

1 Refinement of intraperitoneal injection of sodium pentobarbital for euthanasia in

2 laboratory rats (*Rattus norvegicus*)

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16 Running head: Intraperitoneal euthanasia in rats

17 Key words: CCAC, pentobarbital, killing, refinement, welfare

18 Abbreviations:

19 **LL**: low-dose low-volume

20 **LH**: low-dose high-volume

21 **HH**: high-dose high-volume

22 **LORR**: loss of righting reflex

23 **CHB**: cessation of heartbeat

24 **GIT**: gastrointestinal tract

25 **CV**: coefficient of variation

26 **Abstract**

27 **Background:** The Canadian Council on Animal Care and American Veterinary Medical
28 Association classify intraperitoneal (IP) pentobarbital as an acceptable euthanasia
29 method in rats. However, federal guidelines do not exist for a recommended dose or
30 volume and IP euthanasia has been described as unreliable, with misinjections leading
31 to variable success in ensuring a timely death. The aims of this study were to assess
32 and improve efficacy and consistency of IP euthanasia.

33 **Methods:** In a randomized, blinded study, 51 adult female Sprague-Dawley rats
34 (170-495 g) received one of four treatments: low-dose low-volume (LL) IP pentobarbital
35 (n = 13, 200 mg/kg pentobarbital), low-dose high-volume (LH) IP pentobarbital (n = 14,
36 200 mg/kg diluted 1:3 with phosphate buffered saline), high-dose high-volume (HH, n =
37 14, 800 mg/kg pentobarbital), or saline. Times to loss of righting reflex (LORR) and
38 cessation of heartbeat (CHB) were recorded. To identify misinjections, necropsy
39 examinations were performed on all rats. Video recordings of LL and HH groups were
40 analyzed for pain-associated behaviors. Between-group comparisons were performed
41 with 1-way ANOVA and Games-Howell post hoc tests. Variability for CHB was assessed
42 by coefficient of variation (CV) calculation.

43 **Results:** The fastest euthanasia method (CHB) was HH (283.7 ± 38 s), compared with
44 LL (485.8 ± 140.7 s, $p = 0.002$) and LH (347.7 ± 72.0 s, $p = 0.039$). Values for CV were:
45 HH, 13.4%; LH, 20.7%; LL, 29.0%. LORR time was longest in LL (139.5 ± 29.6 s),
46 compared with HH (111.6 ± 19.7 s, $p = 0.046$) and LH (104.2 ± 19.3 s, $p = 0.01$).

47 Misinjections occurred in 15.7% (8/51) of euthanasia attempts. Pain-associated

48 behavior incidence ranged from 36% (LL) to 46% (HH).

49 **Conclusion:** These data illustrate refinement of this euthanasia technique. Both dose

50 and volume contribute to speed of death with IP pentobarbital and an increase in

51 volume alone does not significantly reduce variability. The proportion of misinjections

52 was similar to that of previous studies.

53

54

55 **Introduction**

56 Over 2 million rats are used in biomedical research in Canada and the European Union
57 annually [1, 2]. The overwhelming majority of laboratory studies employing rodents end
58 with killing the animals upon completion of the study or if a humane endpoint has been
59 reached. While this is a reality of research, efforts to refine killing methods, to achieve
60 “euthanasia”, for rats and other laboratory animals are ongoing, as reflected in recent
61 updates to the Canadian Council on Animal Care (CCAC) and American Veterinary
62 Medical Association (AVMA) euthanasia guidelines [3, 4]. Goals for successful
63 euthanasia include techniques requiring minimal restraint, simplicity of administration,
64 and a swift, painless death [5].

65 A commonly employed technique for euthanasia of laboratory rats is an overdose of
66 carbon dioxide. However, current behavioral and physiologic evidence suggests that
67 this method is aversive and may be painful [6-15]. As a result, the AVMA and CCAC
68 have reclassified killing with carbon dioxide as “conditionally acceptable” [4] and
69 “acceptable with conditions” [3].

70 In contrast, an acceptable method and preferred alternative to carbon dioxide is
71 overdose with a barbiturate such as sodium pentobarbital (PB). An intraperitoneal (IP)
72 route of injection is acceptable when intravenous injection cannot be performed or is
73 impractical [3, 4]. Current guidelines do not indicate a specific dose of sodium
74 pentobarbital for euthanasia, although 200 mg/kg or 3 times the anesthetic dose have
75 been suggested [5]. There are several potential drawbacks associated with IP PB
76 injection, including misinjection, variability in effect and pain [7, 16-22].

77 An important factor contributing to variability of drug effect (speed of onset and success)
78 is misinjection, with deposition of injectate into intra-abdominal fat, abdominal viscera or
79 the subcutaneous space. In the case of IP pentobarbital for euthanasia this results in a
80 delayed time to death or even failure to cause loss of consciousness. Attempts to
81 reduce variability with a two-person injection technique (one to restrain, one to inject)
82 have had variable success, with reported proportions of misinjections ranging from 6 to
83 20% [18-20].

84 Pain, inferred from behavioral observations, necropsy findings and biomarkers, has also
85 been cited as a potential impediment to achieving the principle of euthanasia.

86 Specifically, exhibition of writhing (defined as the contraction of the abdomen and
87 extension of the hind legs), grossly visible inflammation of abdominal viscera at
88 necropsy and a measurable increase in spinal cord cFos have been reported following
89 IP injection of pentobarbital [16, 17, 21, 22].

90 The primary aim of this study was to assess the impact of varying the dose and volume
91 of sodium pentobarbital injected into the intraperitoneal cavity on time to death and
92 consistency of the killing process. Secondary aims were identification of misinjections
93 by necropsy and the quantification of writhing behavior in response to IP PB. We
94 hypothesized that speed and consistency of IP euthanasia would be improved by using
95 a higher dose and higher volume.

96 **Methods**

97 *Ethics Statement*

98 The animal care and use protocol was approved by the Health Sciences Animal Care
99 Committee of the University of Calgary (AC11-0044), in accordance with the guidelines
100 of the CCAC.

101 *Study Design*

102 51 adult female Sprague-Dawley rats (170-495 g), sourced as surplus breeding stock,
103 were included in the study. A sample size of approximately 13 animals, to achieve 80%
104 power with an alpha of 0.05 (with an anticipated 20% misinjection rate) with an effect
105 size of 1.5, was determined from pilot data. All animals remained in paired housing until
106 the time of trial and were not handled prior to the study. Housing consisted of standard
107 micro-filter cages (47 x 25 x 21 cm) with wood shavings and shredded paper bedding
108 and a plastic tube for enrichment. A 12-12 hour lights on-off cycle (lights on at 0700)
109 was maintained in an environmentally controlled room (23°C, 22% humidity). All
110 experiments were performed during the light period (0730-1800).

111 Animals were randomly assigned to one of four treatment groups for IP injection. A low-
112 dose low-volume (LL, n = 13) group received 200 mg/kg sodium pentobarbital
113 (Euthanyl, 240 mg/ml, Bimeda-MTC Animal Health Inc., Cambridge, ON, Canada). A
114 low-dose high-volume group (LH, n = 14) received 200 mg/kg sodium pentobarbital
115 diluted 1:3 with phosphate-buffered saline (PBS). A high-dose high-volume (HH, n = 14)
116 group received 800 mg/kg sodium pentobarbital. A control group (n = 10) received 1 ml
117 of PBS. Each treatment was placed in a 1 ml (LL and control groups) or 3 ml (LH and
118 HH groups) syringe as dictated by the volume of injectate. A new 25 G 5/8" hypodermic
119 needle was attached to each syringe for injection. Blue food coloring (0.01 mL, Club

120 House, Burlington, Ontario) was added to each treatment to facilitate visualization of
121 injectate during necropsy examination.

122 At the beginning of each trial, a single rat was removed from the housing unit and
123 placed in a Plexiglas chamber (L x W x H: 27.5 x 14.5 x 20.5 cm). Two video cameras
124 (Panasonic HC-V720P/PC, Panasonic Canada Inc., Mississauga, ON, Canada) were
125 placed along the long and short axes of the chamber. Prior to each injection, baseline
126 video of the rat was recorded for 10 minutes. Treatments were prepared in a separate
127 room during baseline video recording. Individuals performing the IP injections and
128 behavioral analyses were blinded to treatment.

129 Following baseline video, each rat was removed from the box and restrained for a two-
130 person injection technique. Rats were held in dorsal recumbency at an approximately
131 30-degree angle (head lowermost). The holder (DP) supported each rat and restrained
132 the left pelvic limb. The individual administering each injection (KZ) restrained each rat's
133 right pelvic limb, injecting with the right (dominant) hand (Fig. 1). Each injection was
134 performed in the right caudal quadrant of the abdomen at the level of the coxofemoral
135 joint and approximately 5 mm to the right of midline. The needle was directed cranially
136 at a 45-degree angle to the body wall.

137 Immediately following completion of injection, each rat was returned to the observation
138 chamber. A single blinded observer (KZ) monitored for signs of ataxia (stumbling, falling,
139 crossing feet) following injection. If signs of ataxia were noted, an attempt was made to
140 place the rat in dorsal recumbency to evaluate for a loss of righting reflex (LORR), a
141 surrogate for loss of consciousness [7, 23]. LORR was confirmed if the rat remained in

142 dorsal recumbency for ten seconds. Failure of LORR was established if the rat resisted
143 initial placement on its back or was able to right itself within ten seconds. In cases of
144 initial LORR failure, the test was repeated every 30 seconds until LORR occurred.
145 Following LORR, the animal was monitored for onset of apnea, defined as the animal's
146 chest ceasing to rise and fall. If and when apnea occurred, the rat was placed in left
147 lateral recumbency. The left thoracic wall was then auscultated continuously with a
148 stethoscope to identify cessation of heartbeat (CHB). Following CHB confirmation, video
149 recording was stopped. The observation chamber was cleaned between trials.
150 When CHB did not occur within 20 minutes of IP injection, animals were euthanized with
151 an overdose of carbon dioxide gas using a gradual fill (30% chamber volume per
152 minute) technique. These cases were considered unsuccessful euthanasias.

153 *Necropsy Examination*

154 Following CHB, each animal was carefully removed from the chamber and positioned in
155 dorsal recumbency for necropsy examination. The skin was incised along the midline
156 and the injection site was identified in the abdominal wall musculature. The abdominal
157 wall was incised and the intestines were reflected out of the abdominal cavity.
158 Distribution of blue injectate and any misinjection into hollow viscera were noted. The
159 liver was reflected cranially and any presence of dye within the biliary vessels caused
160 by uptake of injectate from the peritoneal cavity and subsequent biliary excretion was
161 noted. The GIT from the cardia to the descending colon was removed and any intestinal
162 segments with dye-stained serosa were opened to confirm or rule out intraluminal
163 misinjection. Misinjection was defined as the presence of blue injectate within hollow

164 viscera or subcutaneous tissues, or staining the fur. For each rat, the serosal surfaces
165 of the abdominal wall injection site, the caudate liver lobe, and transverse sections of at
166 least three intestinal sections were examined histologically after formalin fixation for
167 evidence of acute inflammation or swelling of mesothelial cells. Evaluation was
168 performed by a single board-certified veterinary pathologist (CK), who was blinded to
169 treatment group assignments.

170 *Off-Line Video Analysis*

171 Videos of the HH and LL trials were analyzed for the incidence of writhing behavior by a
172 single individual blinded to treatment (JR). Baseline recordings were analyzed in their
173 entirety while post-injection videos were analyzed until the rat became ataxic. Writhing
174 was defined as a contraction of the lateral abdominal walls to the extent where the
175 abdomen became concave with concurrent extension of the pelvic limbs [17, 22].

176 *Statistical methods*

177 All statistical analyses were performed using commercial software (GraphPad Prism
178 v.6.03, GraphPad Software, Inc. La Jolla, California, USA and IBM SPSS Statistics 21,
179 IBM, Armonk, NY, USA). Data were considered approximately normal if skewness and
180 kurtosis were less than ± 1.5 and 3, respectively. Between-group comparisons were
181 performed with a one-way ANOVA with a Games-Howell multiple comparisons test.
182 Consistency of the euthanasia process was assessed with a coefficient of variation (CV)
183 calculation. A p-value of < 0.05 was considered significant. Data are presented as mean
184 \pm SD.

185 **Results**

186 Of 51 trials, 43 (84.3%) were successful IP injections and 8 (15.7%) were misinjections.
187 Successful IP injection resulted in death in all PB groups: 34 (79.1%) were given IP PB
188 and 9 were control animals. Successful deaths were distributed as follows: LL (n = 11),
189 LH (n = 12), and HH (n = 11).

190 The fastest killing method from injection to CHB was the HH group ($283.7 \pm 38.0s$),
191 which was significantly faster than both the LL ($485.8 \pm 140.7s$, $p = 0.002$) and LH
192 ($347.7 \pm 72.0s$, $p = 0.039$) groups (Fig. 2). Euthanasia in the LH group was also
193 significantly faster than the LL group ($p = 0.027$).

194 The HH group was not only the fastest, but also the most consistent euthanasia
195 method. The CV for HH was 13.4%, compared with 29.0% for LL and 20.7% for LH
196 groups.

197 The period from injection to LORR was longest in LL ($139.5 \pm 29.6s$), compared with
198 both HH ($111.6 \pm 19.7s$, $p = 0.046$) and LH ($104.2 \pm 19.3s$, $p = 0.01$, Fig 3A). Time from
199 injection to LORR did not differ between LH and HH ($p = 0.64$). The LORR-apnea time
200 period showed the greatest variation between treatment groups and therefore had the
201 greatest influence on the speed of the overall time to death (Fig 3B). LORR-apnea was
202 significantly faster in the HH group ($56.8 \pm 25.1s$) than LL ($253.3 \pm 106.7s$, $p < 0.001$)
203 and LH ($146.6 \pm 66.1s$, $p = 0.002$). LORR-apnea in the LH group was also significantly
204 faster than in the LL group ($p = 0.03$). There was no significant difference from apnea-
205 CHB among treatment groups: HH ($116.2 \pm 19.7s$) versus LH ($93.0 \pm 29.0s$, $p = 0.09$),
206 HH versus LL ($92.9 \pm 24.2s$, $p = 0.06$), LH versus LL ($p = 1.0$).

207 Eight misinjections were identified at necropsy. One misinjection was in a control
208 animal. Seven misinjections were treatment group rats (HH; n = 3, LH; n = 2, LL; n = 2).
209 Of these, euthanasia was unsuccessful (exceeding 20 minutes) in 3 (42.8%) animals
210 (HH [n = 2], LL [n = 1]). In the four animals in which euthanasia was successful (HH [n =
211 1], LH [n = 2], LL [n = 1]), injection-CHB ranged from 318-1200s.
212 The anatomic distribution of the eight misinjections was as follows: four entered the
213 cecal lumen (Fig 4B), two entered the jejunal lumen (Fig 4C), one was entirely within the
214 subcutaneous tissues of the abdominal wall (Fig 4D), and one was predominantly over
215 the fur of the medial thigh, with a small amount in the subcutaneous space. Cecal
216 positions were variable: 14/51 (27.5%) in the right caudal quadrant, 5/51 (9.8%) located
217 in the midline and 32/51 (62.7%) in the left caudal quadrant.

218 *Writhing*

219 Writhing was not observed in either the LL or HH groups in baseline video recordings.
220 Following injection, writhing, assessed in animals with successful injections, was seen
221 in 45.5% (5/11) of HH and 36.4% (4/11) of LL rats.

222 **Discussion**

223 Historically, concerns regarding the variable success of IP euthanasia have revolved
224 around misinjection leading to variability and potential pain [7, 16-20, 22].
225 Our results show that: 1. IP injection with 800 mg/kg sodium pentobarbital (HH group)
226 resulted in the fastest and most consistent killing method; 2. variable cecal position
227 contributed to misinjections; and 3. the incidence of writhing behavior was less than half
228 of that previously reported.

229 Both dose and volume contribute to the speed of euthanasia, and dose in particular
230 appears to have the most dramatic effect on consistency of technique. The speed and
231 consistency of the killing process can be improved through an increase in dose
232 (accompanied by an increase in volume). Increasing injectate volume without increasing
233 dose (LH group) improved the speed of IP euthanasia. However, further improvements
234 in speed and consistency were achieved in the HH group.

235 From these results, several conclusions can be drawn. The mean + 2SD for the period
236 from completion of injection to LORR was 151.0 seconds when 800 mg/kg pentobarbital
237 (HH group) is administered IP. Therefore, it is highly likely that an animal that maintains
238 LORR beyond this time has experienced a misinjection. If using the period from
239 completion of injection to apnea as the indicator of successful injection, the time for
240 mean + 2 SD was 259.1 seconds. Should these times be exceeded, a second injection
241 of pentobarbital or alternative killing method should be performed.

242 Any increase in pentobarbital use is associated with an increased cost. For the
243 formulation used here, this equates to approximately US\$0.13 for the HH technique in a
244 250g rat. While cost is an important consideration, it should be weighed against the
245 labor cost of the slower (approximately 1.7 fold) LL group and potentially prolonged pain
246 experience during the period from injection to LORR.

247 Misinjection is a consistent limitation of IP PB. The rate of misinjection in this study was
248 consistent with the range reported in the literature (for injections given in to the caudal
249 right abdominal quadrant), from 6 to 20% [18-20]. A factor contributing to the
250 misinjection rate is variability in cecal position. IP injection is usually performed in the

251 right caudal abdominal quadrant and previous work has confirmed that the cecum is
252 most commonly located in the left caudal abdominal quadrant (61.9%, right 24.2%,
253 middle 13.8%, total n = 289 adult male and female rats) [19]. Our results are similar to
254 these findings despite using a different injection technique. In the study of Coria-Avila et
255 al. (2007), rats were restrained by a single person and suspended vertically by the
256 thorax with the head up. This suggests that body position during injection has a minimal
257 effect on the incidence of misinjection. Based on the misinjection rates in this and other
258 studies, as well as the positional variation of abdominal viscera noted on necropsy, any
259 suggested method of IP euthanasia is unlikely to prevent completely the possibility of
260 misinjections.

261 Given this inherent obstacle in refining the euthanasia process, we hope that the
262 recommendations described above will facilitate early identification of a misinjection,
263 guiding the decision to repeat the injection or select an alternative euthanasia method.
264 We observed a substantially lower incidence of writhing behavior than previously
265 reported [16, 17]. To facilitate comparison, we used the same definition of writhing as
266 that described by Wadham (1996) and Ambrose (1998, 1999) [16, 17, 22]. The reason
267 for this discrepancy is unclear and may result from several factors.

268 The proposed cause of writhing behavior is the pain resulting from the alkaline pH of the
269 PB solution. The pH of the solution studied here was 11.02 (measured independently by
270 a commercial compounding pharmacy) and that of Wadham (1996) and Ambrose (1998)
271 ranged between 10.9-12.6 [16, 22]. Current suggestions to alter the effect of pH focus
272 on changing solution pH through buffering or the addition of lidocaine to provide

273 analgesia [3, 4]. Wadham (1996) reported that buffering a solution of sodium
274 pentobarbital from an original pH of 12.6 to 9.4 resulted in precipitation [22].
275 Any study combining behavioral observation in the presence of drugs with sedative
276 properties is inherently confounded by a reduced ability to express behaviors as
277 sedation occurs. This is a limitation of the study design. The use of a vehicle control
278 would address this, but one was not readily available as there were restrictions in
279 obtaining formulation information from the manufacturer of PB. Furthermore, the dose
280 we used in the LL group (200 mg/kg), was higher than that of Ambrose (1998, 1999)
281 (150 mg/kg) and selected based on our institutional SOP [16, 17]. This may have
282 contributed to the lower incidence of writhing observed by shortening the time after
283 injection when writhing behavior could be expressed, before sedation occurred.
284 Finally, a lack of habituation to handling may have contributed to our findings. The rats
285 used in this study received little or no handling prior to the experiment. Therefore, the
286 stress associated with handling, injection and the observation chamber may have led to
287 a suppression of normal behaviors.
288 By coupling the effects of volume and dose with the incidence of misinjections we have
289 suggested practical guidelines to refine overdose with IP sodium pentobarbital as a
290 killing method in rats.

291 **Declarations**

292 Ethics approval and consent to participate

293 The animal care and use protocol was approved by the Health Sciences Animal Care
294 Committee of the University of Calgary (AC11-0044), in accordance with the guidelines
295 of the Canadian Council on Animal Care.

296 Consent for publication

297 Not applicable

298 Availability of data and materials

299 The datasets generated and analysed during the current study are available in the
300 Harvard Dataverse repository: Pang, Daniel, 2016, "rat IP PB dataset", [doi:10.7910/](https://doi.org/10.7910/DVN/PMGCHG)
301 [DVN/PMGCHG](https://doi.org/10.7910/DVN/PMGCHG), Harvard Dataverse, V1 [UNF:6:A8TqlyFIJXya5kXJ5eW0kQ==]

302 Competing interests

303 The authors declare that they have no competing interests

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312 Authors' contributions

313 KZ contributed to study design, collected and analyzed data and wrote the first draft of
314 the paper. CGK collected and analyzed data and revised the manuscript. JR analyzed

315 data and revised the manuscript. DSJP conceived the study, contributed to study
316 design, collected and analyzed data and revised the manuscript. All authors read and
317 approved the final manuscript.

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320

321 **Legends**

322 Fig 1: a cartoon showing the two-person injection technique used in the study, with one
323 person holding the rat in dorsal recumbency (head down) and the second person gently
324 restraining the right pelvic limb to facilitate intraperitoneal injection in to the right caudal
325 quadrant.

326 Fig 2: Time from delivery of the intraperitoneal injection to cessation of heart beat was
327 fastest in the high-dose high-volume group (HH). LL = low-dose low-volume group, LH =
328 low-dose high-volume group. * $p < 0.05$ ** $p = 0.002$

329 Fig 3A: Time from delivery of the intraperitoneal injection to loss of the righting reflex
330 was longest in the low-dose low-volume group (LL). LH = low-dose high-volume group,
331 HH = high-dose high-volume group. * $p < 0.05$ ** $p = 0.01$. Fig 3B: Time from loss of the
332 righting reflex to apnea was shortest in the high-dose high-volume group (HH). * $p < 0.05$
333 ** $p = 0.002$ *** $p < 0.001$

334 Fig 4: Abdominal cavities of four rats after confirmation of death; ventral view. Diffuse
335 blue dye staining of serosal surfaces following successful intraperitoneal injection (A).
336 Restricted dye distribution following inadvertent cecal (B), intestinal (C), and

337 subcutaneous (D) misinjection. The insets in panels B and C show dye-stained ingesta,
338 confirming inadvertent luminal misinjection.

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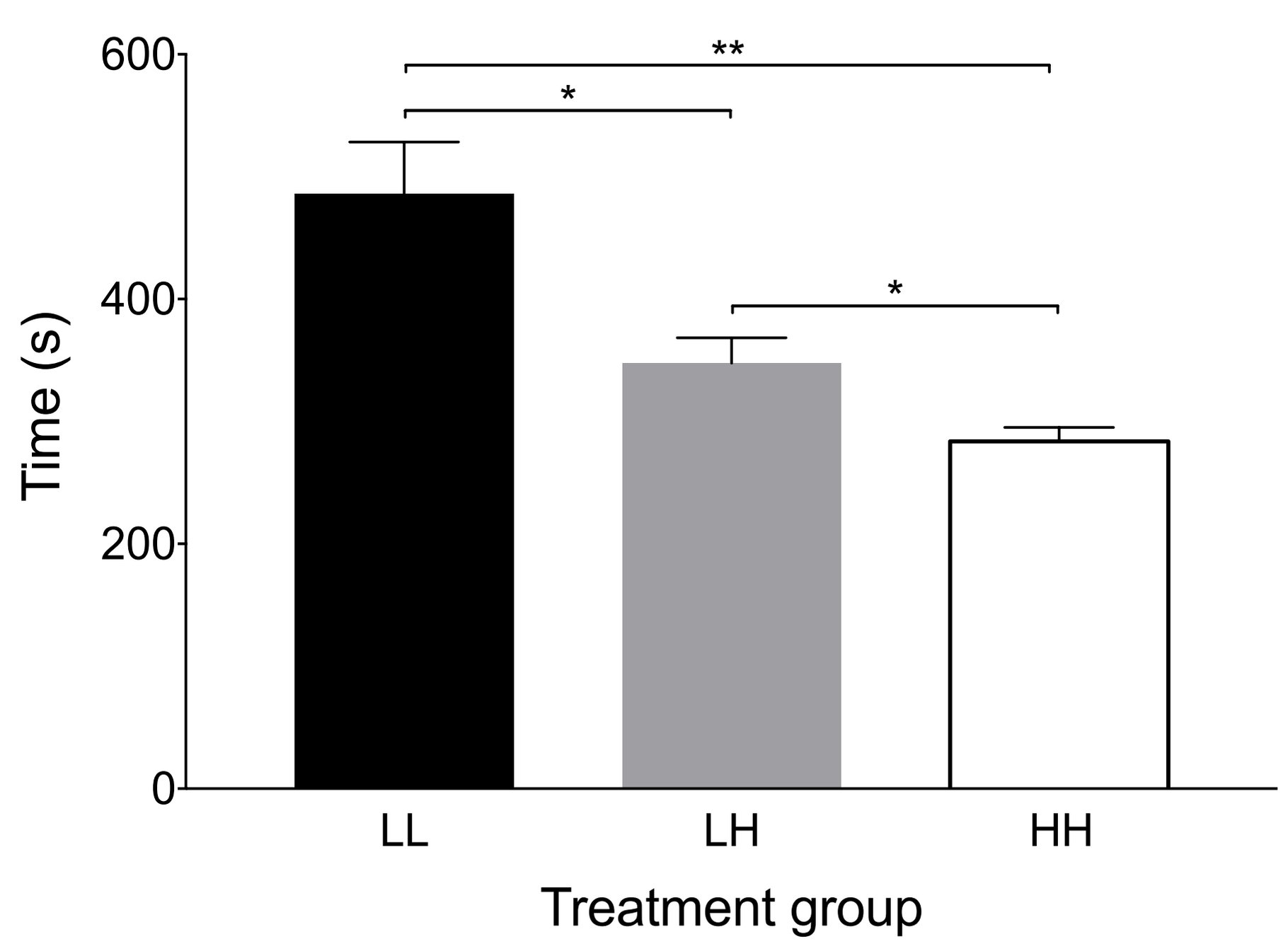
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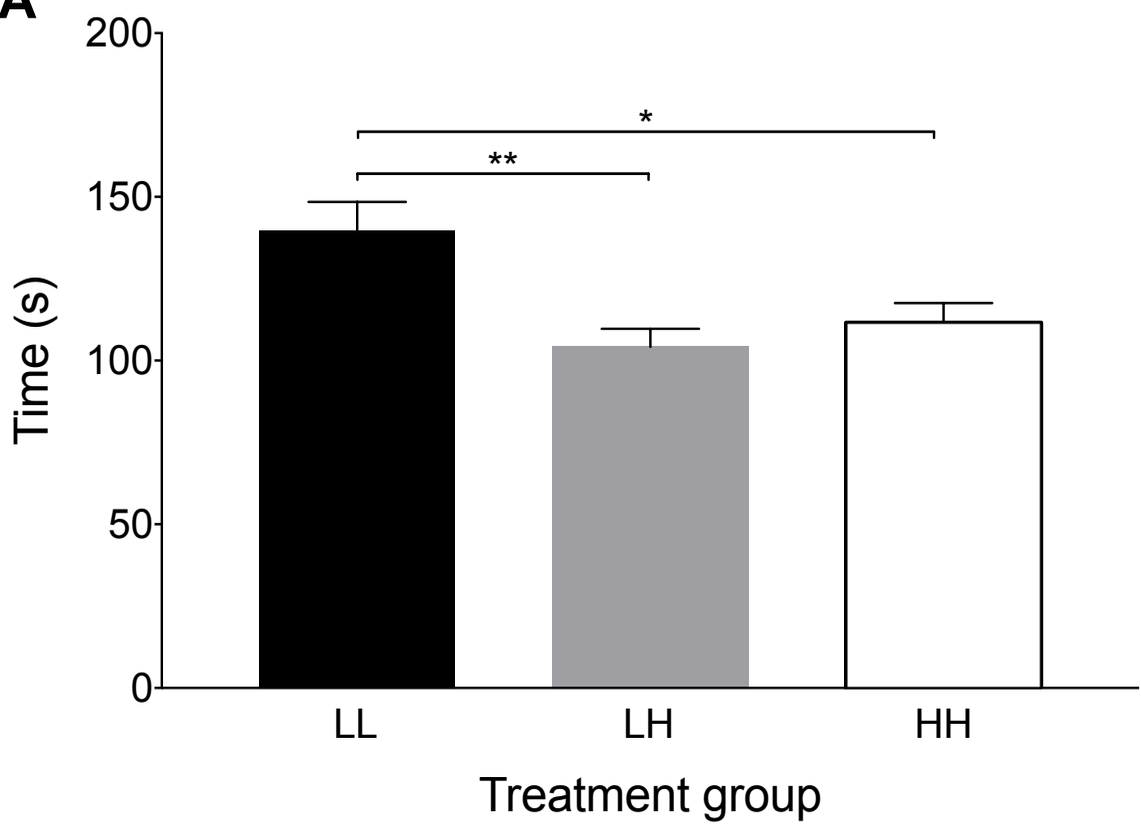
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A**B**