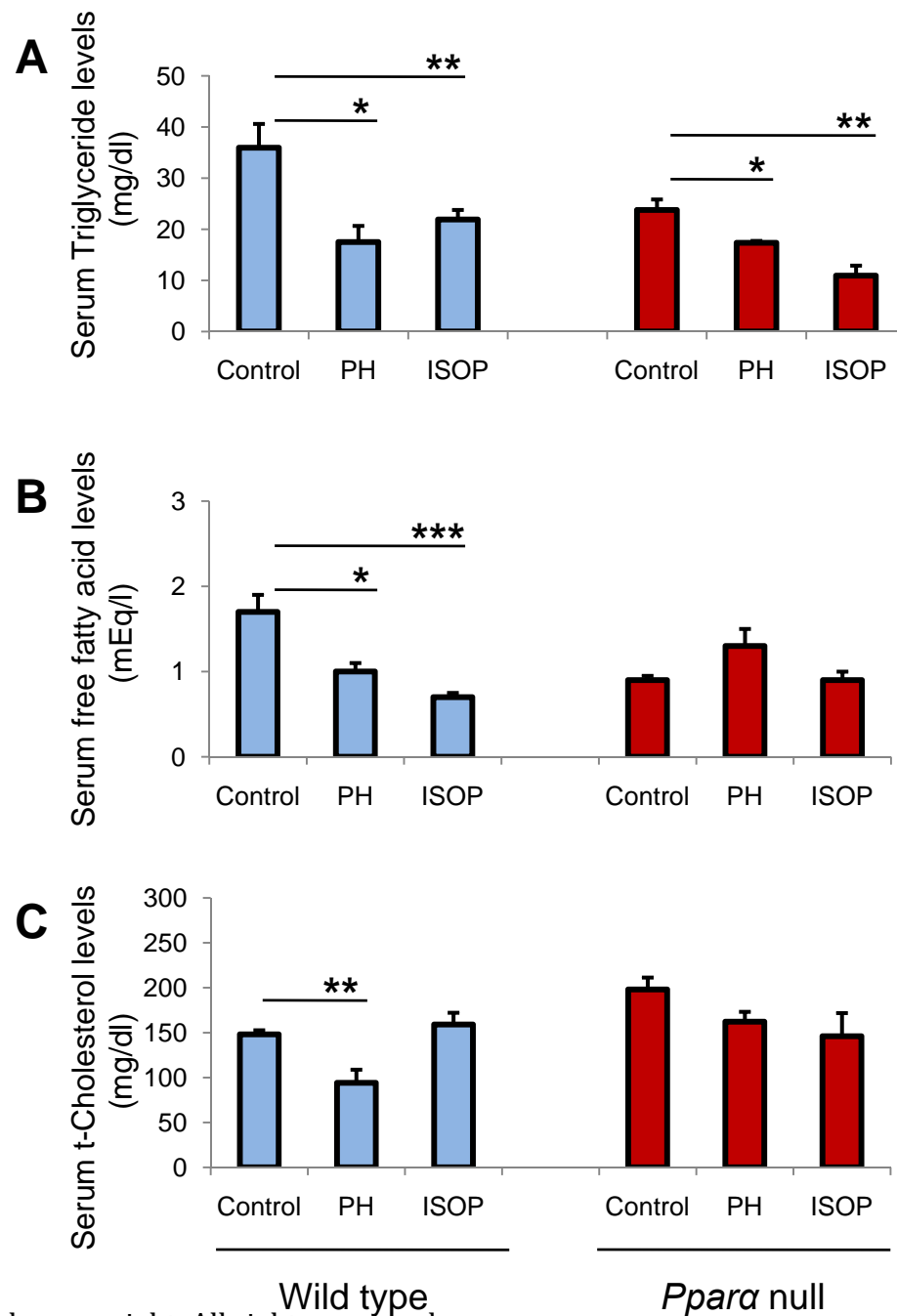
1060 MK conceived and coordinated the study. MK, KKy, TM, EX, YS, KKr, CA, AK and
1061 FJG designed and performed the experiments, and analyzed the data. MK, KKy and
1062 FJG wrote the paper.

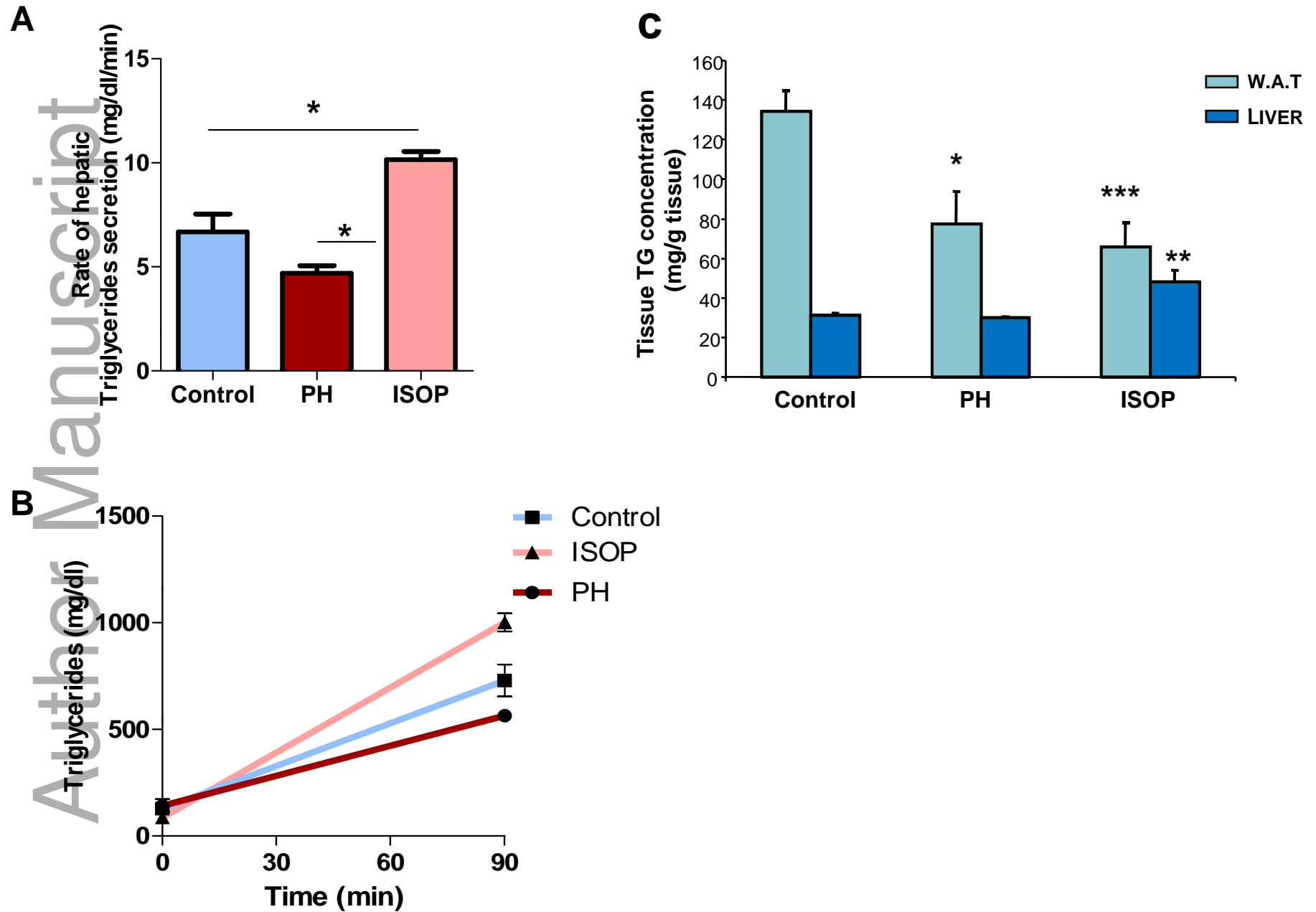
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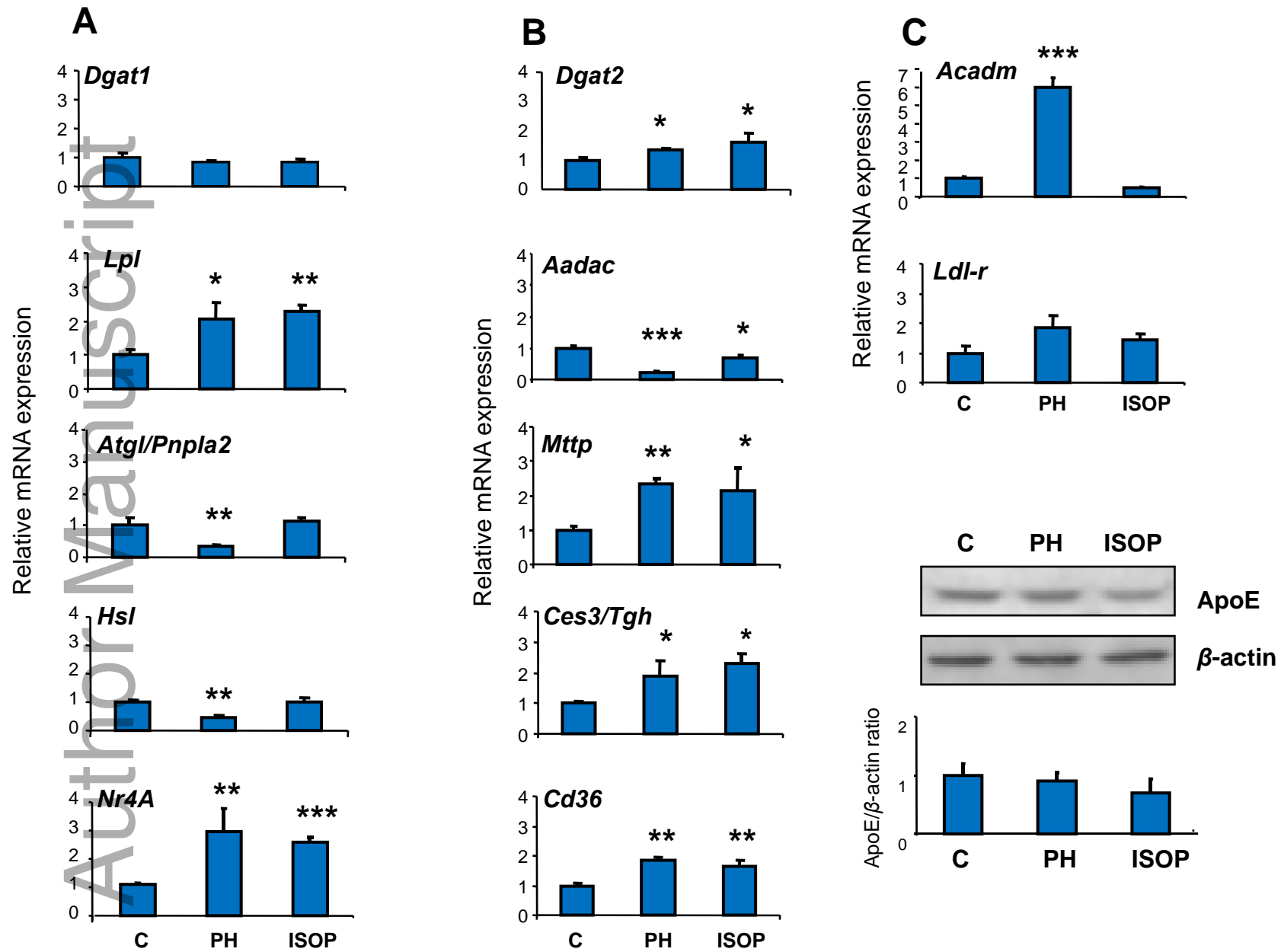
1064 **Conflict of interest**

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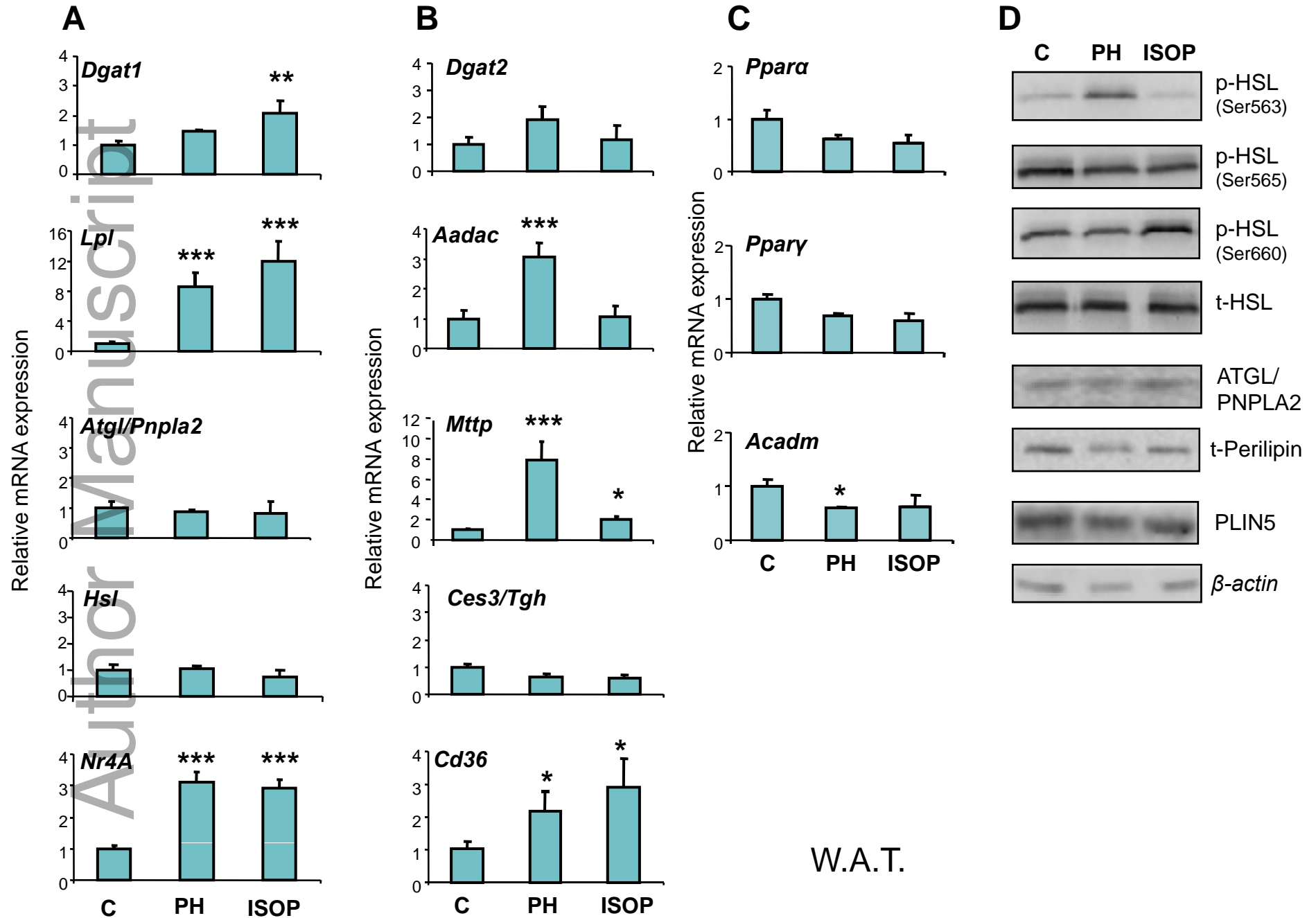
1066 The authors declare no conflict of interest.

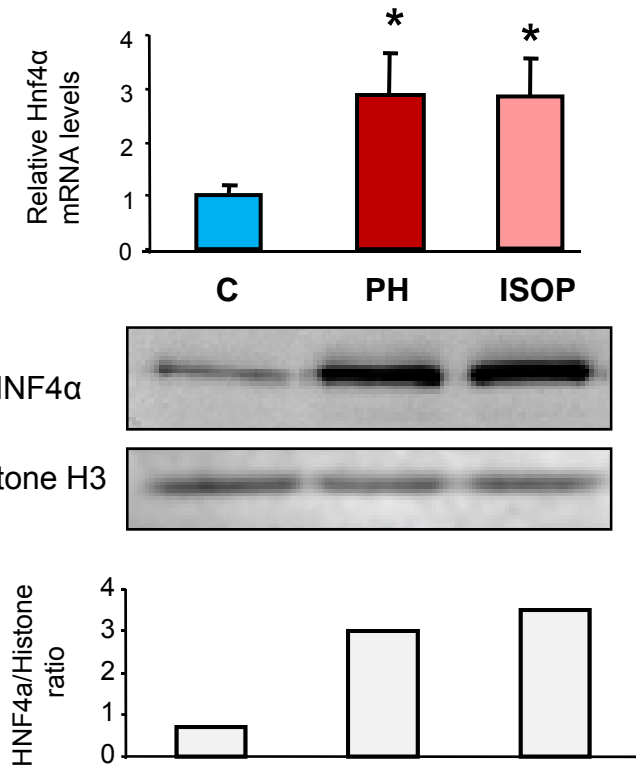
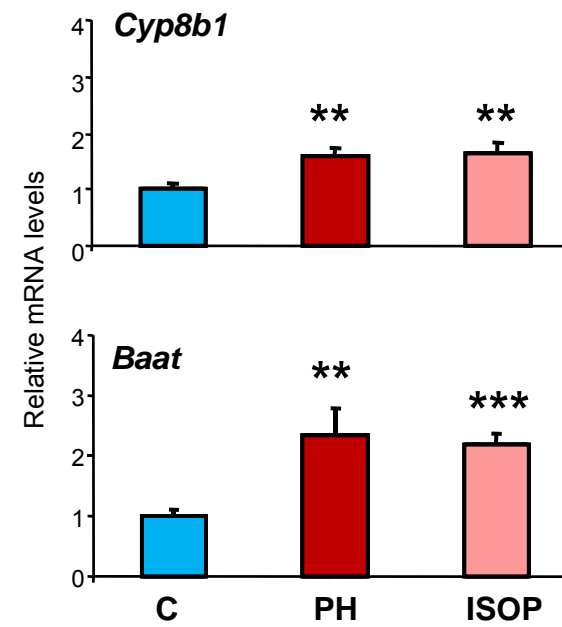






LIVER



A. In vivo**B. In vivo****C. In vitro**