

1 **Cancer causes metabolic perturbations associated with reduced insulin-stimulated glucose
2 uptake in peripheral tissues and impaired muscle microvascular perfusion.**

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16 **Conflict of Interest:** There are no conflicts to disclose.

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20 **Abstract**

21 **Background:** Redirecting glucose from skeletal muscle and adipose tissue, likely benefits the
22 tumor's energy demand to support tumor growth, as cancer patients with type 2 diabetes have 30%
23 increased mortality rates. The aim of this study was to elucidate tissue-specific contributions and
24 molecular mechanisms underlying cancer-induced metabolic perturbations.

25 **Methods:** Glucose uptake in skeletal muscle and white adipose tissue (WAT), as well as hepatic
26 glucose production, were determined in control and Lewis lung carcinoma (LLC) tumor-bearing
27 C57BL/6 mice using isotopic tracers. Skeletal muscle microvascular perfusion was analyzed via a
28 real-time contrast-enhanced ultrasound technique. Finally, the role of fatty acid turnover on
29 glycemic control was determined by treating tumor-bearing insulin-resistant mice with nicotinic
30 acid or etomoxir.

31 **Results:** LLC tumor-bearing mice displayed reduced insulin-induced blood-glucose-lowering and
32 glucose intolerance, which was restored by etomoxir or nicotinic acid. Insulin-stimulated glucose
33 uptake was 30-40% reduced in skeletal muscle and WAT of mice carrying large tumors. Despite
34 compromised glucose uptake, tumor-bearing mice displayed upregulated insulin-stimulated
35 phosphorylation of TBC1D4^{Thr642} (+18%), AKT^{Ser474} (+65%), and AKT^{Thr309} (+86%) in muscle.
36 Insulin caused a 70% increase in muscle microvascular perfusion in control mice, which was
37 abolished in tumor-bearing mice. Additionally, tumor-bearing mice displayed increased (+45%)
38 basal (not insulin-stimulated) hepatic glucose production.

39 **Conclusions:** Cancer can result in marked perturbations on at least six metabolically essential
40 functions; i) insulin's blood-glucose-lowering effect, ii) glucose tolerance, iii) skeletal muscle and
41 WAT insulin-stimulated glucose uptake, iv) intramyocellular insulin signaling, v) muscle
42 microvascular perfusion, and vi) basal hepatic glucose production in mice. The mechanism causing
43 cancer-induced insulin resistance may relate to fatty acid metabolism.

44 **Keywords:** Lewis lung carcinoma, cancer, insulin resistance, glycaemic regulation, microvascular
45 perfusion

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48 **1.1. Introduction**

49 Epidemiological and clinical studies show an association between several types of cancers and poor
50 glycemic control in humans. For example, one-third of cancer patients are glucose intolerant [1],
51 which is of clinical relevance as cancer patients with type 2 diabetes have 30% increased mortality
52 rates [2,3]. Furthermore, recent studies have suggested that insulin resistance could be an
53 underlying cause of cancer-associated loss of muscle and fat mass, called cancer cachexia [4–
54 6]. Cachexia occurs in 60–80% of cancer patients and is associated with poor prognosis [7].
55 Moreover, glucose uptake was significantly lower in lung cancer patients with cachexia during
56 hyperinsulinemia [8]. Despite the emerging link between cancer, cancer cachexia and insulin
57 resistance, the tissue-specific mechanistic and molecular cause(s) of insulin resistance in cancer is
58 largely unexplored.

59 Skeletal muscle and adipose tissue are essential for maintaining whole-body glucose homeostasis
60 by taking up the majority of glucose in response to insulin in healthy humans [9]. Insulin promotes
61 glucose disposal in skeletal muscle and adipose tissue by increasing microvascular perfusion as well
62 as mediating glucose transport across the plasma membrane [10–12]. Whether insulin resistance in
63 cancer is associated with molecular malfunctions in skeletal muscle and adipose tissue, and whether
64 impaired muscle microvascular perfusion is a potential cause of reduced insulin-stimulated glucose
65 uptake have, to our knowledge, not previously been determined.

66 Fatty acid oxidation and/or adipose tissue lipolysis are increased in many cancers [13–15].
67 Excessive fatty acids in the circulation can cause insulin resistance and have been suggested as a
68 cause of insulin resistance in obesity and type 2 diabetes [16–18]. Moreover, literature suggests

69 that excessive fatty acid turnover is a leading cause of cancer cachexia because blockade of fatty
70 acid oxidation or suppression of lipolysis in adipose tissue prevents cachexia in tumor-bearing mice
71 [19–21]. However, a role for altered fatty acid turnover in cancer-associated impaired glycemic
72 regulation is unexplored.

73 The aim of the present investigation was to elucidate tissue-specific contributions and molecular
74 mechanisms underlying impaired glycemic regulation in cancer.

75 In a mouse model of lung cancer, we found significant whole-body insulin resistance and glucose
76 intolerance that was restored by blockage of whole-body fatty acid oxidation or adipose tissue
77 lipolysis. Insulin resistance in tumor-bearing mice was associated with i) impaired glucose uptake
78 in adipose tissue and skeletal muscle despite augmented muscle insulin signaling, ii) abrogated
79 muscle microvascular perfusion in response to insulin, and iii) increased basal hepatic glucose
80 production.

81

82 **2. Material and methods**

83 *2.1. Cell culture.* Lewis lung carcinoma cells (LLC, ATCC® CRL1642™) were cultured in DMEM,
84 high glucose (Gibco #41966-029, USA) supplemented with 10% fetal bovine serum (FBS, Sigma-
85 Aldrich #F0804, USA), 1% penicillin-streptomycin (ThermoFisher Scientific #15140122, USA) (5%
86 CO₂, 37°C). Prior to inoculation into mice, LLC cells were trypsinized and washed twice with PBS.
87 LLC cells were suspended in PBS with a final concentration of $2.5 * 10^6$ cells/ml.

88 *2.2. Animals.* Female C57BL/6 (Taconic, Lille Skensved, DK) mice were group-housed at ambient
89 temperature (21-23°C) with nesting materials and kept on a 12 h:12 h light-dark cycle with access
90 to a standard rodent chow diet (Altromin no. 1324, Brogaarden, DK) and water *ad libitum*. At the
91 age of 12-14 weeks, mice were randomly assigned into control (Control) or LLC tumor-bearing
92 (LLC) groups and subcutaneously injected with 100 µl PBS with or without $2.5 * 10^5$ LLC cells

93 into the right flank. Control and LLC tumor-bearing mice were sacrificed at two time points: 15
94 days (LLC Day 15) and 21-27 days (LLC Day 21-27) after tumor transplantation. Tumor volume
95 (V) was monitored by caliper measurement and defined by $V [\text{mm}^3] = (\text{length} [\text{mm}]) \times (\text{width} [\text{mm}])^2 \times 0.52$ every 2 to 5 days [22]. Food intake was measured continuously every second day
96 during the intervention in non-tumor-bearing, tumor-bearing and etomoxir-treated (described below)
97 tumor-bearing mice. Body weight was monitored before and throughout the interventions. All
98 experiments were approved by the Danish Animal Experimental Inspectorate (Licence: 2016-15-
99 0201-01043).

100 *2.3. Etomoxir and nicotinic acid administration.* Mice housed at ambient temperature are mildly
101 cold stressed and preferentially metabolize more lipids than carbohydrate [23], thus inhibiting fat
102 metabolism might cause undue metabolic stress at ambient temperature and therefore we performed
103 this part of the experiment at thermoneutrality (30°C). After 3 weeks of acclimatization and
104 mimicking intraperitoneal (i.p) injection every other day using empty syringes, LLC transplantation
105 was performed as described above. Half of the tumor-bearing mice and half of the non-tumor-
106 bearing mice were i.p. injected daily with 5 mg/kg body weight ethyl-2-[6-(4-
107 chlorophenoxy)hexyl]-oxirane-2-carboxylate (etomoxir) (Sigma-Aldrich, US), dissolved in 5% (2-
108 Hydroxypropyl)- β -cyclodextrin solution from day 8 following tumor transplantation (LLC-Eto).
109 The other half of the tumor-bearing mice (LLC) and non-tumor-bearing control mice (Control) were
110 i.p. injected with 5% (2-Hydroxypropyl)- β -cyclodextrin solution. Mice were sacrificed 18 days
111 following tumor transplantation.

112 The nicotinic acid-administrated mice (LLC-Nico) were maintained and treated similar to etomoxir-
113 administrated mice, with the exception that 50 mg/kg body weight of nicotinic acid (Sigma-Aldrich,
114 USA) dissolved in 5% (2-Hydroxypropyl)- β -cyclodextrin solution was i.p. injected daily from day
115 8 following tumor transplantation. Mice were sacrificed 19 days following tumor transplantation.

117 *Glucose tolerance test.* D-Glucose (2 g/kg body weight) was injected i.p. following a 6 h fast from
118 7:00 AM. Blood glucose levels before (0 minutes), 20 minutes, 40 minutes, 60 minutes and 90
119 minutes following glucose injection were measured using a glucometer (Bayer Contour,
120 Switzerland). For measurements of plasma insulin concentration, blood was collected from the tail
121 vein at time points 0 and 20 minutes. Plasma insulin was analyzed by ELISA in duplicates (Mouse
122 Ultrasensitive Insulin ELISA, #80-INSTRU-E10, ALPCO Diagnostics, USA).

123 *2.4. Body composition analysis.* Fat mass and lean body mass were determined by quantitative
124 magnetic resonance imaging (EchoMRI-4in1TM, Echo Medical System LLC, USA) 0-2 days
125 before termination. Quantitative magnetic resonance imaging of the tumor itself disclose the tumor
126 as 5% “fat mass” and 95% as “lean body mass” (unpublished observations), and therefore 5% and
127 95% of the dissected tumor mass was subtracted from the fat mass and lean body mass, respectively.

128 *In vivo 2-deoxy-glucose uptake experiments.* To determine glucose uptake in skeletal muscle,
129 perigonadal white adipose tissue (WAT), and the tumor, ³H-labelled 2-deoxy-glucose ([³H]2DG)

130 (Perkin Elmer, USA) was injected retro-orbitally (r.o.) in a bolus of saline (6 μ l/g body weight)
131 containing 66.7 μ Ci/ml [³H]2DG in mice as described [24]. The injectate also contained 0.3 U/kg
132 body weight insulin (Actrapid; Novo Nordisk, DK) or a comparable volume of saline as well. Mice
133 fasted for 3-5 h from 07:00 AM and were anesthetized (i.p. injection of 7.5 mg pentobarbital
134 sodium per 100 g body weight) 15 minutes before the r.o. injection. Blood samples were collected
135 from the tail vein and analyzed for glucose concentration using a glucometer (Bayer Contour,
136 Switzerland). Sampling was performed immediately prior to insulin or saline injection and either
137 after 5 and 10 minutes, or after 3, 6, 9, and 12 minutes as indicated in the figures. After 10 or 12
138 minutes, mice were humanely euthanized by cervical dislocation and the tumor, perigonadal WAT,
139 tibialis anterior (TA), and gastrocnemius muscles were excised and quickly frozen in liquid
140 nitrogen and stored at -80°C until processing. Once tissues were removed, blood was collected by

141 punctuation of the heart, centrifuged (13,000 rpm, 5 minutes) and plasma frozen at -80°C. Plasma
142 samples were analyzed for insulin concentration, IL-6, TNF- α , and specific [3H]2DG tracer activity.
143 Plasma insulin was analyzed as described above. Tissue-specific 2DG uptake was analyzed as
144 described [25,26]. Plasma IL-6 and TNF- α were analyzed using V-PLEX Custom Mouse Cytokine
145 Proinflammatory Panel1 (Mesoscale, #K152A0H-1) according to the manufacturer's
146 recommendation, loading 12.5 μ l of plasma to each well.
147 Calculations of the whole-body glucose uptake index were performed as follows based on the body
148 mass compositions obtained from the MRI scans: $y = a * b$, where y is 2DG uptake index (μ mol/h), a
149 is 2DG uptake rate (μ mol/g/h), b is either tumor, fat, or 0.5*lean mass (g). Half of the lean mass
150 was estimated to be muscle mass based on the study of Rolfe and Brown [27]. It was an assumption
151 that all fat depots on average displayed glucose uptake similar to our measured WAT depot.
152 *2.5. Microvascular perfusion in muscle.* Mice were anesthetized with an i.p. injection of 11 μ l/g
153 body weight of Fentanyl (0.05 mg/ml, Dechra, DK), Midazolam (5 mg/ml, Accord Healthcare, UK)
154 and Acepromazine (10 mg/ml, Pharmaxim, SE), and placed on a heating pad. In control (n=6) and
155 large (tumor size > 800 mm³) tumor-bearing mice (n=6,), microvascular perfusion (MVP) was
156 measured across the adductor magnus and semimembranosus muscles, with real-time contrast-
157 enhanced ultrasound technique using a linear-array transducer connected to an ultrasound system
158 (L9-3 transducer, iU22, Philips Ultrasound, Santa Ana, CA, USA) as described [28]. In short, a
159 transducer was positioned over the left hindlimb and secured for the course of the experiment. A
160 suspension of Optison microbubbles (Perflutren Protein-Type A Microspheres Injectable
161 Suspension, USP, GE Healthcare, USA) was infused intravenously (15 μ l/minute) using a Harvard
162 11 Plus low volume infusion pump (Harvard instrument Co., Holliston, MA). The infusion tube was
163 attached to a vortex mixer to ensure a homogeneous microbubble solution entering the animal. An
164 infusion time of 4 minutes was used where the first 2 minutes was to ensure systemic steady-state

165 conditions before three consecutive MVP recordings were performed. Data were exported to
166 quantification software (QLab, Philips, Andover, MA, USA) for analysis. Regions of interest were
167 drawn clear of connective tissue and large vessels and copied into each file to ensure that regions
168 were identical for each recording. Calculations were made in accordance with Wei et al. [29]. In
169 short, acoustic intensity (AI) versus time curves were fitted to the exponential function: $y = A(I -$
170 $exp(-\beta(t - Bt))$, where t is time (seconds), Bt is the time used for background subtraction, y is the
171 acoustic intensity at any given t , A is the plateau AI defined as MVP, and β is the flow rate constant
172 (liters·s-1) that determines the rate of rising AI.

173 *2.6. Basal and insulin-stimulated hepatic glucose production.* Mice (control, n=5, or tumor-bearing
174 mice (tumor size > 800 mm³), n=6) were clamped in randomized order after a 4 h fasting period
175 from 10:00 AM. Mice were anesthetized with an i.p. injection of 11 µl/g body weight of Fentanyl
176 (0.05 mg/ml, Dechra, DK), Midazolam (5 mg/ml, Accord Healthcare, UK) and Acepromazine (10
177 mg/ml, Pharmaxim, SE), and placed on a heating pad. A polyethylene cannula (PE50, Intramedic,
178 USA) was inserted into a jugular vein for administration of anesthetics, insulin, and glucose.
179 Anesthesia was maintained by constant infusion of the anesthetics (0.03 µl/g/min). After surgery, a
180 60 minutes continuous infusion (0.83 µl/minute, 1.2 µCi/h) of D-[3-3H]-glucose (Perkin Elmer)
181 was administrated. Then, a 120 minutes hyperinsulinemic-euglycemic clamp was initiated, with a
182 primed (4.5 mU) infusion of insulin (7.5 µU/kg/minute) (Actrapid, Novo Nordisk, DK) and D-[3-
183 3H]-glucose (0.83 µl/minute, 1.2 µCi/h). Blood glucose was clamped at 6 mmol/l and maintained
184 by a variable infusion of 20% glucose solution. Blood was sampled from the tail at -10, 0, 105, and
185 120 minutes for determination of plasma glucose, plasma 3H activity by scintillation counting, and
186 thereby the plasma specific activity. Basal and insulin-stimulated HGP were calculated based on the
187 equation described [30]. At 120 minutes, blood for plasma insulin concentration was also obtained
188 from the tail. Mice were humanely euthanized by cervical dislocation.

189 2.7. *Indirect calorimetry*. After seven days of acclimation to single-housing, the mice were
190 transferred to metabolic cages. Here, oxygen consumption and CO₂ production were measured by
191 indirect calorimetry in a CaloSys apparatus for 20 hours (TSE LabMaster V5.5.3, TSE Systems,
192 GER) to calculate respiratory exchange ratio (RER). Non tumor-bearing mice were injected with a
193 single dose of etomoxir (5 mg/kg body weight) or a vehicle control in a cross-over design with
194 injections separated by 24 hours (n=6). All mice were acclimatized to thermoneutrality (30°C) for 3
195 weeks prior to the measurements.

196 2.8. *Immunoblotting*. Mouse muscles was pulverized in liquid nitrogen and homogenized 2 × 0.5
197 minutes at 30 Hz using a TissueLyser II bead mill (Qiagen, USA) in ice-cold homogenization
198 buffer, pH 7.5 (10% glycerol, 1% NP-40, 20 mM sodium pyrophosphate, 150 mM NaCl, 50 mM
199 HEPES (pH 7.5), 20 mM β-glycerophosphate, 10 mM NaF, 2 mM phenylmethylsulfonyl fluoride
200 (PMSF), 1 mM EDTA (pH 8.0), 1 mM EGTA (pH 8.0), 2 mM Na₃VO₄, 10 µg/ml leupeptin, 10
201 µg/ml aprotinin, 3 mM benzamidine). After rotation end-over-end for 30 min at 4°C, supernatants
202 from muscle tissue were collected by centrifugation (10,000 rpm) for 20 minutes at 4°C. Lysate
203 protein concentrations were measured using the bicinchoninic acid method with bovine serum
204 albumin (BSA) as a standard (Pierce). Total proteins and phosphorylation levels of relevant proteins
205 were determined by standard immunoblotting techniques loading equal amounts of protein with a
206 standard curve included for all proteins to ensure quantifications within the linear range.
207 Polyvinylidene difluoride membranes (Immobilon Transfer Membrane; Millipore) were blocked in
208 Tris-buffered saline (TBS)-Tween 20 containing 2% milk for 10-20 minutes at room temperature.
209 Membranes were incubated with primary antibodies (Table 1) overnight at 4°C, followed by
210 incubation with HRP-conjugated secondary antibody for 45 minutes at room temperature.
211 Coomassie brilliant blue staining was used as a loading control [31]. Bands were visualized using
212 the Bio-Rad ChemiDoc MP Imaging System and enhanced chemiluminescence (ECL+; Amersham

213 Biosciences). Bands were quantified using Bio-Rad's Image Lab software 6.0.1
214 (RRID:SCR_014210).

215 Table 1. Primary antibodies

Antibody	Dilution (Primary)	Catalogue number	Company	RRID
AKT2	1:1000 (in 2% milk)	#3063	Cell Signaling Technology (CST)	AB_2225186
P-AKT1/2 ^{Ser473/474}	1:1000 (in 2% milk)	#9271	CST	AB_329825
P-AKT1/2 ^{Thr308/309}	1:1000 (in 2% milk)	#9275	CST	AB_329828
TBC1D4	1:1500 (in 2% milk)	ab189890	Abcam	–
P-TBC1D4 ^{Thr642}	1:1000 (in 2% milk)	#D27E6	CST	AB_2651042
Hexokinase II	1:1000 (in 2% milk)	#2867	CST	AB_2232946
GLUT4	1:1000 (in 2% milk)	#PA1-1065	Thermo Fisher Scientific	AB_2191454

216
217 2.9. *RNA extraction and real-time PCR.* RNA was isolated from 15 mg w.w. tumor tissue by
218 guanidinium thiocyanate-phenol-chloroform extraction method with modifications [32], and tissue
219 homogenized for 2 min at 30 Hz in a TissueLyser II (Qiagen, NL). cDNA was produced by reverse
220 transcribing 3 µg of RNA using superscript II reverse transcriptase (Invitrogen, USA), and the
221 samples diluted to 0.01 µg/ul. Content of interleukin 6 (IL-6), tumor necrosis factor α (TNF- α) and
222 TATA box binding protein (TBP) were determined by real-time PCR (ABI 7900 Prism, Applied
223 Biosystems, US). Sequences used to amplify a fragment of IL-6 were FP:
224 5'GCTTAATTACACATGTTCTCTGGAAA 3', RP: 5'CAAGTGCATCATCGTTGTTCATAC 3',
225 Taqman probe: 5'ATCAGAATTGCCATTGCACAACTCTTTCTCAT 3', and for TNF- α FP:
226 5'ATGGCCCAGACCCTCACA 3', RP: 5'TTGCTACGACGTGGGCTACA 3', Taqman probe:
227 5'TCAGATCATCTCTCAAAATTGAGTGACAAGC 3'. TBP was used as housekeeping gene.
228 The TBP probe was a pre-developed assay reagent from Applied Biosystems, US. TBP was similar
229 between groups. All TaqMan probes were 5'-6-carboxyfluorescein (FAM) and 3'-6-carboxy-

230 N,N,N',N'-tetramethylrhodamine (TAMRA) labeled (Applied Biosystems, US) except TATA-
231 binding protein (TBP) which was 5' FAM with minor groove binding.

232

233 *2.10. Statistical analyses.* Results are shown as mean \pm standard error of the mean (SE) with the
234 individual values shown for bar graphs or mean \pm SE for curve graphs. Statistical testing was
235 performed using t-test, one-way or two-way (repeated measures when appropriate) ANOVA as
236 applicable. Pearson's correlation was used to test the relationship between the tumor volume and
237 insulin-stimulated 2DG uptake or hepatic glucose production. Sidak post hoc test was performed for
238 all ANOVAs to test the difference between LLC groups and the non-tumor control group. A log-
239 transformation was performed in data-sets that were not normally distributed. Statistical analyses
240 were performed using GraphPad Prism, version 7 (GraphPad Software, La Jolla, CA, USA, RRID:
241 SCR_002798). For time-adjusted models of insulin-stimulated glucose uptake (related to Fig. 2) an
242 adjusted model (Pearson's correlations) was applied in IBM SPSS statistics 25 (IBM, USA, RRID:
243 SCR_002865). The significance level for all tests was set at $\alpha = 0.05$.

244

245

246 **3. Results**

247 **3.1. LLC caused adipose tissue wasting and increased spleen weight, indicative of mild 248 cachexia and inflammation**

249 Lewis lung carcinoma tumor-bearing day 15 mice had an average tumor volume of $530 \text{ mm}^3 \pm 285$
250 mm^3 . LLC day 21-27 mice had an average tumor volume of $1345 \text{ mm}^3 \pm 946 \text{ mm}^3$ (Fig. 1A). LLC
251 day 21-27 tumor-bearing mice displayed reduced body weight (-5%; Fig. 1B), which was due to fat
252 loss (Fig. 1C and D) rather than changes in lean body mass (Fig. 1E). Body weight was not
253 associated with tumor-size (Fig. 1F). Spleen weight increased in LLC day 15 (+40%) and LLC day

254 21-27 (+101%) (Fig. 1G) tumor-bearing mice, indicating elevated immunomodulatory activity
255 compared with control mice. This was confirmed by an average 7% increase in plasma levels of IL-
256 6 (Fig. 1H) and TNF- α (Fig. 1I) in tumor-bearing mice. Thus, LLC induced adipose tissue
257 wasting, increased spleen weight, and slightly elevated plasma cytokines, indicative of mild
258 cachexia and inflammation.

259

260 **3.2. LLC tumor-bearing mice displayed reduced blood glucose-lowering effect of insulin**

261 We next investigated the effect of cancer on whole-body insulin action by retro-orbitally injecting a
262 submaximal dose of insulin [33] and analyzing blood glucose in control and LLC tumor-bearing
263 mice (Fig. 2A). In contrast to control mice, insulin did not lower blood glucose in LLC day 15 and
264 LLC day 21-27 tumor-bearing mice 5 minutes following injection (Fig. 2B). Furthermore, the
265 blood-glucose-lowering effect of insulin was markedly reduced (by 2-2.5 mM) at 10 minutes in
266 LLC tumor-bearing mice compared to control mice (Fig. 2B). Accordingly, the area over the curve
267 (AOC) was 60-80% lower in tumor-bearing mice (Fig. 2B), indicative of marked insulin resistance.
268 Given the reduced whole-body insulin action, we measured glucose uptake in skeletal muscle and
269 WAT, tissues that account for the majority of glucose utilization involved in whole-body glycemic
270 control [34,35]. In gastrocnemius muscle, insulin-stimulated glucose uptake was reduced in LLC
271 day 15 tumor-bearing mice (-23%, Fig. 2C) and day 21-27 tumor-bearing mice (-28%, Fig. 2C)
272 compared with control mice. Likewise, in TA muscle, insulin-stimulated glucose uptake was
273 reduced in LLC day 21-27 tumor-bearing mice (-32%, Fig. 2D), but surprisingly, increased in LLC
274 day 15 tumor-bearing mice (+45%, Fig. 2D). In perigonadal WAT, insulin-stimulated glucose
275 uptake tended ($P=0.0784$) to be decreased by LLC at day 21-27 (-37%, Fig. 2E), while being
276 increased in LLC at day 15 (+48%, Fig. 2E). Basal non-stimulated glucose uptake (Fig. 2C-E) and

277 blood glucose (Fig. S1A) were unaffected by cancer. These findings show that cancer induces
278 obvious insulin resistance and markedly reduces muscle and adipose tissue glucose uptake.

279

280 **3.3. Insulin-stimulated glucose uptake in skeletal muscle and white adipose tissue negatively
281 correlated with tumor size**

282 As tissue-specific insulin resistance was more pronounced at day 21-27, we next investigated
283 whether insulin resistance was related to the tumor size. We re-divided our current data set into
284 small vs large tumors (cut off 800 mm³) across the LLC day 15 and LLC day 21-27 groups. In mice
285 with large tumors (LLC-L), the blood-glucose-lowering effect of insulin was almost abrogated (Fig.
286 2F). However, in mice with small tumors (LLC-S), the blood glucose-lowering effect of insulin was
287 only modestly reduced (Fig. 2F). Accordingly, AOC was 96% reduced in mice with large tumors
288 and tended (P=0.051) to be 47% decreased in mice with small tumors (Fig. 2F).

289 In LLC-S tumor-bearing mice, insulin-stimulated glucose uptake tended (P=0.051) to be 19%
290 decreased in gastrocnemius (Fig. 2G), increased in TA (+35%, Fig. 2H), while unaltered in WAT
291 (Fig. 2I) compared to control mice. In contrast, LLC-L tumor-bearing mice displayed a marked
292 reduction in insulin-stimulated glucose uptake in gastrocnemius (-32%, Fig. 2G), TA (-41%, Fig.
293 2H), and WAT (-44%, Fig. 2I). A negative correlation between tumor size and insulin-stimulated
294 glucose uptake in both skeletal muscle and WAT (Fig. 2J) confirmed that cancer can significantly
295 reduce insulin-stimulated glucose uptake in skeletal muscle and WAT in a tumor size-dependent
296 manner. As the size of the tumor increases with time, we adjusted the above model for the duration
297 of tumor-bearing. The observed negative correlation between tumor size and insulin-stimulated
298 glucose uptake persisted when taking tumor burden duration into account (Supplementary Table 1).
299 We observed no correlations between plasma IL-6 (Fig. S1B) or TNF- α (Fig. S1C) and insulin-
300 stimulated glucose uptake in skeletal muscle or WAT. In the tumor tissue the mRNA expression of

301 IL-6 and TNF- α was 60% and 20% higher, respectively, in day 15 compared to day 21-27 tumor-
302 bearing mice (Fig. S1D). Surprisingly, we observed a positive correlation between tumor IL-6 (Fig.
303 S1E) and TNF- α (Fig. S1F) expression and insulin-stimulated glucose uptake in TA and WAT.
304 Nonetheless, with no correlation between plasma cytokines and glucose uptake and a positive
305 correlation with the expression of tumor cytokines, systemic inflammation is likely not a main
306 driver of insulin resistance in tumor-bearing mice in the present study.

307 Previous studies have reported that tumors take up a substantial amount of glucose to support tumor
308 growth, migration, and invasion [36,37]. However, to the best of our knowledge, no study has to
309 date compared glucose uptake into muscle and tumor in the same mouse. We, therefore, analyzed
310 tumor glucose uptake and found that tumor glucose uptake per gram tumor mass was 5.9- and 1.7-
311 fold higher in LLC-S and LLC-L, respectively compared with average basal muscle glucose uptake
312 (Fig. 2K, $P<0.05$). Similar results were obtained in the insulin-stimulated state, where the average
313 muscle glucose uptake was 40% that of glucose uptake in tumors of LLC-S tumor-bearing mice ($p<$
314 0.01). Insulin-stimulated glucose uptake was similar in the tumor and muscle of LLC-L tumor-
315 bearing mice (Fig. 2K). Those findings suggest that the tumor significantly competes with skeletal
316 muscle for glucose. In order to understand how much the tumor contributed to whole-body glucose
317 disposal during the 10 minutes of stimulations, we produced an index of whole-body glucose
318 disposal in fat, skeletal muscle, and the tumor. It showed that in cancer, a substantial part of the
319 available glucose is directed from muscle and fat into the tumor (Fig. 2L).

320

321 **3.4. Reduced skeletal muscle glucose uptake in tumor-bearing mice was not due to impaired
322 AKT signaling**

323 Accounting for the majority of insulin-stimulated glucose disposal [35], we next determined
324 canonical insulin signaling in skeletal muscle to elucidate the molecular mechanisms underlying

325 LLC-induced insulin resistance. We focused our investigation on mice with tumor size of more than
326 800 mm³ (average tumor volume 1755 ± 763 mm³) as insulin resistance was pronounced in this
327 group. To our surprise, we found upregulated phosphorylation of AKT^{Ser474} (+65%, Fig. 3A),
328 AKT^{Thr309} (+86%) (Fig. 3B), and TBC1D4^{Thr642} (+18%, Fig. 3C), indicative of enhanced insulin
329 sensitivity. Protein expression of AKT2 (Fig. 3D), TBC1D4 (Fig. 3E), glucose transporter 4
330 (GLUT4) (Fig. 3F), and Hexokinase II (Fig. 3G) remained unaltered in tumor-bearing mice,
331 although we did detect a tendency (p=0.053) to increased protein expression for TBC1D4 in tumor-
332 bearing mice (representative blots are shown in in Fig. 3H-I).
333 These findings suggest that reduced insulin-stimulated muscle glucose uptake in tumor-bearing
334 mice is not due to decreased myocellular canonical insulin signaling, on the contrary, we observed
335 augmented phosphorylation of both AKT and TBC1D4.

336

337 **3.5. Cancer abrogated muscle microvascular perfusion in response to insulin**

338 Muscle microvascular perfusion (MVP) is essential for insulin to fully stimulate glucose uptake in
339 muscle, however, it is unknown whether cancer influences muscle MVP. Thus, we determined
340 muscle MVP in mice with tumor sizes > 800 mm³ (LLC-L, averaged tumor volume 1283 ± 207
341 mm³). In control mice, insulin increased muscle MVP (+70%; Fig. 4A and B) in accordance with
342 previous studies [38,39]. Remarkably, this increase was completely abrogated in LLC-L tumor-
343 bearing mice (Fig. 4C and D), showing that cancer negatively affects insulin-stimulated muscle
344 microvascular perfusion, which could contribute to muscle insulin resistance.

345

346 **3.6. Tumor-bearing mice exhibit increased basal hepatic glucose production**

347 Increased hepatic glucose production (HGP) is another hallmark of insulin resistance. Therefore, we
348 measured basal and insulin-stimulated HGP in mice with tumor sizes > 800 mm³ (LLC-L, averaged

349 tumor volume $3916 \pm 2196 \text{ mm}^3$) during a hyperinsulinemic-euglycemic clamp. Following 120
350 minutes of continuous insulin infusion (7.5 $\mu\text{U}/\text{kg}/\text{minute}$), blood glucose in both control and LLC-
351 L tumor-bearing mice was maintained at a steady level of 6 mM (Fig. 5A). Steady-state glucose
352 infusion rate (GIR) during the clamp was similar between control and tumor-bearing mice (Fig. 5B).
353 This is in contrast to our findings during the 10 minutes insulin stimulation where tumor-bearing
354 mice displayed reduced insulin response. This discrepancy might reflect the fact that the tumor
355 takes up a large proportion of the glucose in the tumor-bearing mice as indicated in Fig. 2K and L.
356 Over time that could mask smaller reductions in muscle and adipose tissue glucose uptake. It could
357 also reflect the longer stimulation time (the last 30 minutes of the clamp used to calculate GIR), or
358 that the insulin dose used to estimate GIR was higher compared with the r.o. injection.
359 Interestingly, basal HGP was 45% higher in LLC-L tumor-bearing mice compared to controls (Fig.
360 5C). In addition, basal HGP positively correlated with tumor volume (Fig. 5D). Insulin suppressed
361 HGP similarly in LLC-L tumor-bearing and control mice (Fig. 5C). Nevertheless, within the tumor-
362 bearing group, the inhibitory effect of insulin on HGP was negatively correlated with tumor volume
363 (Fig. 5E), although this should be interpreted cautiously, given the low number of mice.
364 Collectively, these findings show that cancer increases basal HGP, but does not affect insulin-
365 stimulated GIR or insulin-suppressed HGP at supra-physiological insulin levels.

366

367 **3.7. Inhibition of fatty acid oxidation partially restored LLC-induced insulin resistance and**
368 **inhibition of lipolysis improved glucose tolerance**

369 Augmented fatty acid metabolism, a hallmark of many insulin resistance conditions [16,40,41], has
370 been reported in human [42] and murine [19,20] cancer models. Therefore, we tested the hypothesis
371 that cancer might reduce insulin action via its effect on fatty acid metabolism in LLC tumor-bearing
372 mice (averaged tumor volume $645 \pm 386 \text{ mm}^3$). To inhibit fatty acid oxidation, we used the CPT1

373 inhibitor, etomoxir. In non-tumor-bearing mice, etomoxir treatment increased plasma free fatty
374 acids (Fig. 6A) and acutely increased RER (Fig. 6B) as would be expected by inhibition of fatty
375 acid oxidation [43]. In agreement with previous reports in cancer patients [13–15] and mouse
376 cancer models [19,20], we observed increased plasma free fatty acids (FFA) (+125%, Fig. 6C),
377 triacylglycerol (+23%, Fig. 6D) and glycerol (+40%, Fig. 6E) concentrations in LLC tumor-bearing
378 compared to control mice. Daily administration of etomoxir, showed that inhibition of whole-body
379 fatty acid oxidation restored blood free fatty acid concentrations to levels of control mice (Fig. 6C–
380 E). Plasma IL-6 (+17%, Fig. 6F) and TNF- α (p=0.09, +10%, Fig. 6G) were increased in tumor-
381 bearing mice. In contrast to the plasma lipid profile, etomoxir treatment did not affect the plasma
382 levels of the cytokines IL-6 and TNF- α (Fig. 6F–G). Similar spleen weight was observed tumor-
383 bearing mice with or without etomoxir treatment (Fig. S2A). Thus, inhibition of fatty acid oxidation,
384 at least partially, corrects abnormal lipid metabolism induced by cancer in mice as measured by
385 plasma triacylglycerol, free fatty acids, and glycerol.

386 Next, we investigated whether the inhibition of fatty acid oxidation improved the metabolic
387 dysfunction of tumor-bearing mice. In control mice, insulin lowered blood glucose by 2 mM and 4
388 mM following 5 and 10 minutes of stimulation, respectively (Fig. 6H). In agreement with our
389 previous observations in the present study, insulin action in LLC tumor-bearing mice was reduced
390 (Fig. 6H). Remarkably, etomoxir rescued insulin action in LLC tumor-bearing mice evidenced by
391 restored blood glucose levels (Fig. 6H) and 2.5-fold increase of AOC (Fig. 6H). We also analyzed
392 plasma insulin concentration 10 minutes following the r.o. insulin injection, as a marker of insulin
393 clearance. Interestingly, LLC tumor-bearing mice showed increased plasma insulin (+237%) (Fig.
394 6I), an indication of reduced insulin clearance, which is also observed in diet-induced insulin-
395 resistant mice [24,44]. Etomoxir administration normalized plasma insulin levels in LLC tumor-
396 bearing mice (Fig. 6I). In order to evaluate glycemic regulation, we undertook a glucose tolerance

397 test and found that tumor-bearing mice were glucose intolerant (Fig. 6J). Glucose intolerance was
398 not rescued by etomoxir (Fig. 6J), despite the improvements in circulating fatty acid levels and
399 insulin action. The glucose challenge increased plasma insulin levels 100-150% similarly in all
400 groups (Fig. 6K). Tumor volume was not affected by etomoxir treatment (Fig. S2B) and food intake
401 was similar between all groups (Fig. S2C). Etomoxir did not affect insulin's blood glucose lowering
402 effect in non-tumor-bearing mice (Fig. S2D), suggesting that the benefits of inhibiting fatty acid
403 oxidation on insulin sensitivity were specific to the context of cancer-induced accelerated fatty acid
404 metabolism.

405

406 We next inhibited adipose tissue lipolysis by a potent inhibitor, nicotinic acid. Nicotinic acid did not
407 increase insulin's blood glucose-lowering effect in non-tumor-bearing mice (Fig. S2E), nor did it
408 affect glucose tolerance (Fig. S2F). Tumor size and growth rate were not affected by nicotinic acid
409 treatment (Fig. S2G) and spleen weight increased similarly in tumor-bearing mice with or without
410 nicotinic acid treatment (Fig. S2H). In contrast to etomoxir, the blood glucose response to r.o.
411 injected insulin was not improved by nicotinic acid (Fig. 6L). However, nicotinic acid did restore
412 glucose tolerance in LLC-tumor-bearing mice (Fig. 6M). The glucose challenge increased plasma
413 insulin levels similarly (100-150%) in all groups (Fig. 6N), suggesting that altered insulin
414 sensitivity rather than insulin levels caused the improvement in glucose tolerance by nicotinic acid.
415 Taken together, these findings demonstrate that altered fatty acid metabolism, but likely not
416 inflammation, is involved in LLC-induced insulin resistance and glucose intolerance.

417

418 **4. Discussion**

419 Cancer resulted in marked perturbations on at least six metabolically essential functions; i) insulin's
420 blood-glucose-lowering effect, ii) glucose tolerance, iii) skeletal muscle and WAT insulin-

421 stimulated glucose uptake, iv) intramyocellular insulin signaling, v) muscle microvascular perfusion,
422 and vi) basal hepatic glucose production in mice (depicted in Fig. 7). Additionally, we show that the
423 mechanism causing cancer-induced insulin resistance may relate to altered fatty acid metabolism
424 but is likely not related to inflammation.

425 A major finding in the current study was the significantly impaired insulin-stimulated glucose
426 uptake in both skeletal muscle and adipose tissue in LLC tumor-bearing mice. These findings
427 suggest that skeletal muscle and adipose tissue are major players in dysregulated glucose
428 metabolism often observed in human cancers and murine cancer models [45–47]. In accordance,
429 reduced insulin-stimulated glucose disposal has also been reported in cachexic patients with lung
430 cancer [8] and lymphoma [48], suggesting that our findings are clinically relevant and translatable
431 to the human situation. Tumor size seemed to be a key factor in peripheral insulin resistance, as we
432 observed the skeletal muscle and WAT glucose uptake to be negatively correlated with tumor size.
433 Conversely, inflammation seemed not to be involved in cancer-associated insulin resistance, as
434 plasma IL-6 and TNF- α did not correlate with glucose uptake rates. Interestingly, the decreased
435 insulin-stimulated glucose uptake in mice with large tumors was not due to diminished proximal
436 insulin signaling in muscle. On the contrary, insulin-stimulated AKT/TBC1D4 signaling was
437 upregulated in skeletal muscle of tumor-bearing mice. This is surprising, given that tumor-bearing
438 mice displayed increased whole-body inflammation as indicated by increased spleen volume and
439 increased plasma levels of IL-6 and TNF- α , consistent with other investigations showing increased
440 IL-6 in patients with non-small cell lung cancers [49,50]. Inflammation would be expected to
441 reduce insulin signaling in muscle [51]. Tumorkines, such as VEGF and HIF-1, are reported to be
442 upregulated in the LLC model of cancer [52,53]. Although not analyzed in our study, those
443 tumorkines could increase insulin signaling in muscle, as they have been reported to increase
444 PI3K/AKT signaling in cancer cells [37,54]. The causes of upregulated muscle insulin signaling in

445 cancer warrants further investigation but increased insulin signaling in muscles with reduced
446 insulin-stimulated glucose uptake has been reported in other models [55]. Nevertheless, the
447 mechanisms by which cancer causes insulin resistance seems to be different from the mechanisms
448 causing insulin resistance in for example obesity and type 2 diabetes, where muscle AKT and
449 TBC1D4 signaling is either unaffected [56,57] or reduced [58–60]. Furthermore, inflammation did
450 not seem to be a main driver of insulin resistance, as spleen volume, plasma IL-6, and plasma TNF-
451 α were similarly increased in mice with small and large tumors, of which only the mice with large
452 tumors displayed reduced insulin-stimulated glucose uptake in muscle and WAT.

453 Another major finding of the present investigation was that insulin-stimulated muscle microvascular
454 perfusion was abrogated in tumor-bearing mice. To our knowledge, these data are the first to show
455 that dysregulated muscle microvascular perfusion is involved in a common model of cancer and
456 cachexia. Insulin-stimulated microvascular perfusion in muscle is a critical facet in glucose uptake
457 regulation [10,39,61–63]. In agreement, genetic or pharmacological inhibition of microvascular
458 perfusion impaired insulin-stimulated muscle glucose uptake by 40% in otherwise healthy mice
459 [38]. In addition, insulin-stimulated microvascular perfusion is reduced in different insulin-resistant
460 conditions, including obesity and type 2 diabetes [64,65]. In cancer, impaired microvascular
461 perfusion could be caused by elevated circulating fatty acid levels, as experimentally elevated
462 circulating fatty acids reduced insulin-stimulated muscle microvascular perfusion by 40% [66,67]
463 without causing impairments in intracellular insulin signaling in healthy humans [17]. Our findings
464 show that decreased microvascular perfusion could contribute to cancer-induced impaired muscle
465 glucose uptake in response to insulin.

466 In our study, a reduction in skeletal muscle and WAT glucose uptake likely contributed to the
467 attenuated blood-glucose-lowering effect of insulin in tumor-bearing mice. On the other hand, Lang
468 et al [68] have previously reported that insulin resistance in tumor-bearing rats was due to an

469 impaired ability of insulin to suppress hepatic glucose production, although that study did not
470 analyze muscle and adipose tissue glucose uptake. In the present study, insulin's inhibitory effect
471 on hepatic glucose production was not impaired, while basal hepatic glucose production was 45%
472 increased. Increased basal hepatic glucose production has also been reported in patients with lung
473 cancer [69] but the mechanisms remain to be established.

474 Elevation of fatty acids is often associated with insulin resistance [16–18] and increased plasma free
475 fatty acid concentrations are reported in cancers [19,20,46,70]. For example, the release of fatty
476 acids and glycerol from WAT explants was 30-40% increased in LLC or B16 tumor-bearing mice
477 [19]. Increased circulating fatty acids have been shown to induce insulin resistance [71]. Indeed, we
478 found that whole-body insulin action was restored by blocking fatty acid oxidation via etomoxir
479 administration in tumor-bearing mice. Furthermore, lipolysis inhibition via nicotinic acid
480 administration rescued glucose intolerance in tumor-bearing mice. Interestingly, etomoxir
481 normalized plasma triacylglycerol, fatty acids, and glycerol concentrations in tumor-bearing mice,
482 which could benefit insulin action. The plasma fatty-acid-lowering effect of etomoxir in tumor-
483 bearing mice is somewhat surprising, as fatty acid oxidation inhibition in non-tumor-bearing mice
484 increased circulating fatty acid levels in our present and other studies [72,73], and prolonged FA
485 oxidation inhibition thus increased plasma triacylglycerol and liver fat content [43]. However,
486 etomoxir inhibits lipolysis in adipocytes and can increase re-esterification of fatty acids to
487 triacylglycerol in the liver, thereby diminishing the release of fatty acids from the adipose tissue and
488 triacylglycerol from the liver [74]. In the cancer condition with accelerated lipolysis [13,15], it is
489 possible that etomoxir's effect on triacylglycerol metabolism seen in liver and lipolysis overrules
490 the contrary effect of etomoxir on reduced fatty acid oxidation. Etomoxir has also been reported to
491 reduce inflammation [75,76], which could also contribute to the amelioration of cancer-induced
492 insulin resistance. However, spleen weight and plasma IL-6 and TNF- α levels increased similarly in

493 tumor-bearing mice with or without etomoxir treatment, suggesting that etomoxir did not prevent
494 cancer-induced inflammation. Indeed, our results indicate that the impact of cancer on lipid
495 metabolism exerts a critical impact on the pathology of cancer-induced insulin resistance.

496 Based on the present investigation and relevant literature [13–18,77], we hypothesize a model
497 where the tumor secretes tumorkines that increase fatty acid metabolism, which in turn leads to
498 peripheral insulin resistance. Redirecting glucose from skeletal muscle and adipose tissue, likely
499 benefits the tumor’s energy demand to support tumor growth, migration, and invasion [36]. The
500 clinical relevance of this is suggested, as cancer patients with type 2 diabetes have increased
501 mortality rates [2].

502 There are certain limitations to the present investigation and unresolved questions. Firstly, our
503 investigation was undertaken in a mouse model of lung cancer with a subcutaneously implanted
504 tumor. Whether our results apply to the human condition and other types of cancer remain to be
505 determined. However, patients with lung cancer have been reported to display reduced insulin-
506 stimulated glucose uptake [8] and accelerated hepatic glucose production [69], suggesting that our
507 findings do mimic the human condition. Secondly, insulin resistance was observed without loss of
508 body mass. While this provides a good model for investigating the effect of cancer isolated from the
509 presence of cachexia, we cannot draw any conclusions regarding the involvement of insulin
510 resistance in cancer cachexia or *vice versa* [4–6]. Thirdly, our study shows that MVP was markedly
511 compromised by cancer. However, whether this was due to altered fatty acid metabolism remains
512 undetermined and future studies should investigate the effect of etomoxir and nicotinic acid on
513 MVP. Treatment with pharmacological vasodilators have shown similar [78] or increased [79]
514 muscle glucose uptake, thus, whether vasodilators could improve insulin sensitivity in cancer is also
515 an unresolved question to be answered by future investigations.

516 In conclusion, cancer impaired the blood-glucose-lowering effect of insulin, caused glucose
517 intolerance, and reduced glucose uptake in muscle and WAT. Furthermore, tumor-bearing mice
518 displayed increased basal hepatic glucose production. Cancer-associated insulin resistance was
519 neither due to inflammation nor impaired proximal muscle insulin signaling, but was associated
520 with a complete abrogation of insulin-stimulated muscle microvascular perfusion. Finally, we
521 identify fatty acid metabolism as a player in cancer-associated insulin resistance, providing
522 potential therapeutic targets for cancer-induced insulin resistance. These findings suggest that
523 insulin resistance is likely of key importance in the therapy of cancer.

524

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547
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761 **Figure legends:**

762 **Figure 1.** Characteristics of Lewis lung carcinoma (LLC) tumor-bearing mice. **A)** Tumor volume,
763 **B)** body weight, **C)** perigonadal white adipose tissue (WAT) weight, **D)** fat mass, **E)** lean body
764 mass, **F)** correlation between tumor volume and body weight, and **G)** spleen weight in control mice
765 (n=25-28) and LLC tumor-bearing mice following 15 (n=14-15) or 21-27 days (n=19-22) tumor
766 inoculation. The weights of all tissues were normalized to body weight with tumor weight
767 subtracted. Plasma concentration of interleukin-6 (IL-6) (**H)** and tumor necrosis factor α (TNF- α) (**I**)
768 in control and tumor-bearing mice. Statistically significant effect of LLC on body composition or
769 tissue weights is indicated by (*) $P<0.1$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$. Values are shown as
770 mean \pm SE with individual values.

771 **Figure 2.** Insulin sensitivity in Lewis lung carcinoma (LLC) tumor-bearing mice. **A)** Experimental
772 overview **B)** Blood glucose levels measured before (0 minutes), 5 minutes, and 10 minutes
773 following retro-orbital (r.o.) insulin injection (0.3 U/kg body weight) and area over the curve (AOC)
774 (n=22 in control group; n=13/14 in LLC groups) during the 10 minutes insulin stimulation. **C)**
775 Basal (n=2-8) and insulin- (n=16-21 in control group; n=11-14 in LLC groups) stimulated 2-deoxy-
776 glucose (2DG) uptake in gastrocnemius (Gas) muscle, **D)** tibialis anterior (TA) muscle, and **E)**
777 perigonadal white adipose tissue (WAT) following 15 days or 21-27 days tumor inoculation.
778 Data were re-divided into mice with large tumor volume (>800 mm 3 ; LLC-L) and small tumors
779 (<800 mm 3 ; LLC -S). **F)** Blood glucose levels measured before (0 minutes), 5 minutes, and 10
780 minutes following r.o. insulin injection and AOC (n= 22 in control group; n= 13/14 in LLC groups)
781 during the 10 minutes of insulin stimulation. **G)** Basal (n=4/6) and insulin (n= 16-21 in control
782 group; n= 10-14 in LLC groups) stimulated 2DG uptake in Gas, **H)** TA, and **I)** WAT. **J)**
783 Correlation between tumor volume and insulin-stimulated 2DG uptake in Gas, TA and WAT
784 (n=24-26). **K)** Basal (n=4-5) and insulin- (n=12-14) stimulated glucose uptake in the tumor

785 compared with muscle. **L)** Index of the contribution of muscle, fat, and tumor tissue to whole-body
786 glucose uptake. Statistically significant effect of LLC on whole-body insulin action at each
787 timepoint and 2DG uptake is indicated by (*)P<0.1; *P < 0.05; **P < 0.01; ***P < 0.001.
788 Statistically significance on whole-body insulin action at 10 minutes between LLC-S and LLC-L is
789 indicated by ††† P < 0.001. Statistically significant effect of insulin on 2DG uptake is indicated by
790 ☒ P < 0.01; ☒☒P < 0.001. Values are shown as mean± SE with or without individual values.

791 **Figure 3.** Effect of Lewis lung carcinoma on insulin-stimulated signaling in gastrocnemius muscle
792 of mice with large tumors (>800mm³; LLC-L). **A)** Basal (n=6) and insulin- (n=10-12) stimulated
793 phosphorylated (P)-AKT^{Ser474}, **B)** P-AKT^{Thr309}, **C)** P-TBC1D4^{Thr642}. **D)** Protein expression of AKT2,
794 **E)** TBC1D4, **F)** GLUT4, and **G)** Hexokinases II (n=16-18). **H)** Representative phospho-blots. **I)**
795 Representative blots of total proteins. Statistically significant effect of LLC-L on insulin signaling
796 is indicated by (*)P<0,1; *P < 0.05; **P < 0.01; ***P < 0.001. Statistically significant effect of
797 insulin is indicated by ☒ P < 0.05; ☒☒P < 0.001. Values are shown as mean ± SE with individual
798 values. A.U., arbitrary units.

799 **Figure 4.** Effect of Lewis lung carcinoma on skeletal muscle microvascular perfusion in mice with
800 large tumors (>800mm³; LLC-L). **A)** Microvascular refilling curves after microbubbles destruction
801 **B)** microvascular perfusion presented as the plateau AI value in adductor magnus and
802 semimembranosus muscles in control (**A** and **B**) and LLC-L (**C** and **D**) tumor-bearing mice at
803 baseline and after 60 minutes of insulin (7.5 μ U /kg/minute) infusion (n=6). Statistically significant
804 effect of insulin on microvascular perfusion is indicated by #P < 0.05; ###P < 0.001. Values are
805 shown as mean or mean ± SE with individual values.

806 **Figure 5.** Effect of Lewis lung carcinoma on hepatic glucose production (HGP) in mice with large
807 tumors (>800mm³; LLC-L). **A)** Blood glucose, **B)** glucose infusion rate (GIR), **C)** basal or insulin-
808 stimulated HGP in control mice (n=5) and LLC-L tumor-bearing mice (n=6), **D)** correlation

809 between tumor volume and basal or **E**) insulin-stimulated HGP during hyperinsulinemic-
810 euglycemic clamp (7.5 μ U /kg/minute). Statistically significant effect of LLC-L on basal HGP is
811 indicated by **P < 0.01. Values are shown as mean \pm SE with or without individual values.

812 **Figure 6.** Effect of etomoxir on insulin sensitivity and glucose tolerance. **A)** acute effect of
813 etomoxir treatment in non-tumor-bearing mice (n=8). **B)** acute effect of etomoxir treatment on
814 respiratory exchange ration (RER) (n=6). **C)** Plasma free fatty acid (FFA), **D)** triacylglycerol, **E)**
815 glycerol, **F)** plasma interleukin-6 (IL-6), and **G)** plasma tumor necrosis factor α (TNF- α) in control
816 mice and LLC tumor-bearing mice with or without etomoxir (Eto) administration (n=11-15). **H)**
817 Blood glucose levels before (0 minutes), 5 minutes and 10 minutes following retro-orbital insulin
818 injection (0.3 U/kg body weight) and area over the curve (AOC). **I)** Plasma insulin (n=9-12). **J)**
819 Blood glucose levels before (0 minutes), 20 minutes, 40 minutes, 60 minutes and 90 minutes
820 following an intraperitoneal glucose tolerance test (2 g/kg body weight) and area under the curve
821 (AUC) (n=6-9). **K)** Plasma insulin levels at 0 minutes and 20 minutes following intraperitoneal
822 glucose tolerance test (2 g/kg body weight) (n=6-9). **L)** Blood glucose concentration before (0
823 minutes), 5 minutes and 10 minutes following retro-orbital insulin injection (0.3 U/kg body weight)
824 in control mice or Lewis lung carcinoma (LLC) tumor-bearing mice with or without nicotinic acid
825 (Nico) administration (n=8-12), and area over the curve (AOC). **M)** Blood glucose concentration
826 before (0 minutes), 20 minutes, 40 minutes, 60 minutes and 90 minutes following intraperitoneal
827 glucose tolerance test (GTT; 2 g kg⁻¹ body weight) in control and LLC tumor-bearing mice with or
828 without nicotinic acid (Nico) administration and area under the curve (AUC). **N)** plasma insulin
829 concentration at 0 minutes and 20 minutes into the GTT (n=6-8). Statistically significant effect of
830 LLC is indicated by *P < 0.05; **P < 0.01; ***P < 0.001. Statistically significant effect of Eto/Nico
831 is indicated by #P < 0.05; ##P < 0.01; ###P < 0.001. Values are shown as mean \pm SE with or
832 without individual values.

833 **Figure 7.** Graphic overview of Lewis lung cancer-induced metabolic perturbations identified by the
834 present study. Created with BioRender.com.

835 **Figure supplementary 1.** **A)** Blood glucose levels measured before (0 minutes), 5 minutes, and 10
836 minutes following retro-orbital saline injection in control (n=6) or Lewis lung carcinoma (LLC)
837 tumor-bearing mice following 15 days (n=2) or 21-27 days (n=8) tumor inoculation. **B)** and **C)**
838 Pearsons correlations between insulin-stimulated 2-DG uptake and plasma levels of interleukin-6
839 (IL-6) and tumor necrosis factor α (TNF- α) in gastrocnemius muscle (Gas), tibialis anterior muscle
840 (TA), and white adipose tissue (WAT). **D)** mRNA expression of tumor tissue IL-6 and TNF- α . **E)**
841 Pearson's correlations between insulin-stimulated 2-DG uptake in peripheral tissues and tumor
842 expression of IL-6. **E)** Pearson's correlations between insulin-stimulated 2-DG uptake in peripheral
843 tissues and tumor expression of TNF- α . Values are shown as mean \pm SE.

844 **Figure supplementary 2.** **A)** Spleen weight in control or LLC tumor-bearing mice following
845 etomoxir (Eto) administration (n=11-14). **B)** Tumor volume of LLC tumor-bearing mice with Eto
846 administration (n=11-14). **C)** Food intake in control or LLC tumor-bearing mice with Eto
847 administration (n=3-6). **D)** Blood glucose concentration before (0 minutes), 5 minutes, 10 minutes
848 and 15 minutes following retro-orbital insulin injection (0.3 U/kg body weight) in non-tumor-
849 bearing control mice treated with or without Eto. **E)** Blood glucose concentration before (0 minutes),
850 5 minutes, 10 minutes and 15 minutes following retro-orbital insulin injection (0.3 U/kg body
851 weight) in non-tumor-bearing control mice treated with or without Nicotinic acid (Nico). **F)** Blood
852 glucose concentration before (0 minutes), 20 minutes, 40 minutes, 60 minutes and 90 minutes
853 following intraperitoneal glucose tolerance test (GTT; 2 g kg⁻¹ body weight) in control mice with
854 or without Eto or Nico administration. **G)** Tumor volume and **H)** spleen weight in control or LLC
855 tumor-bearing mice with Nico administration (n=8-12). Statistically significant effect of LLC is
856 indicated by *P < 0.05; **P < 0.01; ***P < 0.001. Statistically significant effect of Nico in tumor-

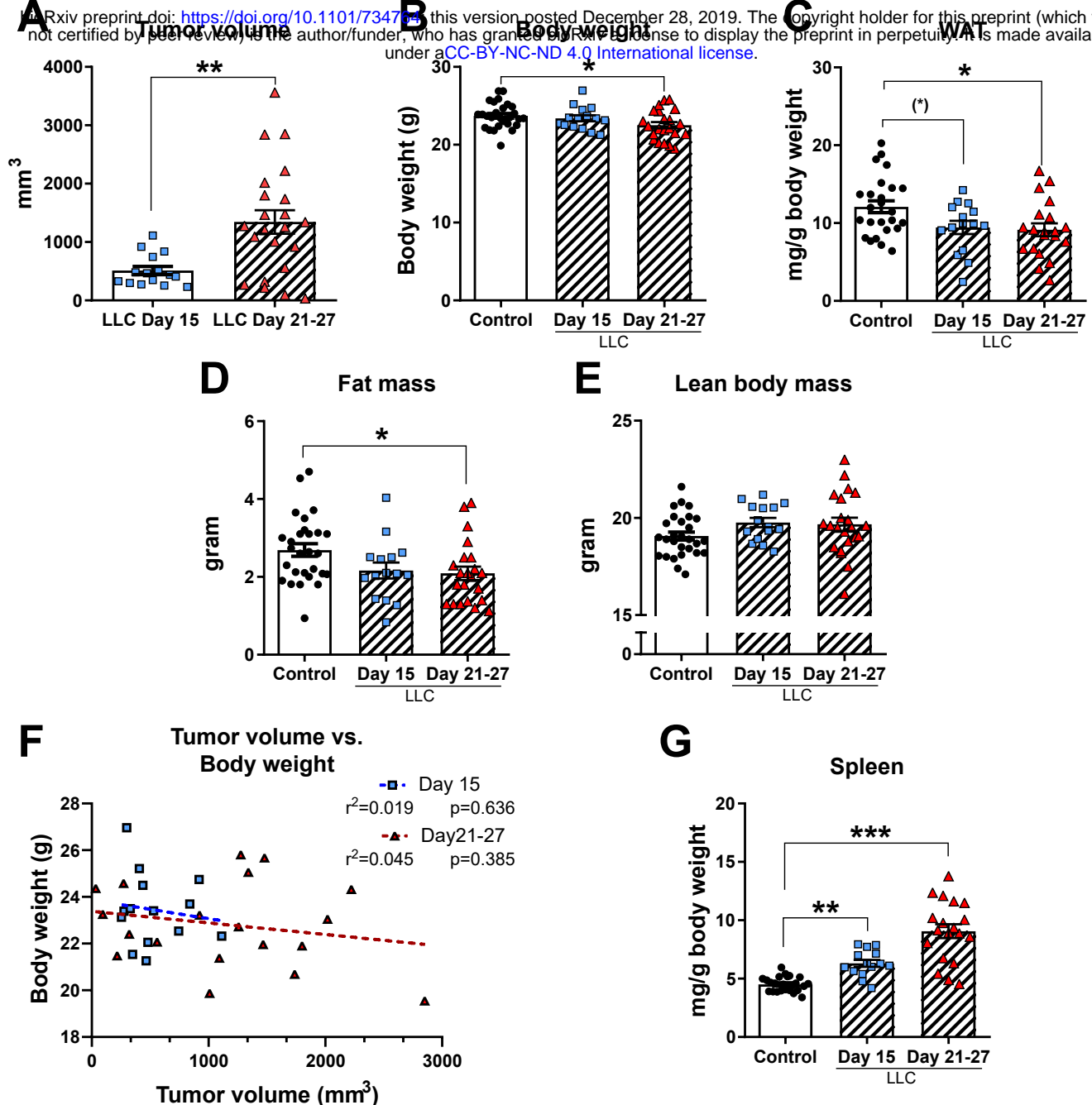
857 bearing mice is indicated by ##P < 0.01. Statistically significant effect of glucose injection on
858 plasma insulin is indicated by \$\$\$P < 0.001. Values are shown as mean \pm SE with or without
859 individual values.

860 **Supplementary Table 1**

861 Pearson`s correlations of tissue-specific glucose uptake, tumor volume, body weight, and time with
862 (“Time”) or without (“none”) adjustment for time.

Figure 1

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Cytokines

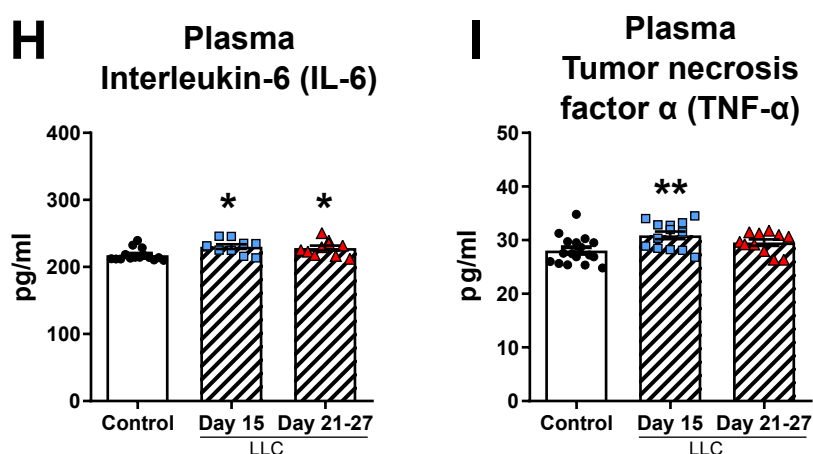
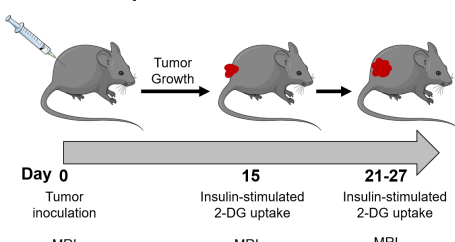


Figure 2

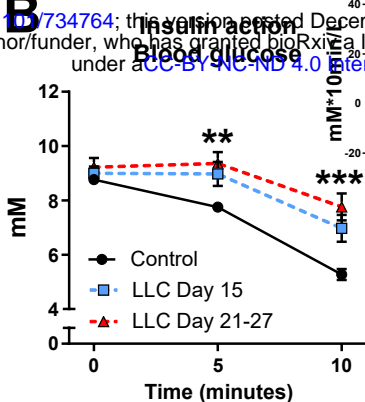
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Experimental overview



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AOC

Blood glucose

$\text{mM} \times 10 \text{ min/L}$

Time

Control Day 15 Day 21-27

LLC

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Figure 3

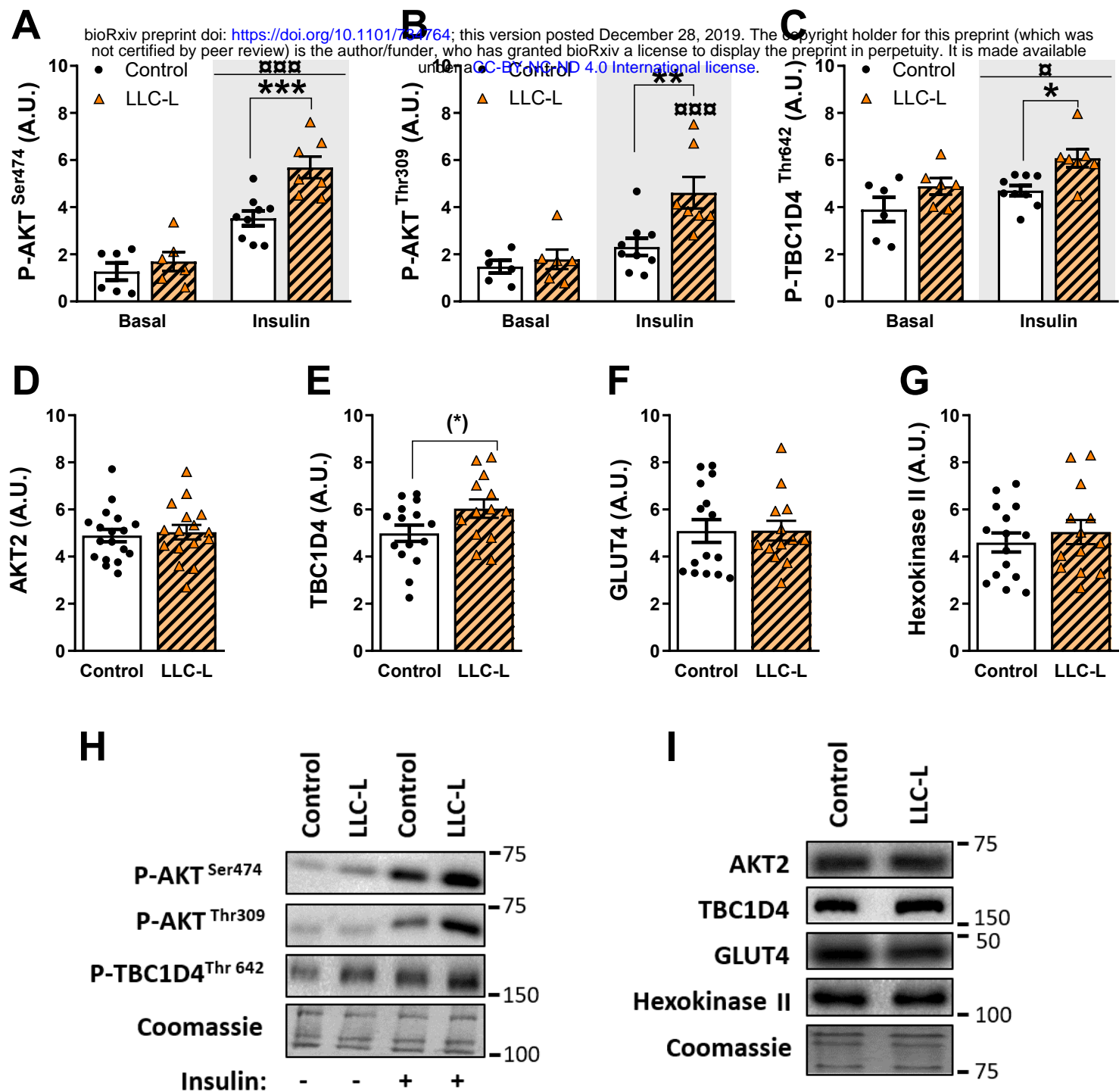


Figure 4

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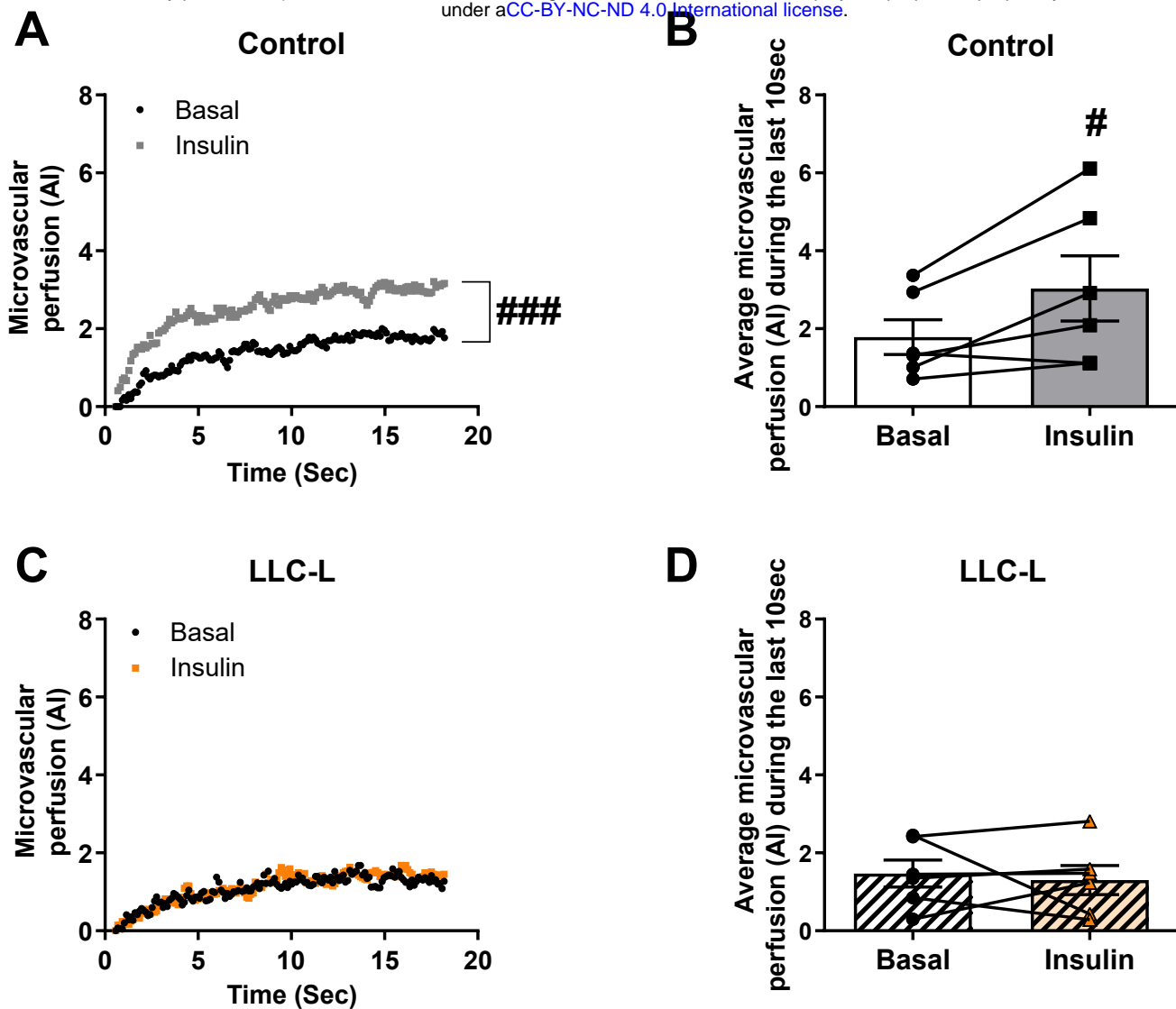
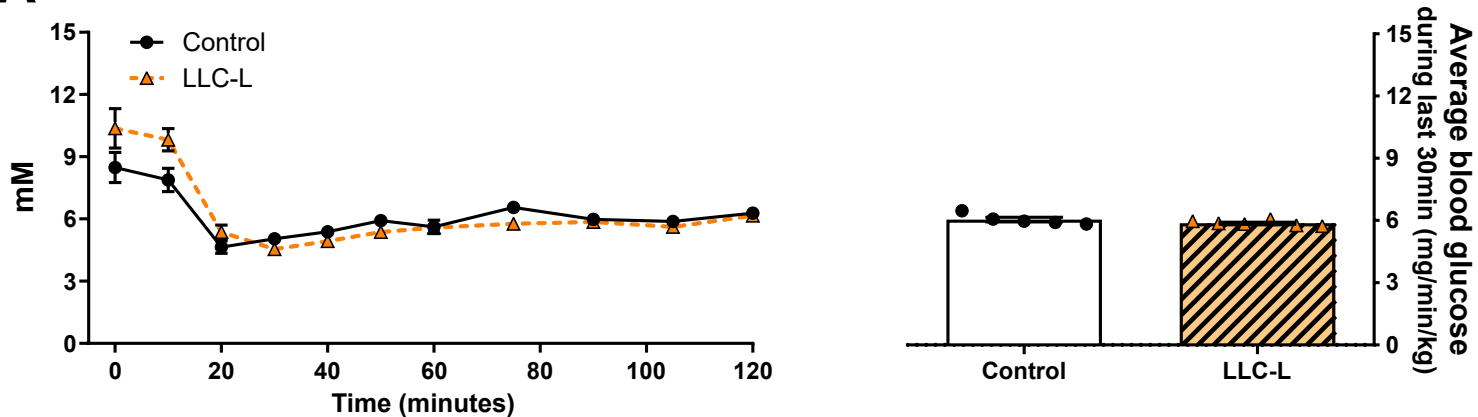


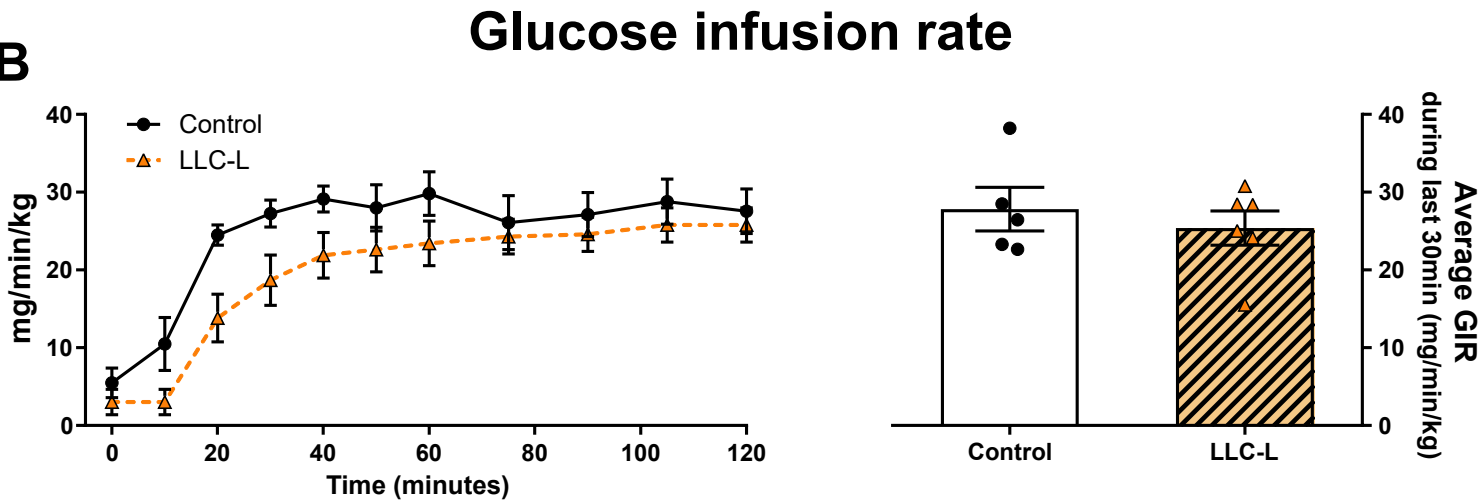
Figure 5

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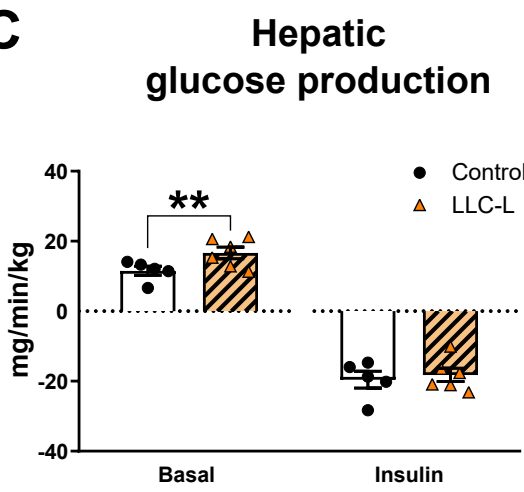
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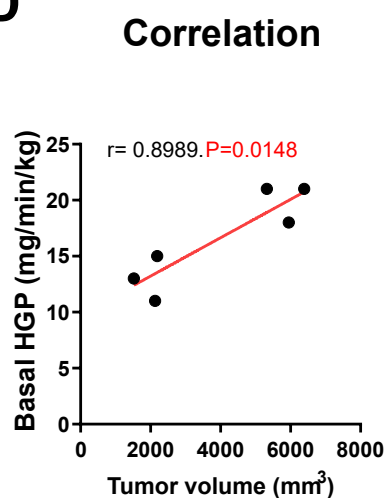
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D



E

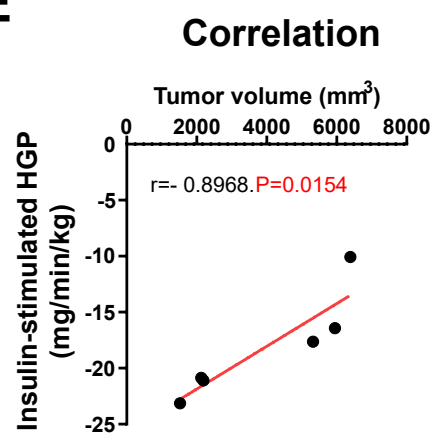
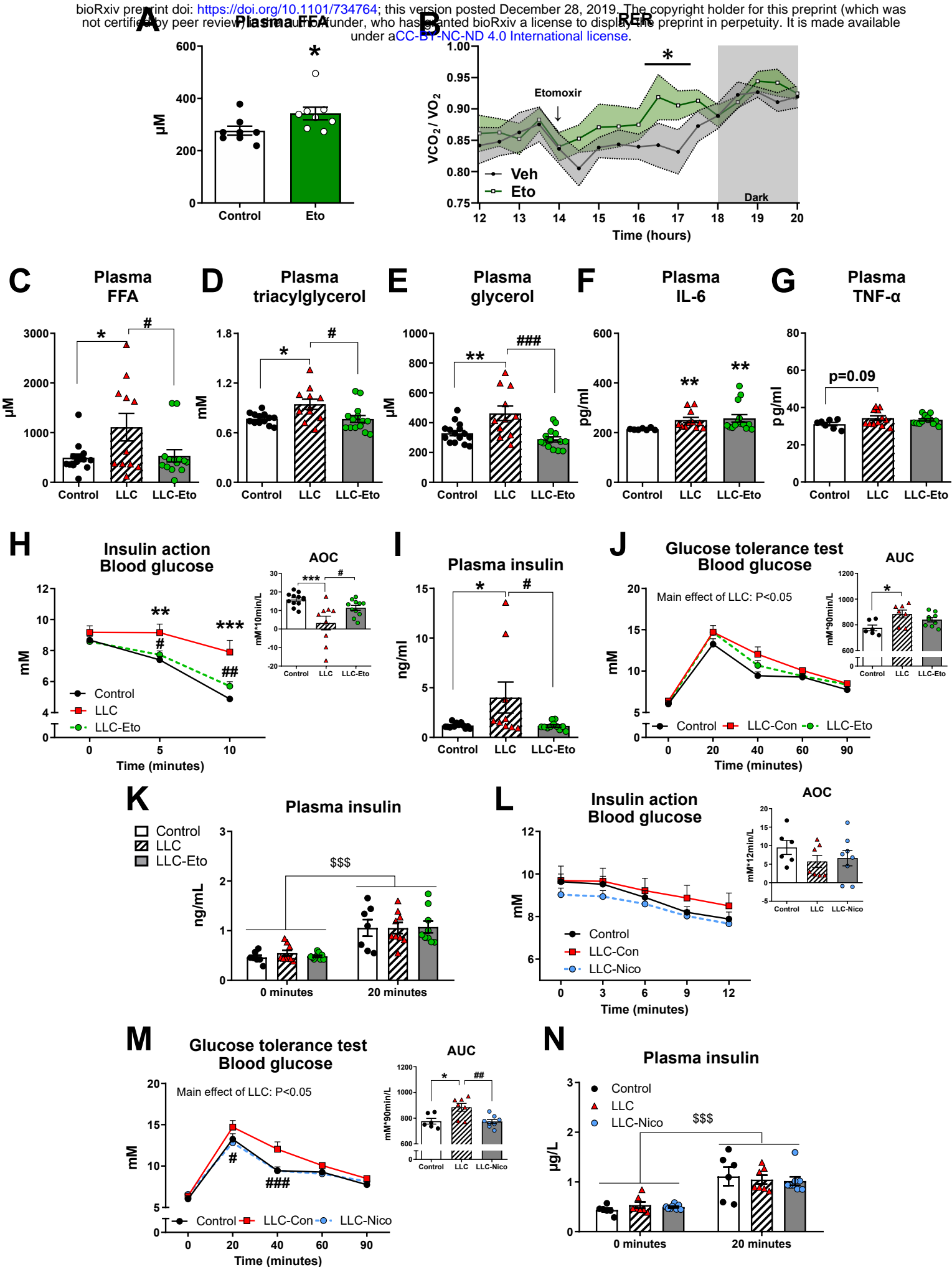
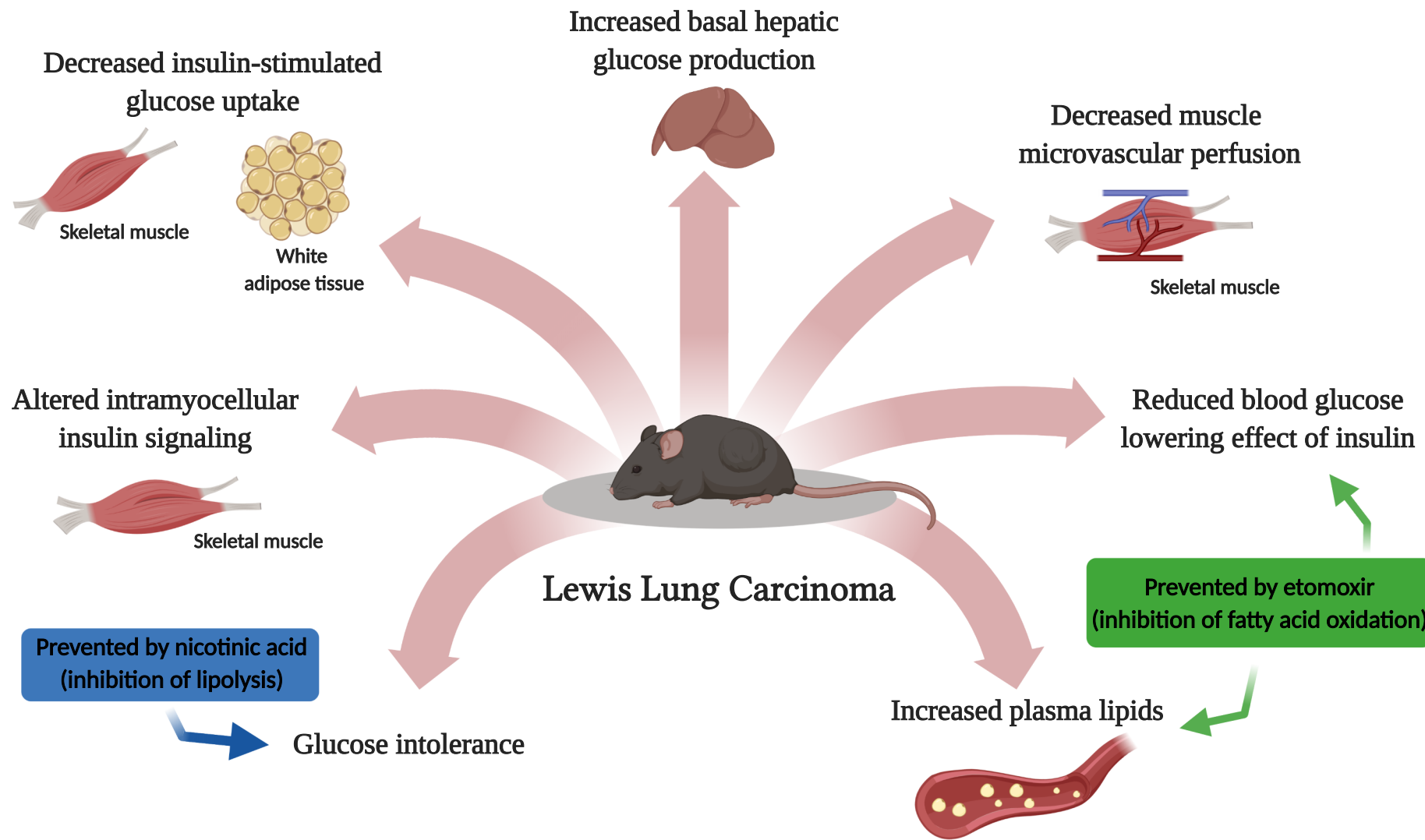


Figure 6

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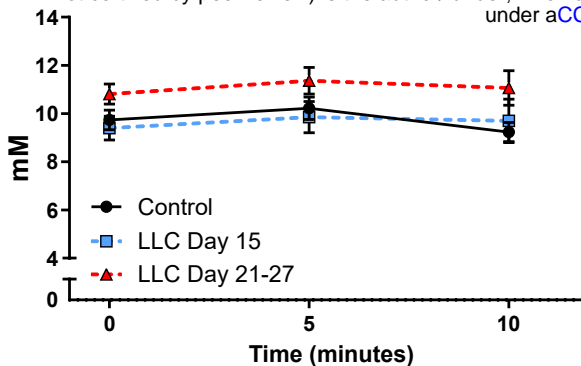
Cancer-induced metabolic perturbation



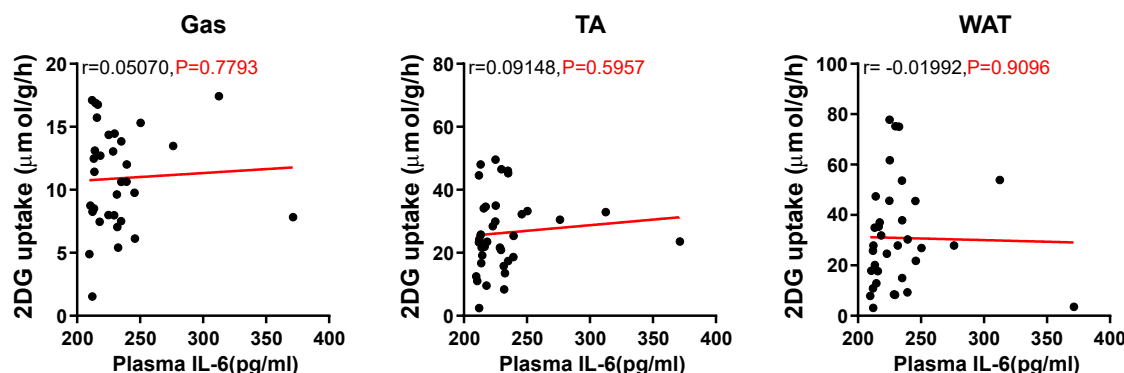
Supplementary Figure 1 (S1)

A

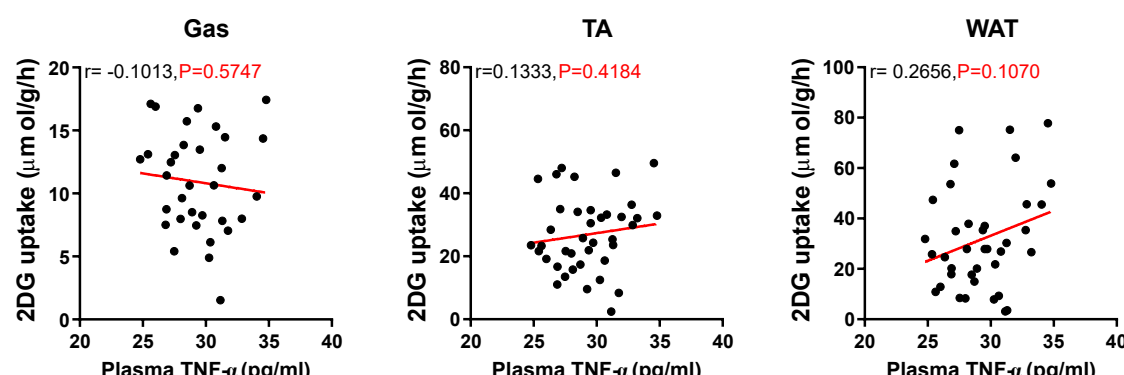
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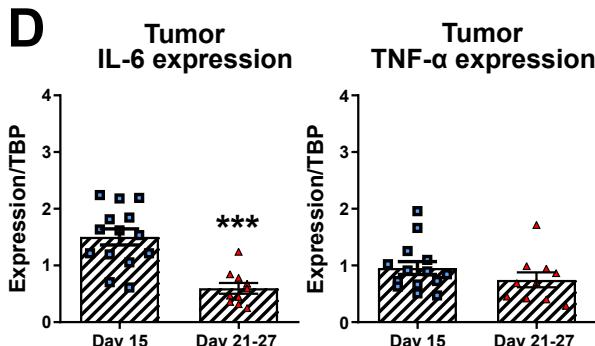
B



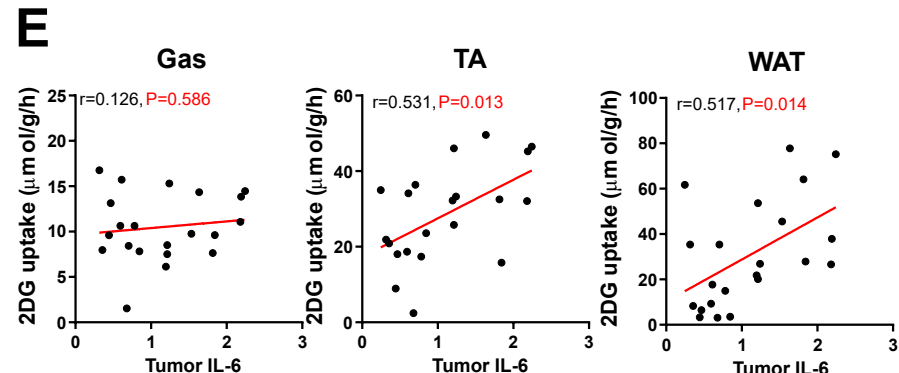
C



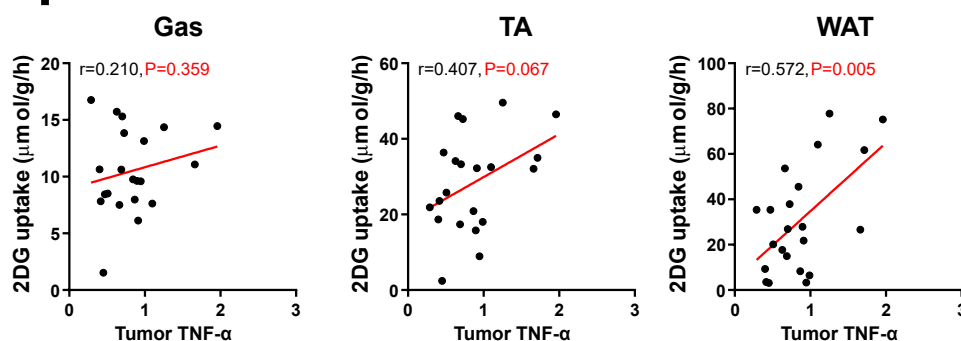
D



E



F

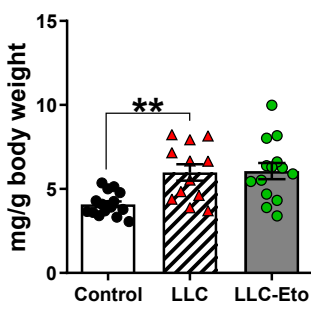


Supplementary Figure 2 (S2)

A

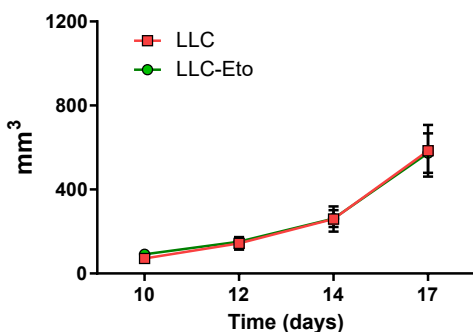
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Spleen



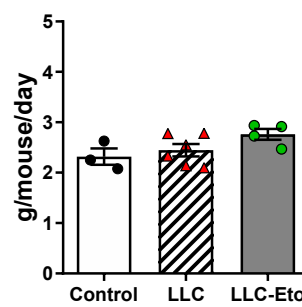
B

Tumor volume



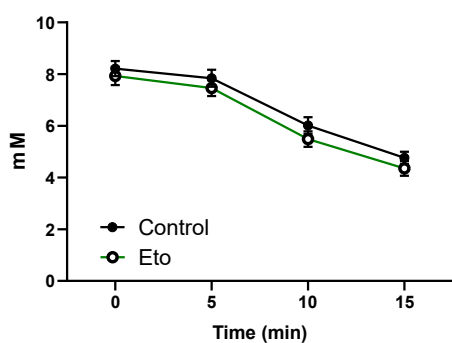
C

Food intake



D

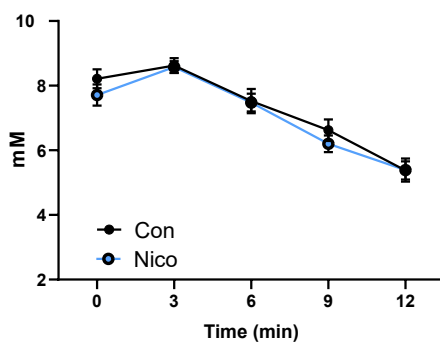
Insulin action
Blood glucose



Chronic Nicotinic Acid Treatment (inhibition of lipolysis)

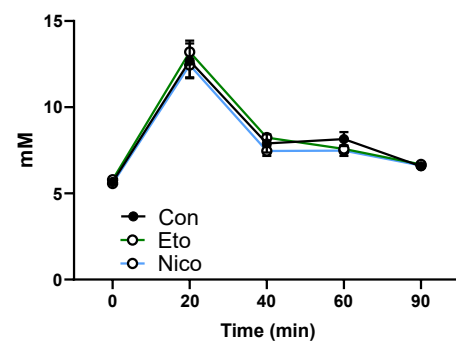
E

Insulin action
Blood glucose



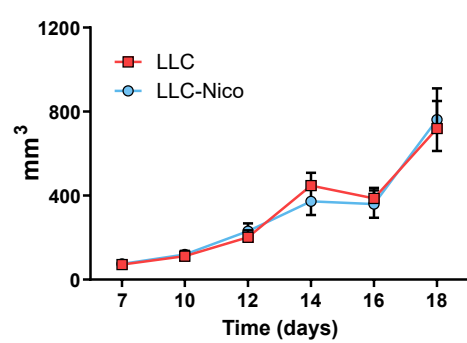
F

Glucose tolerance test
Blood glucose



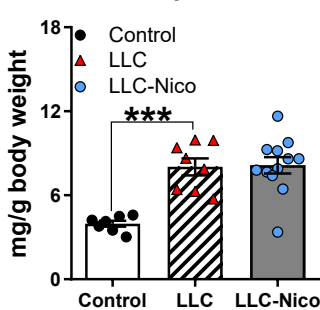
G

Tumor volume



H

Spleen



Supplementary Table 1

Control		Variables		Pearson correlations					
				Glucose uptake_TA	Tumor volume	Glucose uptake_Gas	Glucose uptake_WAT	Body weight	Time
-none- ^a	Glucose uptake_TA	Correlation	1,000	-,726	,463	,783	,013	-,617	
		p-value	.	,000	,026	,000	,948	,001	
		df	0	24	21	21	24	24	
	Tumor volume	Correlation	-,726	1,000	-,401	-,583	,042	,391	
		p-value	,000	.	,052	,003	,834	,044	
		df	24	0	22	22	25	25	
	Glucose uptake_Gas	Correlation	,463	-,401	1,000	,281	,260	-,033	
		p-value	,026	,052	.	,206	,221	,879	
		df	21	22	0	20	22	22	
	Glucose uptake_WAT	Correlation	,783	-,583	,281	1,000	,013	-,442	
		p-value	,000	,003	,206	.	,951	,031	
		df	21	22	20	0	22	22	
	Body weight	Correlation	,013	,042	,260	,013	1,000	,080	
		p-value	,948	,834	,221	,951	.	,693	
		df	24	25	22	22	0	25	
	Time	Correlation	-,617	,391	-,033	-,442	,080	1,000	
		p-value	,001	,044	,879	,031	,693	.	
		df	24	25	22	22	25	0	
Time	Glucose uptake_TA	Correlation	1,000	-,669	,563	,723	,080		
		p-value	.	,000	,006	,000	,705		
		df	0	23	20	20	23		
	Tumor volume	Correlation	-,669	1,000	-,422	-,497	,012		
		p-value	,000	.	,045	,016	,953		
		df	23	0	21	21	24		
	Glucose uptake_Gas	Correlation	,563	-,422	1,000	,297	,263		
		p-value	,006	,045	.	,191	,225		
		df	20	21	0	19	21		
	Glucose uptake_WAT	Correlation	,723	-,497	,297	1,000	,054		
		p-value	,000	,016	,191	.	,806		
		df	20	21	19	0	21		
	Body weight	Correlation	,080	,012	,263	,054	1,000		
		p-value	,705	,953	,225	,806	.		
		df	23	24	21	21	0		