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Voluntary intake of paracetamol-enriched drinking water and its influence on the success of embryo transfer in mice

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Abstract: Embryo transfer (ET) in mice is a key technique in biomedical research, and is carried out mostly via surgery by transferring founder embryos into pseudo-pregnant recipient females. To cover post-operative analgesic requirements in surrogate mothers, oral self-administration of painkillers has several advantages, but its effectiveness has also been criticized as voluntary ingestion of the drug can be uncertain. Additionally, concerns about potential negative side effects of analgesics on embryo viability and development have been raised. In this regard, we investigated the impact of orally administered analgesia by comparing the outcome of ET with and without paracetamol in the drinking water (3.5 mg/ml) of surrogate mothers. Water intake increased significantly when paracetamol, as a sweet-tasting formulation (children's syrup), was added to the drinking water. Measurements of paracetamol concentrations in blood serum confirmed reasonable drug uptake. Success rate of ETs and the body weight of newborn offspring were not different whether paracetamol was administered for two days after surgery or not. In conclusion, paracetamol in drinking water was consumed voluntarily in substantial doses, without detectable side-effects, by freshly operated surrogate mothers, and can therefore be recommended as a feasible method for providing analgesic treatment for surgical ET in mice.

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1 **Title:**

2 **Voluntary intake of paracetamol-enriched drinking water and its influence on**
3 **the success of embryo transfer in mice**

4
5 **Short title:** Paracetamol for embryo transfer

6
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33 **Abstract**

34 Embryo transfer (ET) in mice is a key technique in biomedical research, and is
35 carried out mostly via surgery by transferring founder embryos into pseudo-pregnant
36 recipient females. To cover post-operative analgesic requirements in surrogate
37 mothers, oral self-administration of painkillers has several advantages, but its
38 effectiveness has also been criticized as voluntary ingestion of the drug can be
39 uncertain. Additionally, concerns about potential negative side effects of analgesics
40 on embryo viability and development have been raised. In this regard, we
41 investigated the impact of orally administered analgesia by comparing the outcome
42 of ET with and without paracetamol in the drinking water (3.5 mg/ml) of surrogate
43 mothers. Water intake increased significantly when paracetamol, as a sweet-tasting
44 formulation (children's syrup), was added to the drinking water. Measurements of
45 paracetamol concentrations in blood serum confirmed reasonable drug uptake.
46 Success rate of ETs and the body weight of newborn offspring were not different
47 whether paracetamol was administered for two days after surgery or not. In
48 conclusion, paracetamol in drinking water was consumed voluntarily in substantial
49 doses, without detectable side-effects, by freshly operated surrogate mothers, and
50 can therefore be recommended as a feasible method for providing analgesic
51 treatment for surgical ET in mice.

52

53 **Keywords:**

54 Acetaminophen; Paracetamol; Embryo transfer; Water intake; Mice

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57

58 **1. Introduction**

59 The transfer of isolated embryos into pseudo-pregnant surrogate mothers
60 represents a basic routine procedure for establishing new genetically modified
61 mouse lines, and is used routinely for rederivation of pathogen-contaminated lines
62 or revitalization of archived strains. Embryo transfer (ET) in mice is commonly
63 conducted by a surgical approach (Nagy et al., 2003). Therefore, a laparotomy is
64 performed, i.e. the abdominal cavity is opened under sterile conditions, and the
65 oviduct or uterus horns are exposed for the transfer of embryos, aided by visual
66 control using a microscope. Surgical ET is performed under general anaesthesia,
67 and intra- as well as post-operative pain relief in animals undergoing such invasive
68 surgery is an essential refinement to avoid unnecessary pain and suffering of the
69 affected animals.

70 Recently, new ET techniques, avoiding the need for surgery and post-operative pain
71 treatment, have been published (Bin Ali et al., 2014; Cui et al., 2014; Steele et al.,
72 2013). In these procedures, embryos (only blastocysts are recommended) are
73 transferred with specialized instruments through the cervix into the uterine horn in
74 conscious surrogate mothers. Although the specifically designed devices required
75 are already available on the market, most laboratories prefer a surgical approach,
76 as it allows for reliable evidence of pseudo-pregnancy by the direct observation of
77 a swollen ampulla or corpora lutea in recipient ovaries. Furthermore, the accurate
78 transfer of the appropriate developmental stages into the oviduct or uterus horns
79 can be confirmed visually. Surgical ET has proven to be a reliable and efficient
80 technique for decades. However, when a surgical approach is chosen, post-
81 interventional analgesia needs to be applied for alleviation of post-operative pain,
82 which can persist for 1–2 days after surgery. The choice of an appropriate pain
83 control should take into account an easy and reliable mode of application, preferably

84 with the longest possible analgesic affect, but without any negative effects on
85 embryo development or gestation.

86 Regarding any potential influence on the success rate of ET, different effects have
87 been reported for opioid analgesics. While morphine treatment hampered blastocyst
88 implantation and decreased uterine receptivity (Tang et al., 2015), administering a
89 single dose of buprenorphine during ET surgery did not increase embryonic loss
90 compared to untreated animals (Goulding et al., 2010), and the number of
91 successfully implanted embryos was even greater compared to untreated mice
92 (Krueger and Fujiwara, 2008). Also, application of tramadol after ET in mice did not
93 affect success rate outcomes, and may even have improved pup survival as birth
94 rates and body weight in animals receiving tramadol did not differ from untreated
95 animals, whereas the number of offspring was slightly increased in animals treated
96 with this type of analgesic (Koutroli et al., 2014).

97 Non-steroidal anti-inflammatory drugs (NSAIDs) are generally not recommended for
98 pain treatment during pregnancy (Nolan, 2000). However, in mice, flunixin treatment
99 was not associated with increased embryonic loss after ET (Goulding et al., 2010).

100 In another study, application of tolfenamic acid or flunixin led to a higher pregnancy
101 rate and higher numbers of offspring than in animals undergoing ET without
102 analgesic treatment (Schlapp et al., 2015). A report on multimodal analgesia
103 (recipient female mice received carprofen together with buprenorphine) also
104 showed no significant adverse effects on the results of ET in mice (Parker et al.,
105 2011).

106 Besides potential side-effects on gestation and embryo development, duration of
107 analgesic action and route of application are the main criteria when choosing an
108 appropriate pain relief protocol for mice. In small rodents, analgesics are applied
109 mainly by intraperitoneal (i.p.) or subcutaneous (s.c.) injection. With opioids,

110 however, the duration of action is rather short, and injections have to be repeated
111 several times per day to ensure constant analgesic efficacy (Jirkof et al., 2015).
112 Some NSAIDs are known to induce longer lasting pain relief compared to opioids
113 and may need to be injected only once or twice per day (Flecknell, 1984; Miller and
114 Richardson, 2011). However, mice generally experience stress in response to
115 immobilization and injections (Cinelli et al., 2007; Meijer et al., 2005; Meijer et al.,
116 2006). Therefore, oral self-medication represents a promising alternative to provide
117 stress-free post-operative analgesia. The advantages of oral self-administration via
118 drinking water (or food items) are the considerable reduction in stress and potential
119 pain that might be caused by handling and restraining of mice with fresh wounds.
120 However, food neophobia, where animals abstain from the consumption of
121 unfamiliar substances or food, is a well-known behaviour in small rodents (Bauer et
122 al., 2003). Moreover, food or water intake can be decreased after surgery, thus
123 latency to consume analgesics voluntarily could be prolonged, resulting in insufficient
124 post-operative pain relief. Consequently, when adding drugs to food or drinking
125 water, it is advisable to examine whether sufficient amounts of the medicated food
126 or water are in fact consumed voluntarily over time.

127 In human medicine, paracetamol (acetaminophen) has become a popular and
128 widely used non-opioid drug for treatment of fever, as well as for acute and chronic
129 pain management (Allegaert et al., 2014; Mattia and Coluzzi, 2009; Raffa et al.,
130 2004). While the mechanism of action remains partly unknown, selective inhibition
131 of cyclooxygenase enzymes, as well as interaction with endogenous opioid
132 pathways are unique features of paracetamol. Paracetamol is considered to have
133 analgesic and antipyretic, rather than anti-inflammatory, effects compared to typical
134 NSAIDs (Mattia and Coluzzi, 2009). Its intake in therapeutic dosages is generally
135 regarded as safe in a variety of patients, also in pregnant women, where the use of

136 other NSAIDs is contraindicated due to potential risk to the unborn child
137 (Aminoshariae and Khan, 2015). However, when overdosed, paracetamol can
138 cause liver injuries, triggered by the hepatotoxic effect of its metabolites (Mattia and
139 Coluzzi, 2009).

140 Paracetamol is also recommended for pain relief in laboratory animals (Flecknell,
141 1984; Miller and Richardson, 2011). Acetaminophen was shown to increase the pain
142 threshold in rats (Mickley et al., 2006) and to be effective on bone cancer pain (Saito
143 et al., 2005) or to show a potent, synergistic effect when combined with morphine or
144 NSAIDs in mice (Miranda et al., 2006; Saito et al., 2005). The drug can be
145 administered easily by various routes, e.g. by adding to drinking water (Hayes et al.,
146 2000; Mickley et al., 2006). This makes it an ideal drug for broad application in
147 laboratories when opioids are not considered necessary, or are not available.

148 In the present study, we investigated the analgesic paracetamol as a means of pain
149 management after surgical ET in mice by adding it to the drinking water. The aim of
150 the present study was to determine whether paracetamol in drinking water would be
151 taken up voluntarily by mice in amounts sufficient to cover post-operative analgesic
152 requirements after laparotomy without any detrimental effect on the ET success
153 rate.

154

155 **2. Materials and Methods**

156 *2.1. Ethics statement*

157 Animal housing and the experimental protocols were approved by the Cantonal
158 Veterinary Office, Zurich, Switzerland, and were in accordance with Swiss Animal
159 Protection Law. Housing and experimental procedures were also conform to
160 *European Directive 2010/63/EU of the European Parliament, and of the Council of*

161 *22 September 2010 on the Protection of Animals used for Scientific Purposes* and
162 *to the Guide for the Care and Use of Laboratory Animals* (2010/63/EU, 2010;
163 Balingier et al., 2011).

164 A preliminary investigation was undertaken to exclude adverse effects of a
165 standardized pain treatment protocol with paracetamol in surrogate mothers during
166 ET. Later, at the request of animal welfare officers and authorities, further
167 investigation was performed to confirm the usefulness and reliability of the
168 administration route, i.e. offering the drug for voluntary uptake. Mice used in the
169 present study were surplus animals from our in-house breeding colony. To reduce
170 animal numbers, no dose response studies or analgesiometric testing were
171 conducted. Since experiments were performed at different time points, surrogate
172 mothers or naïve female mice involved in the study varied with respect to their
173 genetic background, i.e. mice of different outbred stocks were used in the two parts
174 of the study.

175

176 *2.2. Animals and housing conditions*

177 The animal facility provided standardized housing conditions, with a mean room
178 temperature of $21 \pm 1^\circ\text{C}$, relative humidity of $50 \pm 5\%$, and 15 complete changes of
179 filtered air per hour (HEPA H 14 filter); air pressure was controlled at 50 Pa. The
180 light/dark cycle in the animal rooms was set to a 12h/12h cycle (lights on at 07:00,
181 lights off at 19:00) with artificial light of approximately 40 Lux in the cage. Mice were
182 housed in a barrier-protected specific pathogen-free unit and were kept in
183 Eurostandard Type III open-top plastic cages (425 mm × 266 mm × 155 mm, floor
184 area 820 qcm; Techniplast, Indulab, Gams, Switzerland) with autoclaved dust-free
185 sawdust bedding (80–90 g per cage, LTE E-001 Abedd; Indulab, Gams,

186 Switzerland). A standard cardboard house (Ketchum Manufacturing, Brockville,
187 Canada) served as a shelter, and tissue papers were provided as nesting material.
188 The animals had unrestricted access to sterilized drinking water, and ad libitum
189 access to a pelleted and extruded mouse diet in the food hopper (Kliba No. 3436;
190 Provimi Kliba, Kaiseraugst, Switzerland). To avoid any possible interference from
191 external factors, all necessary husbandry and management procedures were
192 completed in the room at least 1 day before starting the experiment, and
193 disturbances (e.g., unrelated experimental procedures) were not allowed.

194 The specific pathogen-free status of the animals was monitored frequently and
195 confirmed according to FELASA guidelines throughout the experiments by a
196 sentinel program. The mice were free of all viral, bacterial, and parasitic pathogens
197 listed in FELASA recommendations (Mahler et al., 2015).

198 For measurements of water intake and paracetamol concentrations in blood serum,
199 40 female, naïve Crl:CD-1 mice, 8–16 weeks old, were used. Naïve mice were
200 housed in groups of four to eight prior to the study. During baseline measurements
201 and experiments mice were housed individually.

202 To determine the impact of paracetamol on the outcome of ET, 15 female Zbz:FM
203 mice were used as embryo recipients. The surrogate mothers were 8–16 weeks old
204 when ET was performed. They were housed in groups of two to six animals until
205 mating with vasectomized Zbz:FM males. Mating took place between 16:00 to 17:00
206 to induce pseudo-pregnancy. Vaginal plug positive females were isolated on the
207 next morning and subsequently housed individually. Two-cell stage embryos were
208 obtained after standard superovulation of B6D2F1 females, mated with Zbz:FM
209 males according to standard protocols (Rulicke, 2004). Briefly, female mice were
210 treated at about 16:00 by intraperitoneal injection of 5 IU pregnant mare serum
211 gonadotrophin (PMSG, Folligon; Intervet, Boxmeer, the Netherlands), followed 48

212 hrs later by 5 IU human chorionic gonadotrophin (hCG, Pregnyl; Organon AG,
213 Pfäffikon SZ, Switzerland) and mated. About 40 hrs later, treated females were killed
214 by cervical dislocation and two-cell embryos were flushed from both excised
215 oviducts; embryos were stored in an incubator at 37°C, 5% CO₂ in air using M16
216 medium (Sigma-Aldrich, St. Louis, Missouri, USA) until embryo transfer on the same
217 day.

218

219 *2.3. Experiment set up and data acquisition*

220 The schedule for the experimental procedure of both parts of the study is shown in
221 Fig. 1.

222

223 *2.3.1. Naïve mice: Water intake and paracetamol in blood serum*

224

225 *Treatment groups:*

226 Forty naïve female mice were randomly allocated into five groups, each group
227 consisting of eight animals: three groups received paracetamol in the drinking water
228 (PW 1–3). In order to compare serum concentrations between voluntary uptake in
229 drinking water and other ascertained administration routes, two further groups
230 received paracetamol either via oral gavage (group G) or via i.p. injection (group I).
231 These two groups, with paracetamol administered as bolus, served as control
232 groups.

233

234 *Treatment protocol:*

235 Paracetamol was provided in the drinking water according to the recommended
236 published dosage (Flecknell, 2009; Miller and Richardson, 2011). The amount of
237 paracetamol in drinking water was calculated with the intention to provide the mice

238 with 200 mg/kg body weight (BW) paracetamol over 24 hrs. Assuming that the water
239 consumption of adult outbred mice is at least 3 ml per day, 28 ml paracetamol syrup,
240 formulated to be applied per orally in children (Dafalgan® Children's Syrup, 30
241 mg/ml; Bristol-Myers Squibb SA, Steinhausen, Switzerland) was diluted in 212 ml
242 tap water, resulting in a final concentration of 3.5 mg paracetamol per ml drinking
243 water. One hour after onset of the light phase (08:00), mice were provided with a
244 freshly prepared bottle of paracetamol-containing water for 6 hrs (group PW1), 11
245 hrs (group PW2) or 24 hrs (group PW3).

246 In control group G, the same dose of paracetamol (200 mg/kg BW) (Dafalgan®
247 Children's Syrup, 30 mg/ml; Bristol-Myers Squibb SA, Steinhausen, Switzerland)
248 was given at 12:00 per gavage as bolus with a tube directly into the stomach of the
249 mice. In control group I, the same dose of paracetamol (200 mg/kg BW) was given
250 i.p. at 12:00 by using a formulation intended for injection delivery (Perfalgan®, 500
251 mg/ml; Bristol-Myers Squibb SA, Steinhausen, Switzerland). Mice in the control
252 groups were provided with untreated drinking water *ad libitum*.

253

254 *Water intake:*

255 Water intake was determined by weighing the drinking bottle at 6 hrs (PW1), 11 hrs
256 (PW2) and 24 hrs (PW3) after provision of paracetamol-containing water (day 1).
257 Baseline measurements of water intake without paracetamol in the drinking water
258 were taken in the same mice at the identical time points the day before (day 0) (Fig.
259 1).

260

261 *Blood sampling and paracetamol serum concentration measurements:*

262 In groups PW1–3, blood was sampled once per animal, either at 6 hrs (14:00; PW1),
263 11 hrs (19:00; PW2), or 24 hrs (08:00; PW3) after paracetamol was provided for

264 voluntary intake with drinking water. In control groups I and G, blood was sampled
265 at 2 hrs (14:00) after administering paracetamol in a single dose via gavage (G) or
266 i.p. injection (I). All mice were bled under sevoflurane anaesthesia and killed after
267 the procedure. Blood was centrifuged and the serum stored at -20°C until further
268 analysis. Paracetamol serum concentrations were determined by DRI®
269 Paracetamol-Serum-Tox-Assay by our in-house laboratory (Institute for Clinical
270 Chemistry, University Hospital Zurich, Switzerland).

271

272

273 *2.3.2. Surrogate mothers: Water intake and reproductive parameters*

274

275 *Treatment groups:*

276 Fifteen female mice were randomly allocated to either the untreated (n=8) or the
277 paracetamol treated (n=7) group.

278

279 *Embryo transfer procedure:*

280 After monogamous mating with vasectomized males at 16:00 to 17:00 on the day
281 prior to ET (day 0), females were checked for successful mating on the following
282 morning (day 1) between 07:00 and 07:30 by vaginal plug control. Plug positive
283 females were assumed to be pseudo-pregnant, and were housed individually in
284 fresh cages. At 08:00 on day 1, they received either a fresh water bottle without
285 medication (n=8) or a preemptive bottle with medicated drinking water (n=7) (Fig.
286 1).

287 At 13:00 on day 1, pseudo-pregnant females were transferred from the animal room
288 to the nearby laboratory. Anaesthesia and ET were carried out in a biosafety
289 cabinet, which was equipped with a water-bath-heated operating surface (38°C) and

290 an inhalation anaesthesia device, as described in detail previously (Rulicke, 2004).
291 Briefly, anaesthesia was induced by restraining the mouse and holding its nose in a
292 cone delivering Sevoflurane ($\leq 8\%$ in oxygen at a flow of 200 ml/min). After 15–
293 20 seconds, loss of protective reflexes was checked (e.g., pedal withdrawal reflex)
294 and the back of the anesthetized animal was shaved and disinfected.
295 ET was conducted bilaterally under aseptic conditions. The skin was cut in the
296 midline of the back, the abdomen was opened by a small incision in the peritoneum
297 near the ovary, and the reproductive tract was pulled out. Six two-cell stage embryos
298 were transferred in M2 medium (Sigma-Aldrich, St. Louis, Missouri, USA) via the
299 infundibulum in the ampulla on each side, so that each recipient received 12
300 embryos. After placing the tract back in the abdominal cavity, the peritoneum was
301 sutured with absorbable threads and the skin closed with staples. The anaesthetic
302 gas was then stopped and 100% oxygen was supplied to the animal, which
303 subsequently regained reflexes within 2–3 minutes and started to move away from
304 the face mask. Anaesthesia and ET were completed within 15–20 minutes. The
305 animal was allowed to recover for approximately 10 minutes on the warm surface in
306 the biosafety hood, under a filter cup to prevent it escaping. After recovery, the
307 mouse was returned to its cage and brought back to the animal room. Preparation
308 of embryos, anaesthesia and ET was performed by the same technician, who was
309 blinded to the treatment regimens. Animals of the untreated and treated groups were
310 delivered in a randomized manner by the care taker to the lab technician each day.
311 All ETs were completed within a week.

312

313 *Data acquisition in surrogate mothers:*

314 Water intake was determined at different time points as shown in Fig. 1.

315 Water bottles were weighed at 08:00 and 13:00 of day 1, at 08:00 of day 2 and at
316 08:00 of day 3, i.e. 5, 24 and 48 hrs after starting the experiment. Surrogate mothers
317 were monitored for pregnancy after 9 and 12 days of gestation by checking the
318 appearance of the abdominal girth. From day 20 post coitum onwards, they were
319 checked twice daily for birth, and offspring (including still births) were counted. All
320 newborn offspring were weighed using an analytical balance within 12 hrs after birth,
321 and counted daily until weaning at 21 days of age.

322

323 *2.4. Statistical analyses:*

324 Statistical analyses were performed using SPSS 22 software (IBM, Armonk, NY,
325 USA). All data were tested for normal distribution and homogeneity of variance, and
326 are presented as mean +/- standard deviation.

327 Baseline and experimental water intake in ml/h of naïve mice was compared with a
328 paired t-test. Water intake of surrogate mothers (in ml/h), as well as litter size and
329 offspring weight after ET, were compared between treatment groups by independent
330 t-tests. Mean serum concentrations of paracetamol were compared between
331 different administration routes with one-way ANOVA. Post hoc analysis with the
332 Games Howell test was carried out to identify significant differences between
333 groups. Significance for all statistical tests was established at $p \leq 0.05$.

334

335 **3. Results**

336 *3.1. Intake of water with and without paracetamol in naïve mice and surrogate* 337 *mothers*

338 Fluid intake in ml per hour in naïve mice with and without paracetamol in the drinking
339 water is presented in Fig. 2.

340 During baseline measurements, naïve mice drank approximately 4.5 ± 1.46 ml per
341 day in total. Water intake increased significantly when paracetamol was added to
342 the drinking water. Intake of paracetamol-treated water versus untreated water was
343 2.0 ± 0.74 ml vs. 1.2 ± 0.53 ml ($p = 0.017$) after 6 hrs, 3.4 ± 0.82 ml vs. 1.8 ± 0.13
344 ml ($p = 0.001$) after 11 hrs, and 5.8 ± 1.40 ml vs. 3.6 ± 0.71 ml ($p = 0.001$) after 24
345 hrs of administration.

346 Fluid intake in ml per hour in surrogate mothers with and without paracetamol in the
347 drinking water is shown in Fig. 3.

348 Total intake of untreated water within the 5 hrs prior to ET varied between individual
349 animals, ranging from 0.7 to 2.1 ml. The intake of paracetamol-containing water
350 during this period ranged from 1.3 to 2.9 ml. The difference between both groups
351 was non-significant preemptive to ET ($p = 0.243$). Total water intake after surgery
352 was notably higher with paracetamol treatment, with a significant increase during
353 the first (6–24 hrs, $p = 0.023$) and the second (24–48 hrs, $p = 0.008$) day after ET.

354

355 3.2. *Estimated paracetamol intake calculated from water intake*

356 From the amount of paracetamol-containing water consumed by naïve mice, the
357 following consequential doses can be calculated: group PW1 mice consumed 104–
358 357 mg/kg BW (mean 208 ± 90) paracetamol within 6 hrs. In the first 6 hrs, five mice
359 consumed less than the target dose of 200 mg/kg BW, namely 104–177 mg/kg BW,
360 whereas three mice consumed more than the target dose. In group PW 2, mice
361 consumed 273–506 mg/kg BW (mean 351 ± 85) paracetamol within 11 hrs, i.e. all
362 group PW2 mice consumed more than the target dose. In group PW3, mice
363 consumed 331–636 mg/kg BW (mean 517 ± 106) paracetamol within 24 hrs, and
364 calculated doses exceeded target dose of 200 mg/kg BW in all mice. From the

365 amount of paracetamol-containing water consumed by surrogate mothers, the
366 following consequential doses can be calculated: before ET started, doses of
367 between 154 and 343 mg/kg BW were consumed within 5 hrs (mean 236 ± 68). Two
368 out of 7 animals consumed doses less than 200 mg/kg BW prior to ET, namely
369 154 mg/kg BW and 170 mg/kg BW. Following ET, in the remaining 18 hrs of day 1,
370 the amount of paracetamol additionally consumed ranged from 379 to 766 mg/kg
371 BW (mean 590 ± 123). On day 2 after ET, doses from 680 to 1077 mg/kg BW (mean
372 820 ± 132) were consumed within 24 hrs.

373

374 3.3. Serum paracetamol concentrations

375 After 6 hrs, the mean serum concentration of naïve mice receiving paracetamol with
376 drinking water (PW1) was $11.1 \pm 3.0 \mu\text{mol/L}$ ($1681.6 \pm 460.1 \text{ ng/ml}$). Serum
377 concentrations of naïve mice receiving paracetamol with drinking water were
378 similarly increased after 11 h and 24 h (PW2: $18.3 \pm 5.7 \mu\text{mol/L}$, 2777.6 ± 870.0
379 ng/ml ; PW3: $18.5 \pm 10.7 \mu\text{mol/L}$, $2796.5 \pm 1620.0 \text{ ng/ml}$). In control groups, the
380 serum concentration was high 2 hrs after bolus application in the injection group (I),
381 with $29.1 \pm 8.14 \mu\text{mol/L}$ ($4402.6 \pm 1152.3 \text{ ng/ml}$), as well as in the gavage group (G)
382 with $37.5 \pm 14.60 \mu\text{mol/L}$ ($5668.5 \pm 2208.3 \text{ ng/ml}$).

383 Mean serum concentrations differed significantly [$F(4,35) = 9.85$, $p \leq 0.0001$]. Post
384 hoc tests revealed significant differences between the i.p. injection group (I) and
385 PW1 ($p = 0.002$), as well as between the gavage group (G) and PW1 ($p = 0.008$)
386 and PW2 ($p = 0.044$).

387 Individual serum concentrations of mice of different treatment groups are shown in
388 Fig. 4.

389

390 *3.4. Outcome from ET: comparison of reproductive parameters*

391 The results are summarized in Table 1.

392 ET was successful in all surrogate mothers of the untreated group, i.e. without
393 paracetamol in the drinking water. All mice became pregnant and delivered litters of
394 2–6 pups (chronological order: 6, 5, 3, 2, 3, 6, 6, 4).

395 In the paracetamol-treated group, one recipient was detected not to be pregnant at
396 day 9 and 12 of gestation. We assume that pseudo-pregnancy in this female,
397 although with a vaginal plug, had not been appropriately induced. However, this
398 negative result was included for calculations and analysis. The remaining 6
399 recipients of the paracetamol-treated group delivered litters of 3–8 pups
400 (chronological order: 8, 3, 6, 6, 6, 3). The treated surrogate mothers delivered on
401 average slightly more pups per litter; however, differences in the final success of ET
402 were not significant ($p = 0.864$).

403 The body weight of newborns was not significantly different between the two groups
404 ($p = 0.330$). No dead offspring (or parts of pups) were found in cages around the
405 time of birth, and all pups were reared and developed well, i.e. no losses or
406 aberrations of growth or health were noticed at weaning.

407

408 **4. Discussion**

409 This study found no evidence of adverse effects on gestation or embryonic
410 development after administration of 3.5 mg paracetamol per ml drinking water for
411 2 days post-surgery. Interestingly, the water intake of surrogate mothers and naïve
412 mice increased when paracetamol was added to the drinking water in the form of a
413 children's syrup. Measurements of serum concentration of paracetamol in naïve
414 mice confirmed substantial drug uptake after 6 hrs preemptive application (i.e. the

415 approximate time point of the ET), and drug levels increased further after 11 and 24
416 hrs (i.e. correlating with the post-operative phase after ET). In summary, mice
417 obviously consumed considerable amounts of paracetamol voluntarily with their
418 drinking water before and after surgery, and the outcome of ET was unaffected by
419 the treatment.

420 Paracetamol, also known as acetaminophen, is one of the most widely used
421 analgesic and antipyretic drugs in human medicine. It is considered safe in
422 therapeutic dosages to treat fever and pain, and is one of the few pain medications
423 recommended during pregnancy (de Fays et al., 2015; Thiele et al., 2013). For pain
424 treatment in adult human patients, dosages of 325–650 mg paracetamol
425 administered per orally or parenteral every 3–4 hrs (max. 4000 mg within 24h) are
426 generally considered to be effective and safe. In laboratory mice, doses of 110–305
427 mg/kg BW (Fish et al., 2008; Flecknell, 1984; Hawk et al., 2005) have been used for
428 decades. The most common dose recommended by textbooks for pain treatment in
429 mice is 200 mg paracetamol/kg BW (Flecknell, 2009; Miller and Richardson, 2011).
430 According to these recommendations, for our study, the amount of paracetamol in
431 the drinking water was calculated to be 3.5 mg/ml, with the intention to provide the
432 mice with approximately 200 mg per kg BW. This target dose was reached within
433 the first 5–6 hrs in some of the naïve mice and surrogate mothers after providing
434 paracetamol-enriched drinking water. However, several mice stayed beneath the
435 target dose (104–177 mg/kg BW) of 200 mg/kg BW after 5–6 hrs, i.e. just before the
436 intended ET. Low water intake during the pre-operative phase could have been due
437 to the still unfamiliar taste of the water, and to generally lower water intake at the
438 beginning of the light period. Water consumption during the day time tends to be
439 less and more sporadic than during night time due to circadian rhythmicity (Sauer
440 et al., 2016).

441 After 11 and 24 hrs, all naïve mice voluntarily consumed more than the target dose.
442 The consumption of medicated water also increased in surrogate mothers during
443 the post-surgery treatment phase of 24 and 48 hrs, resulting in an ingested dose
444 significantly higher than the target dose of 200 mg/kg BW (Figs. 2 and 3). This is
445 likely to be attributed to the fact that paracetamol was added to the drinking water
446 as a children's syrup, which, due to its sweet taste, could have stimulated animals
447 to drink more than usual, even after surgery.

448 It is well known that paracetamol can cause severe liver damage when overdosed.
449 Damage to the liver is not induced by the drug itself but by the build-up of a toxic
450 metabolite due to oversaturated glucuronidation in the liver (Mattia and Coluzzi,
451 2009). Due to its hepatotoxic characteristics, paracetamol is used widely in
452 experimental models of acute liver injury in mice. According to safety data sheets
453 for paracetamol, the oral lethal dose (LD) 50 in mice is 338 mg/kg BW (see for
454 example www.caymanchem.com/msdss/10024m.pdf). However, it has been
455 reported that experimentally induced liver injury is also sex- as well as strain-
456 dependent (Mohar et al., 2014; Mossanen and Tacke, 2015). Male mice seem to be
457 more susceptible than female mice (Taguchi et al., 2015), and C57BL/6 mice are
458 more responsive than BALB/c (Mossanen and Tacke, 2015). Mossanen and Tacke
459 recommend a dose of 300 mg/kg BW paracetamol with i.p. injection after a fasting
460 period of 12 hrs to reliably induce acute liver injury in mice. Taguchi et al.
461 administered doses of 300 mg/kg BW or 600 mg/kg BW paracetamol, with i.p.
462 injection after 12 hrs fasting to induce liver injury in 4- to 12-week-old mice (Taguchi
463 et al., 2015). Additionally, a recent study showed that pregnant mice were more
464 sensitive to paracetamol-induced hepatotoxicity (Karimi et al., 2015). In this latter
465 study, a dose of 250 mg/kg BW paracetamol administered as a single bolus injection
466 after 16 hrs of fasting at gestation day 12.5 induced hepatocellular injury and

467 inflammation, while a dose of 450 mg/kg BW induced lethal effects in pregnant but
468 not in non-pregnant mice. Although paracetamol administration did not affect the
469 fetal loss rate, decreased body weights were found in offspring in the prenatal and
470 neonatal stage (Karimi et al., 2015).

471 As most of the mice in our study voluntarily consumed, at least during the second
472 part of the experiment, higher doses than the target dose of 200 mg/kg BW, and in
473 some cases even more than the highest recommended dose of 305 mg/kg BW,
474 concern regarding potential liver damage or decreased body weight in offspring due
475 to accidental overdosing arises. However, studies by Hayes et al. and Christy et al.
476 revealed no deaths or apparent signs of liver damage or failure even after mice
477 ingested approx. 320–640 mg/kg BW of paracetamol voluntary via drinking water
478 (Christy et al., 2014; Hayes et al., 2000).

479 To elucidate further the potential for over-dosage and subsequent toxic effects from
480 paracetamol consumption with drinking water in our study, the concentration of
481 paracetamol in blood serum was determined in naïve mice. In both our control
482 groups, after i.p. injection or gavage of 200 mg/kg BW as a bolus, serum
483 concentrations of paracetamol reached 4402.6 ± 1152.3 ng/ml and 5668.5 ± 2208.3
484 ng/ml, respectively, at 2 hrs after treatment. In contrast, serum concentrations were
485 significantly lower in all drinking water groups compared to our controls. Here, the
486 maximum level of 2796.5 ± 1620.0 ng/ml was noted after 24 hrs (PW3).

487 In human patients, if plasma concentrations 4 hrs after drug intake are lower than
488 120023 ng/ml (794 μ mol/L), toxic liver effects are unlikely to result. If plasma
489 concentrations are higher than 120023 ng/ml (794 μ mol/L), liver insufficiency could
490 occur, and if plasma concentrations are higher than 300057 ng/ml (1985 μ mol/L),
491 liver necrosis is likely (DRI® Paracetamol-Serum-Tox-Assay). As data for toxic
492 plasma concentrations in mice are still lacking, we have to rely on data from human

493 studies: In our study, serum concentrations of paracetamol after bolus application
494 as well as after voluntary intake in drinking water, were always far below critical
495 levels from human tox-assays. Moreover, no cases of death occurred, and no
496 obvious aberrations in appearance and behaviour of animals were noticed at regular
497 routine checking. We therefore assume that toxic effects were unlikely at the doses
498 used.

499 Additionally, in our study, doses of up to about 600–1000 mg/kg BW paracetamol
500 per day in the drinking water of mice on days 1 and 2 of gestation did not lead to
501 any significant impairment of our ET success rate. The number of pups born was
502 related to the number of transferred two-cell stage embryos, and was not
503 significantly different between the paracetamol-treated and untreated surrogate
504 mothers. Although one of the surrogate mothers in the paracetamol treated group
505 failed to get pregnant while all untreated animals gave birth, the litters of treated
506 surrogate mothers were on average larger, thus compensating for the lower rate of
507 pregnancy. In addition, the body weight of newborn pups was comparable after
508 paracetamol treatment of recipients at 2 days of gestation. Altogether, our results
509 provided no evidence for any adverse effects of paracetamol treatment on the
510 overall outcome of ET.

511 The observed lack of detrimental effects on animal health and ET outcome may be
512 the result of constant but low intake of the drug via drinking water. Most mice in the
513 present study consumed high levels of paracetamol; however, the animals ingested
514 the medication distributed over a time span of up to 2 days rather than as a high
515 dose bolus after fasting, as carried out in studies to induce liver damage (Corcoran
516 et al., 1988; Karimi et al., 2015; Mossanen and Tacke, 2015; Taguchi et al., 2015)
517 or for traditional LD50 determination. Paracetamol reaches peak concentrations at
518 30–60 minutes after administration, and its half-life in blood plasma is about 2 hrs

519 (Flower et al., 1985; Mickley et al., 2006), thus reducing concerns regarding toxicity
520 in our study.

521 In the present study, the efficacy of paracetamol in regards to post-operative pain
522 relief was not investigated. The focus was rather on whether mice would voluntarily
523 ingest paracetamol-enriched water in amounts sufficient to achieve commonly
524 recommended doses, and whether the drug had any influence on the success rate
525 of ET and offspring survival. Both strains of mice (CrI:CD and Zbz:FM) consumed
526 similar doses of acetaminophen via the drinking water. However, as food and water
527 intake can differ between strains (Bachmanov et al., 2002), the dosage of
528 acetaminophen may also need to be adjusted due to strain variation (Dickinson et
529 al., 2009). Consequently, no evaluation of pain relief can be drawn from the present
530 study, even though plasma levels of paracetamol were comparable to doses
531 effective in analgesiometric tests (Qiu et al., 2007). Future studies are needed to
532 provide evidence for the degree of pain relief after ET with paracetamol, and to
533 elucidate other possible side-effects of the drug when used for this purpose.

534 For transferring this protocol to other laboratories, specific conditions of each
535 country might be considered. It could be necessary to check for availability of
536 acetaminophen and clarify whether a formulation or commercially available drug is
537 permitted by regulative authorities for the use in experimental animals.

538 With regard to surrogate mothers, specifics of strain and age might be considered,
539 although for ET females in a similar age range and mostly outbred strains are used.
540 Thus, differences regarding dose-response and toxic effects might be negligible.
541 This is underpinned by our observation that both outbred strains (CrI:CD and
542 Zbz:FM) consumed similar amounts of water, i.e. doses of acetaminophen.

543 Furthermore, the uptake of paracetamol with the drinking water might be decreased
544 after anaesthesia and surgery, but data obtained in this study and from other

545 publications (Cesarovic et al., 2010; Sauer et al., 2016) show, that no relevant
546 alteration of drinking behaviour occurred after inhalation anaesthesia with or without
547 surgery. However, in case of doubts regarding uptake of the drug in the immediate
548 post-anaesthetic phase, one may administer the analgesic then as a single bolus-
549 injection to compensate for a suspected delay in drinking after the intervention.

550

551 **5. Conclusions**

552 In summary, the animals in our study ingested voluntarily substantial amounts of
553 paracetamol with drinking water that allow the assumption of constant post-
554 operative pain treatment. An extension of the preemptive application phase of the
555 medication in the drinking water or a single i.p. injection of paracetamol might be
556 necessary to assure target plasma concentration immediately before, and during
557 the first hours after ET. High doses of paracetamol were reached already several
558 hours after surgery, supported by the increased consumption of medicated water.
559 The animals received their medication without stress through handling, restraint, or
560 manipulation (e.g. frequent injections), all of which could influence their well-being
561 (Jirkof et al., 2015) and possibly adversely affect pregnancy and the outcome of ET.
562 Although substantial doses of paracetamol were consumed within 2 days after
563 surgery, no side-effects on the overall outcome of ET were detected. Therefore,
564 administering paracetamol in drinking water could be a feasible method for providing
565 pain relief in mice undergoing ET.

566

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568 commercial, or not-for-profit sectors.

569

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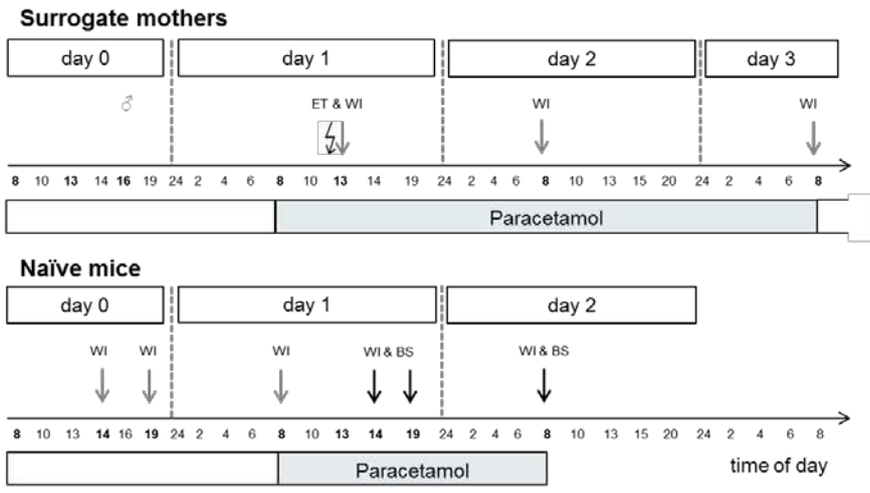
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598 **Legends**

599 **Fig. 1: Experimental schedule for surrogate mothers and naïve mice.**

600 Measurements of intake of either untreated (n = 8) or paracetamol-containing water
 601 (n = 7) in surrogate mothers took place at 13:00 and 08:00 (i.e. after 5, 24, 48 hrs).
 602 ET was performed at 5 hrs after the start of the experiment (13:00 – 14:00).
 603 In naïve mice (n = 8 / group), baseline measurements of untreated water intake took
 604 place at 14:00, 19:00 and 08:00 (i.e. after 6, 11, 24 hrs) on the first day. On the
 605 following day, measurements of paracetamol-treated water intake as well as blood
 606 sampling, took place at 14:00, 19:00 and 08:00 (i.e. after 6, 11, 24 hrs).
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- Surrogate mothers:
- ♂ mating with vasectomized males
 - ☐ medicated water bottle / paracetamol treatment
 - ⚡ embryo transfer surgery (ET)
 - WI ↓ measurement of water intake at 13:00 and 08:00 (i.e. after 5, 24, 48 hrs)
- Naïve mice:
- ☐ medicated water bottle / paracetamol treatment
 - WI ↓ measurement of water intake at 14:00, 19:00 and 08:00 (i.e. after 6, 11, 24 hrs)
 - BS ↓ blood sampling at 14:00, 19:00 and 08:00 (i.e. after 6, 11, 24 hrs)

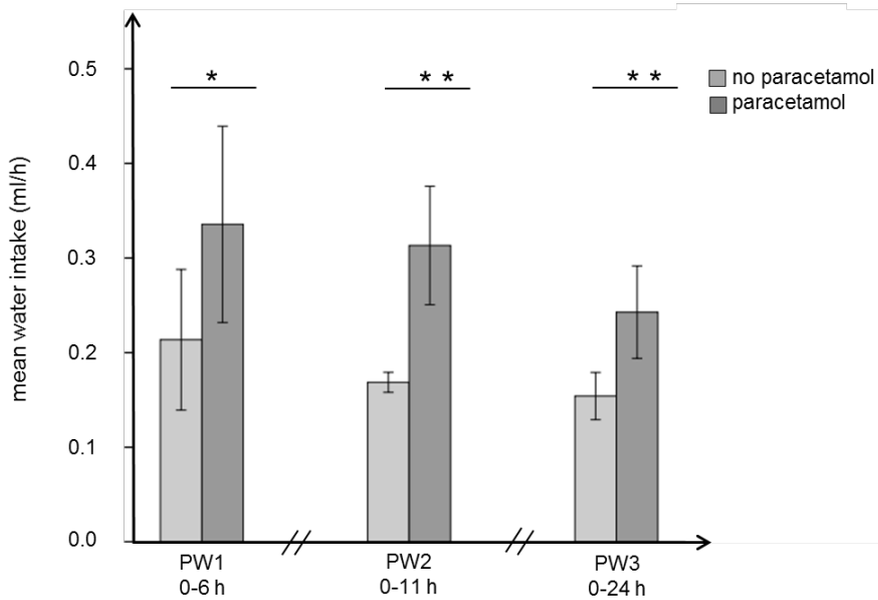
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611 **Fig. 2: Comparison of mean water intake per hour with and without**
612 **paracetamol in naïve mice.**

613 Paracetamol was provided to naïve mice in their drinking water at a concentration
614 of 3.5 mg paracetamol per ml water. Baseline measurements for intake of untreated
615 water were taken the day before. Measurements of water intake was conducted
616 after 6 hrs in PW1, after 11 hrs in PW2, and at 24 h in PW3 (n = 8 / group). Mean
617 values (\pm SD) of water intake in naïve mice with and without paracetamol in drinking
618 water is traced as ml/h. Bars indicate SD. Significant differences between baseline
619 and experiment were found in all three groups (PW1: $p = 0.017$; PW2: $p =$
620 0.001 ;PW3: $p = 0.001$). * $p \leq 0.05$ and ** $p \leq 0.01$.

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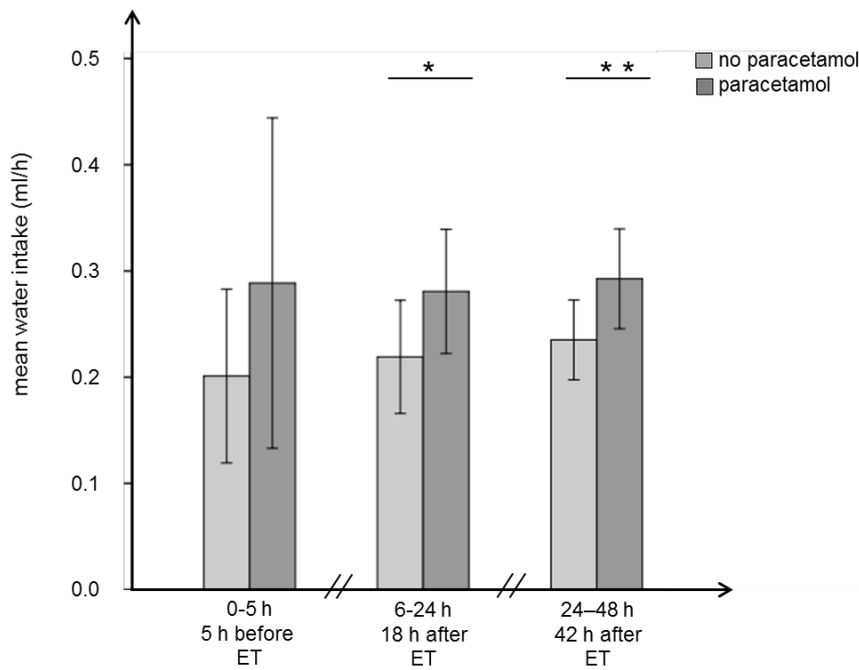
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626 **Fig. 3: Comparison of mean water intake per hour with and without**
627 **paracetamol in surrogate mothers.**

628 Mean water intake in untreated (n = 8) and paracetamol-treated (n = 7) surrogate
629 mothers in the 5 hrs before ET, and during 2 days (\leq 42 hrs) after ET. Water intake
630 was calculated as ml/h. Bars indicate SD. A significant difference was found
631 between treated and untreated groups on the first (18 hrs post ET, $p = 0.023$) and
632 second (42 hrs post ET, $p = 0.008$) day after ET. * $p \leq 0.05$ and ** $p \leq 0.01$.

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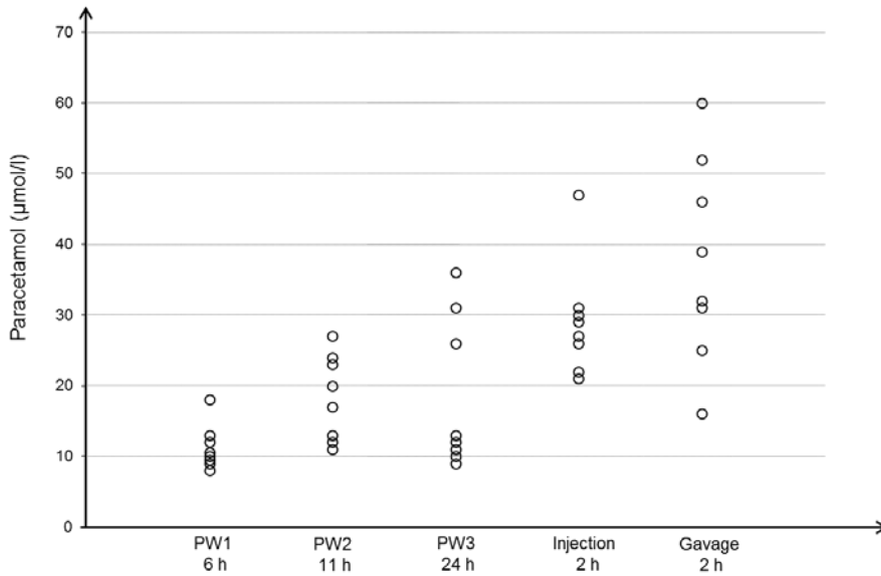
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640 **Fig. 4: Individual serum concentrations of paracetamol in naïve mice.**

641 In PW groups, paracetamol was provided to naïve mice in their drinking water at a
642 concentration of 3.5 mg paracetamol per ml water. Blood serum was taken after 6
643 hrs in PW1, after 11 hrs in PW2, and at 24 hrs in PW3. In control groups,
644 paracetamol was administered as bolus at a dose of 200 mg/kg BW by
645 intraperitoneal injection (I) or gavage (G). Blood was sampled at 2 hrs after bolus
646 application.

647 Individual serum concentrations for all groups (n = 8 / group) are depicted as one
648 dot for each mouse.

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656 **Table 1: Outcome of ET.**

657 One surrogate mother of the paracetamol-treated group was not visibly pregnant
 658 and did not give birth, but was included in the calculation of data. Statistical
 659 comparison of litter size and offspring weight showed no significant difference
 660 whether surrogate mothers received paracetamol or not with their drinking water for
 661 48 hrs (success rate: $p = 0.864$, body weight in newborn offspring: $p = 0.330$).

662

	without treatment		with paracetamol	
number of foster mothers used for ET	8		7	
total number of two cell embryos transferred	96		84	
number of pregnant females at day 9 and 12 of gestation	8		6	
number of litters	8		6	
total number of offsprings	35		32	
mean litter size	4.38 (± 1.60)		4.57 (± 2.70)	
relation between live offsprings and transferred two cell embryos (success rate)	35/96 (36%)		32/84 (38%)	
mean offspring body weight at birth [g], (\pm SD)	1.90 (± 0.19)	n=35	1.86 (± 0.14)	n=32

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