1 2 MISS TEAGAN GALE (Orcid ID : 0000-0003-0259-5071) 3 PROFESSOR ROB BROOKS (Orcid ID : 0000-0001-6926-0781) 4 5 6 Article type : Research Article 7 8 9 Section: Animal Growth and Development 10 Editor: Dr Christine Miller 11 12 Perceived threats of infanticide reduce maternal allocation during lactation and lead to 13 elevated oxidative damage in offspring 14 15 Teagan Gale<sup>1</sup>, Michael G. Garratt<sup>2</sup>, & Robert C. Brooks<sup>1</sup> 16 <sup>1</sup>Evolution and Ecology Research Centre, School of Biological, Earth and Environmental Sciences (BEES), the 17 University of New South Wales, High Street, Kensington, NSW 2052, Australia. 2Department of Pathology, 18 University of Michigan Medical School, Ann Arbor, MI 48109, United States. 19 Corresponding Author: Teagan Gale: t.gale@unsw.edu.au 20 Abstract 21 1. Maternal investment is costly to the mother but essential to offspring survival in 22 altrical species. Infanticide by novel males results in loss of maternal investment, and 23 maternal strategies have evolved to mitigate these losses. One such maternal strategy, 24 the Bruce effect, involves spontaneous abortion by females of some mammal species 25 when exposed to a novel male during pregnancy. 26 2. In mice, the Bruce effect only occurs during early pregnancy, but we have previously 27 found that female mice exposed to a novel male's scent in late pregnancy weaned 28 smaller offspring. Here we replicate that manipulation in order to resolve the cause of 29 the reduced weaning weight and subsequent effects on offspring fitness.

This is the author manuscript accepted for publication and has undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi: 10.1111/1365-2435.13146</u>

- 30 3. Females exposed to an unfamiliar male's scent in late pregnancy spent significantly
  31 less time nursing their pups during lactation, suggesting that reduced maternal
  32 allocation contributes to slower offspring growth. The offspring with a reduced
  33 weaning weight exhibited catch-up growth and reached a normal weight at adulthood.
  34 These offspring, however, were found to bear oxidative damage in adulthood,
  35 revealing long-term effects on offspring condition.
- 4. We conclude that female mice strategically alter their investment in lactation in
  relation to the likelihood of infanticide, but that this results in long term fitness costs
  to their offspring.
- 39

40 Running head: Female mice reduce investment when they perceive a threat of infanticide
41 Key words: catch-up growth, infanticide, maternal effects, oxidative stress.

#### 42 Introduction

43

44 In mammals, reproduction is characterised by large maternal investments in pregnancy, lactation 45 and other behaviours critical for offspring survival, growth and development (Wolff and Sherman 46 2008). This investment such as nutrition, warmth, and protection can be costly to the mother 47 (Bronson 1989; Clutton-Brock and Vincent 1991), and those costs are exacted in the currency of 48 residual reproductive value (König et al. 1988). Generally, the relationship between mother and 49 offspring is a cooperative interaction, with the mother benefitting from the success of her progeny. 50 However, with current and future reproduction known to trade-off (Reznick 1985; Stearns 1992; 51 Kirkwood and Austad 2000), a mother can also benefit from altering, and sometimes reducing, her 52 investment in current offspring according to environmental conditions, in order to optimize her overall fitness. 53

54

55 In mice (as in most mammals) lactation is the most energetically demanding episode of 56 reproduction (Speakman 2008) and is a key time to modify investment in relation to contextual 57 conditions. Maternal investment in lactation is somewhat flexible in mice (König 1985; König and 58 Markl 1987), and offspring do not get more milk than corresponds to the mother's optimal 59 allocation during lactation, despite frequent suckling attempts (König and Markl 1987). Other 60 species, such as wild primates have been observed to alter maternal investment and wean offspring prematurely when an unfamiliar male enters the natal group (Teichroeb and Sicotte 2008; Zhao et 61 62 al. 2011). Such flexibility may significantly enhance maternal fitness. 63

64 Females have also evolved other strategies to optimise their maternal investment when faced 65 with the prospect of infanticide. Infanticide represents one of the more extreme forms of 66 sexual conflict that results in a significant loss (and waste) of maternal investment for 67 females. Many mammalian species have been documented to commit infanticide, which is 68 thought to be an adaptive strategy for the perpetrator to enhance their reproductive success (Agrell et al. 1998), provide nutritional benefits, increase access to limited resources, 69 70 increase reproductive opportunities, or ensure parental care is directed to the perpetrator's 71 own offspring (Ebensperger and Blumstein 2007). Social counter-tactics to infanticide may 72 include forming affiliative relationships with adult males (Smuts 1985; Palombit et al. 1997; 73 Borries et al. 1999; Palombit 2000; Weingrill 2000), cooperative defence (see Van Schaik 74 and Janson 2000; Pradhan and van Schaik 2008; Teichroeb et al. 2012), changes in group 75 sex-ratio (see Van Schaik and Kappeler 1997; Van Schaik and Janson 2000; Pradhan and van 76 Schaik 2008; Teichroeb et al. 2012), or multiple male mating to confuse paternity (see Van 77 Schaik and Janson 2000; Wolff and Macdonald 2004). It is also of note that female 78 infanticide is also common in rodents (Vom Saal et al. 1995; Yl et al. 1997).

79

The threat of infanticide has also been shown to have effects beyond the killing of the 80 81 offspring in mice. Parkes and Bruce (1961) reported that recently inseminated female mice (*Mus musculus*) terminate their pregnancies following exposure to the urinary scent of an 82 83 unfamiliar male, and return to oestrus. This phenomenon, now known as the 'Bruce effect', 84 has been experimentally confirmed in numerous laboratory studies in house mice (Bruce 85 1960; Chipman and Fox 1966) and across a number of murine and microtine rodent species 86 (Clulow and Clarke 1968; Clulow and Langford 1971; Mallory and Clulow 1977). It has also 87 recently been observed in the wild in the primate *Theropithecus gelada* (Roberts et al. 2012). 88

89 The Bruce effect only occurs in mice if the female is exposed in early pregnancy, up to 4-5 90 days post mating (Bruce 1961), before embryo implantation. We have shown that if female 91 mice are exposed to an unfamiliar male in late pregnancy (approximately day 14), offspring 92 are of comparable birth weight, but growth is slower over lactation, resulting in lighter 93 weaning weights (Gale et al. 2013) compared with controls. Following weaning however, 94 the experimental offspring grew faster and caught up to the control offspring by the onset of 95 adulthood. Reduced weaning weight is possibly due to alteration of maternal behaviour 96 during lactation. As unfamiliar males typically kill pups sired by another male (Vom Saal 97 1985), the females may strategically reduce their investment in lactation to prevent wasting it

98 on 'doomed' offspring. If so, females are capable of utilising subtle tactics to optimise their 99 investment in reproduction in relation to their perceived chances of offspring survival. In 100 commensal habitats mice generally live in groups consisting of a number of breeding 101 females, potentially some non-breeding females and subordinate males, and a dominant male 102 who aggressively defends his territory against unfamiliar males (Hurst 1990). As territory 103 turnover does frequently occur (Oakeshott 1974; Bronson 1979), these counteradaptations to 104 the infanticide threat the unfamiliar presents are of great advantage to these females. 105 However, whether reduced offspring growth was a consequence of a strategic alteration of 106 investment by mothers in that study is unknown. Other possible causes of the offspring's 107 reduced weaning weight could include stress triggered by the unfamiliar male (stress has 108 previously been found to result in litters with a reduced weaning weight following lactation 109 (Barlow et al. 1978; Kinsley and Svare 1988; Meek et al. 2000)), or an epigenetic 110 modification in offspring gene expression.

111

112 Accelerating growth requires an increase in metabolic activity that can damage the organism 113 (Morgan et al. 2000). One such cost of accelerated growth, oxidative stress, has been documented in zebra finches (Alonso - Alvarez et al. 2007) and damsel flies (De Block and 114 115 Stoks 2008). Oxidative stress results from an imbalance between the production of harmful 116 reactive oxygen species (ROS) and an organism's ability to mitigate and detoxify the 117 damaging effects (Monaghan et al. 2009). Failure to moderate this balance can result in 118 oxidative damage to key biological molecules such as DNA, proteins and lipids (Monaghan 119 et al. 2009) and can limit investment in other life history stages (Costantini 2008).

120

127

Modifications in maternal investment have been found to have further effects other than just a reduced weaning weight. Mouse pups show great developmental plasticity, with various aspects of their early environment affecting their life-history trajectories and having lasting effects on adult phenotypes. Through the influence of maternal effects, mothers may alter the phenotype of their offspring and affect their physiological, sexual and behavioural responses as adults (Rossiter 1996; Sheldon and West 2004).

- 128 The aims of this investigation were to test if the degree of maternal investment during
- 129 lactation is affected by the threat of infanticide, and whether any such effects on maternal
- 130 investment impose further costs to the offspring later in life. We predicted that:

- 131 1) When females are faced with the threat of infanticide they will strategically alter
- 132 investment in their offspring by changing their behaviour during lactation.
- 1332) Offspring that accelerate their growth rate following a reduced weaning weight will suffer
- 134 oxidative costs as a consequence of the rapid growth.
- 135 3) Offspring that have a reduced weaning weight will suffer costs to components of
- 136 reproductive effort such as scent-marking rates and composition for males and future
- 137 reproduction for females.
- 138

# 139 Material and methods

140 This study aimed to further investigate the cause of the reduced weaning weight we 141 previously observed in offspring whose mothers had been exposed to a novel male and also 142 test for further effects to the offspring's fitness. To do this we replicated the experimental 143 manipulation of our previous experiment (Gale et al. 2013) exposing females to either an 144 unfamiliar male or the paternal male late in their pregnancy on three occasions over the last eight days of gestation. Building on from this previous experiment we added additional 145 measures to attempt to elucidate the cause of the reduced weaning weight observed in the 146 147 offspring and attempt to detect any costs of the altered growth rate. We added a third 148 unmated control treatment, in which females were housed with another female instead of a 149 male. The unmated treatment acted as a control for female weight and our marker of stress (glucocorticoid metabolites). We then conducted scan sampling during lactation where we 150 151 examined female nursing behaviour, collected fecal samples to measure female stress, and 152 measured offspring oxidative stress levels at adulthood to test for oxidative damage 153 associated with compensatory growth. We also tested aspects of reproductive fitness in the 154 offspring including scent-marking rates and major urinary proteins in the males and we 155 mated the female offspring to assess their pup birth and growth rates.

156

# 157 Animal Housing

158 Experimental mice were all sixth-generation captive-bred house mice (*Mus musculus*)

- 159 originally derived from a population of wild mice acquired from a chicken farm in the
- 160 Northwest of Sydney, Australia (as in Gale et al., 2013). Females were weaned at 28 days of
- age and were housed with their female siblings until the beginning of the experiment. Males
- 162 were also weaned at 28 days of age but were housed individually. Mice were maintained on
- 163 a 12:12 hour reversed light cycle. A dim red light was used for experimental procedures,

which were all undertaken in the dark phase as mice are nocturnal. Each mouse was housed
in a 315 x 180 x 125mm cage lined with corncob bedding and provided with tissues and
shredded newspaper for bedding. Vella Rat and Mouse Pellets and water were provided *ad libitum*.

168

#### 169 Experimental protocol

170 To investigate the consequences of a novel male's presence to pregnant females (post-171 implantation of the embryo) we compared two controls (mated and unmated) with the 172 experimental 'novel male' treatment. The unmated control was exposed only to the presence 173 and odour of another unfamiliar female. The mated control or 'familiar male' treatment 174 females were mated and then exposed to the scent and presence of the same mate. Our third 175 treatment was the novel male treatment that was experimentally manipulated so that late 176 gestation the mated females were exposed to the scent and presence of a different male who 177 was unrelated to the mate. Because exposure to this new male was late in pregnancy, well 178 beyond the time frame when a Bruce effect is known to occur in mice, females carried 179 pregnancies through to parturition, therefore allowing us to examine whether females alter 180 their reproductive investment in their offspring and whether this has any long-lasting effects 181 on the offspring (as previously used in Gale et al. (2013).

182

183 To begin each treatment replicate, a male (or another female for the unmated control) 184 unrelated to the focal female (all between 80-120 days of age) was placed into the focal 185 females cage for a two-week mating period. Males were subsequently removed, and the 186 females were transferred to a clean cage. Female weights were recorded every three days, a weight gain of 3g indicated pregnancy (mean days  $\pm$  s.e. from reaching the 3g weight 187 188 threshold to giving birth for females exposed to the male they mated with:  $6.9 \pm 1.7$ ; females 189 exposed to an unfamiliar male:  $7.1 \pm 1.9$ ). As the gestation period of *Mus musculus* is 21 190 days (Jones et al., 2009), when the exposures took place the females were close to parturition and the embryos had undergone substantial development. When the 3g-weight gain threshold 191 192 was reached, females were sequentially allocated into a treatment, to either exposure from 193 the paternal male (n=25) or a novel male (n=24). The unmated control females had already 194 been assigned to be exposed to another female (n=25). As the unmated control females did 195 not gain weight concurrent with pregnancy, one unmated control female was randomly 196 assigned to begin their exposure each day that a female from a different treatment hit the 197 weight gain threshold

199 The mice (males or females) added to the females' cages were separated by a metal divider 200 with nine small holes (5 mm diameter) which only allowed limited contact. The exposures 201 took place for three hours a day for three consecutive days. On each day a handful of the 202 respective male or female's used bedding (approximately 15-20g) was placed into the 203 female's cage. Following exposure, the female (prior to parturition) was placed into a clean 204 cage. There were no significant differences between the treatments in the number of days 205 between the removal of the breeding male and the beginning of the male exposure (mean 206 days  $\pm$  s.e. for females exposed to the male they mated with: 2.16  $\pm$  1.1; an unfamiliar male: 207  $2.08 \pm 1.2$ ). There was also no difference in the time between the exposure and the 208 subsequent birth (mean days  $\pm$  s.e. for females exposed to the male they mated with: 3.2  $\pm$ 209 1.9; an unfamiliar male:  $3.7 \pm 1.5$ ).

210

198

To test for further costs associated with reduced investment during lactation, offspring were tested for either effects to their reproductive fitness or for oxidative damage. At 13 weeks of age half of the male and female offspring from both treatments were randomly assigned to a reproductive fitness or oxidative stress group. The females in the reproductive fitness group were mated to an unrelated male and the size, weight and pup mortality was recorded. The males in the reproductive fitness group were tested for scent-marking rates and concentration of major urinary proteins.

218

#### 219 Maternal Investment and offspring weights

The body weights of the females were recorded every three days throughout the experiment.

221 Offspring were weighed at birth and every three days during lactation (to the nearest 0.1g) as

- a quantification of maternal investment (Ortiz et al. 1984; Ross 1988; Pontier et al. 1989).
- 223 The offspring were weighed collectively as a litter until weaning (4 weeks old), after which
- they were separated from their dam, and an individual weight was recorded. Another
- individual weight was collected for each of the offspring at adulthood (13 weeks old).
- 226

### 227 Maternal behaviour

- 228 Maternal behaviour was evaluated by observing each of the females using a scan –sampling
- technique twice a day, every day for the duration of their lactation (four weeks).
- 230 Observations were conducted during the dark phase using only a head torch with red light.

231 The first observation occurred in the second or third hour after the change from the light to 232

- dark period and the second between four hours and one hour before the change from the dark
- 233 to light period. Each of the females was randomly assigned an observation order at the
- 234 beginning of each observation period and observations begun ten minutes after the red light
- was turned on to allow them to habituate. 235

and the second second

236

237 A single observer sequentially recorded each female's behaviour for a total of eight 238 observations with a five minute gap between each of the eight recordings. As this was done 239 morning and evening each day it totaled 16 scans a day. Behaviours recorded included; in 240 nest, nursing, grooming, licking pups, eating/ drinking, nest building, resting or active (e.g. 241 see (Benus and Rondigs 1996; Koteja et al. 1999; Palanza et al. 2002). The unmated control 242 females were not included in any of the scan sampling of maternal behaviour.

243

#### **Glucocorticoid metabolites** 244

245 To determine whether the novel male elicits a stress response from the female, fecal samples 246 were obtained from the females at three points through the experiment and tested for 247 glucocorticoids metabolites (using methods as described by Palme and Möstl (1997).

- 248 Following a stressful event glucocorticoids are released into circulation (Sapolsky et al.
- 249 2000) and are hence used as an indicator of the stress response. The first sample was
- 250 taken at the beginning of the experiment (three weeks prior to being paired for mating), the

251 second on the second day of male exposure, and the third on day seven of lactation. The

252 unmated females were randomly allocated for fecal sample collection in order to correspond

253 with the timings of the females in the other treatments.

For collection of fecal samples, mice were placed in a large empty cage (565 x 387 x 254

255 203mm) made from H.D. polyethylene with a wire roof, for a maximum of 45 minutes. The

256 cages were placed topside down over another corresponding cage and fecal samples were

- 257 collected from the bottom cage. Immediately after collection, fecal samples were frozen at -80 °C.
- 258

259

260 Fecal samples were homogenized and an aliquot of typically 0.05g faeces (Palme et al. 2013) 261 was extracted with 1ml of 80% methanol for 30min on a vortex. When there was insufficient

- 262 sample the protocol was adjusted accordingly (e.g., 0.25 g faeces in 2.5 mL methanol).
- 263 Samples were placed in a spinner overnight and then the supernatant was diluted (1:1000)
- 264 with assay buffer (Trizma, pH 7.5). Samples were then analysed in a double-antibody 5a-

pregnane- 3b,11b,21-triol-20-one enzyme immunoassay (EIA) which has been validated for
use in mice to assess concentration of glucocorticoid metabolites as described by (Touma et
al. 2003; Touma et al. 2004).

268

#### 269 Oxidative Stress

At 13 weeks of age, half of the female and male offspring from the novel male and familiar male treatment groups were culled humanely by cervical dislocation, and the liver, kidney, heart and gastrocnemius muscle were quickly removed, snap-frozen in liquid nitrogen and stored at -80°C. To assess oxidative stress in the mice two biomarkers of oxidation including protein thiol content and aconitase enzyme activity were analysed in each of the tissues (Gibson et al. 2015). Both of these biomarkers correlate negatively with oxidative stress.

276

277 Protein thiol content was measured by methods described by (Di Monte et al. 1984) but
278 modified for use on a 96 well plate reader (Vasilaki et al. 2006). Protein thiols are essential

279 for stability of and optimum function of proteins, but are highly susceptible to oxidation

- 280 (Halliwell and Gutteridge 1999), and therefore good markers of oxidative stress.
- 281

282 Aconitase is an enzyme of the tricarboxylic acid cycle that is highly susceptible to 283 deactivation by radical oxygen species (specifically superoxide) and therefore used as a 284 marker to indicate levels of reactive oxygen species and concomitantly oxidative stress 285 (Hausladen and Fridovich 1994; Gardner et al. 1995; Hausladen and Fridovich 1996; Gardner 1997). As aconitase is located in part in the mitochondria (Wiegand and Remington 286 287 1986; Gardner et al. 1995), mitochondrial density was also assessed using citrate synthase 288 activity. Citrate synthase is an enzyme commonly used as an indicator of the content of intact 289 mitochondria (Holloszy et al. 1970) and was measured in homogenates according to (Pichaud et al. 2008). 290

291

### 292 Offspring reproductive fitness

293 Dominant adults are known to scent mark more regularly than subordinates (Drickamer

1995) to communicate their territorial and sexual status (Bronson 1979; Hurst et al. 2001)

and competitive ability (Rich and Hurst 1998). These chemical scent marks are of high

- importance to male fitness as they directly influence the attractiveness of a male to a female
- 297 (Rich and Hurst 1998). The main involatile scent component of male mouse urine is major

- 298 urinary proteins (MUPs) that bind volatile components of the urine and slowly release them
- from the scent marks (Hurst et al. 1998). Scent marks may not prevent intruders invading the
- 300 territory, but they do allow males to use long-lasting signals of identity and dominance over
- 301 a territory to alert competitors and potential mates (Hurst et al. 1998; Hurst and Beynon
- 302 2004). Therefore, both the rates and the composition of scent-marks can substantially
- 303 influence male reproductive fitness.
- 304

# 305 Scent-marking rates

306 Scent-marking rates of all of the individually housed male offspring were assessed at 13

307 weeks of age. Scent-marking rates were measured by placing the individual males into an

308 empty (315 x 180 x 125mm) cage lined with Benchkote for one hour a day, for three

309 consecutive days. The scent marks were measured by the number of spatially separate marks

- 310 observed under UV light and the average number of marks for each male over the three trials
- 311 was used for analysis.
- 312

# 313 Major Urinary Protein Concentration

A urine sample was collected from each of the male offspring at 13 weeks of age. Males

were confined in a large empty cage (565 x 387 x 203mm) made from H.D. polyethylene

316 with a wire roof, for maximum of 180 minutes. The cages were placed topside down over

another corresponding cage to allow the mouse urine to pool in the bottom cage. Urine was

then pipetted into an Eppendorf tube and frozen at -20 °C. The concentrations of major

- 319 urinary proteins were established using Coomassie plus® protein assay reagent kit from
- 320 Perbio Science UK Ltd (Cramlington, Northumberland, UK) as described by Cheetham et al.
- 321 (2009). We also measured urinary creatinine (Beynon and Hurst 2004) using the method of

322 Cheetham et al. (2009) to correct for the urinary dilution.

323

# 324 Statistics

325 All statistical analyses were performed using SPSS software package version 2.1 (IBM Corp,

326 Armonk, NY, USA). The analyses were done with dam ID and experimental block fitted as

327 random factors to account for non-independence of individuals originating from the same

328 litter and time differences of each group of experimental mice unless otherwise described.

329 For the oxidative stress data we also fitted plate number as a random factor to control for

between plate variability. Scent marking frequency was transformed to log (x+1) to account

for measures of zero deposits and normalise the data. Significance was determined at  $p \le 0.05$ .

332 **Results** 

#### 333 Offspring weight

334 In the novel male treatment, the females gave birth to pups that were of similar weight and litter size as the familiar male treatment females (i.e. exposed to the paternal male). Out of 335 336 25 females mated in the familiar male treatment, 23 mothers gave birth within the time frame 337 with a mean litter size of 4.217±0.77 and out of 24 mothers in the novel male treatment, 21 338 gave birth within the timeframe with a mean litter size of 4.14±0.89. The mean birth weight 339 of novel male treatment pups  $(1.619 \pm 0.432g)$  was not significantly different from that of the 340 familiar male treatment (1.628  $\pm$  0.433g; ANOVA: weight: F<sub>1,182</sub>= 0.022, p= 0.882; litter size:  $F_{142} = 0.070$ , p= 0.792). Two females in the control treatment and one female in the 341 342 experimental treatment group destroyed their litters within three days of giving birth. Some 343 pup mortality of unknown cause was observed over lactation; however, this is common in 344 captive breeding mice (Weber et al. 2013). The pup mortality was not significantly different 345 between the treatments (ANOVA:  $F_{1,7}$ = 0.778, p= 0.407) with six mothers of the novel male 346 treatment committing maternal infanticide (total 10 pups lost) and three mothers of the 347 familiar male (total seven pups lost).

348

349 Although the novel male treatment females gave birth to pups of a similar weight and litter 350 size as the familiar male treatment females, novel male treatment litters grew more slowly 351 over lactation. To analyse the differences in offspring growth rate we used repeated measures 352 ANOVA reporting within-subjects effects. Mauchly's Test of Sphericity indicated that the 353 assumption of sphericity had been violated ( $\chi 2(2) = 34.210$ , p = <0.001), therefore degrees of 354 freedom were corrected using Greenhouse-Geisser adjusted degrees of freedom. There was a 355 difference in growth rate between the treatments but there was no effect of sex (treatment:  $F_{1,7,310}=7.49$ , p= 0.001; interaction between treatment and time:  $F_{1,7,310}=5668$ , p= <0.001; 356 357 sex x treatment:  $F_{1.6,306}=1.3$ , p= 0.270). This resulted in novel male treatment pups being 358 significantly smaller (7.19 $\pm$ 0.61g), at weaning (4 weeks old) (ANOVA: F<sub>1.182</sub>=27.11, 359 p = < 0.001) than the familiar male treatment pups (7.99 $\pm$ 1.48g) (see Figure 1). This difference 360 did not persist into adulthood however, as both males and females of the novel male 361 treatment exhibited catch-up growth following weaning and there were no weight differences 362 at adulthood (13 weeks old) (ANOVA:  $F_{1, 184}=0.760$ , p=0.384) (see Figure 2). There were

363 also no significant interactions of treatment and sex in weight at weaning (sex x treatment 364  $F_{1,180}$  = 2.14, p=0.145) or at adulthood (sex x treatment  $F_{1,180}$  = 0.044, p=0.868)). 365 366 Insert Figure 1 and 2 367 368 **Maternal behaviour** 369 There was no significant difference in the amount of time the mothers spent in nest, 370 grooming, licking pups, eating/drinking, nest building, resting or active (Table 1). The only 371 behaviour showing significant differences between treatments was nursing, with females 372 exposed to familiar males spent almost twice the time nursing (Table 1). 373 374 Insert Table 1 375 376 **Glucocorticoid** metabolites in mothers 377 Three fecal samples were taken from each female: The first sample was taken before the 378 experiment began to determine the normal level of fecal glucocorticoid metabolites (FGMs) 379 for each female. The second collection was taken on day two of the exposures to a male and 380 the third coincided with mid lactation in the mated treatments. Some of the females did not 381 provide a sample within the timeframe they were in the collection cages (45min) so the 382 sample sizes for the glucocorticoid metabolites are novel male n=20, familiar male n=21 and 383 unmated control n= 18. Using a repeated measures ANOVA we found that FGM's did 384 change significantly over time (effect of time:  $F_{2, 112} = 131$ , p= <0.001; interaction between 385 time and housing companion:  $F_{4,112}$  = 29.5, p<0.001). In the novel male and familiar male 386 treatments the FGM's were higher in the second collection and dropped back down closer to 387 their normal levels of FGM's at the third collection. The was no difference in the FGMs in 388 the females before the experiment began (ANOVA:  $F_{2,56}=0.094$ , p=0.911), but there was 389 following exposure to their partners (ANOVA:  $F_{2.56}$ =43.72, p=<0.001) and at the end of the 390 experiment which coincided with mid lactation for the reproducing treatments ( $F_{2.56}$ =4.6, 391 p=0.014)(See Figure 3). After removing the unmated control from the analysis we found that the effect of time was still significant ( $F_{2,78}$ = 143, p=<0.001) but the interaction between 392 393 time and housing companion was not ( $F_{1,78}$ =1.89, p=0.158). While there was no difference 394 between the FGMs between the novel male treatment and the familiar male treatment, there 395 was a difference between the mating treatments and females who were not mated. Females 396 from both mated treatments experienced a rise in FGM's levels in the second collection 397 following exposures to a male, however, by the third collection at mid lactation their FGM's

398	had returned closer to their normal level (collection one). The unmated control females FGM
399	levels did not show any pronounced variability over the three collections.
400	
401	Insert Figure 3
402	
403	Oxidative stress in offspring
404	Both markers of oxidative stress showed consistent evidence of oxidative damage with the
405	novel male treatment (n=47) in two (liver and heart) out of four organs tested having lower
406	protein thiol concentrations and aconitase enzyme activity (indicating higher levels of
407	oxidative damage) than the familiar male treatment (n=38). There was an effect of sex with
408	novel male treatment offspring males having lower protein thiol concentrations than those of
409	the familiar male treatment males in the heart but there were no other sex differences in any
410	of the analyses for oxidative damage (see Table 2 and Figure 4 and 5).
411	
412	The same trend was observed when testing the aconitase enzyme activity to citrate synthase
413	activity (mitochondrial density) ratio. The liver and heart in the novel male treatment were
414	also found to have a significantly lower aconitase enzyme activity to citrate synthase activity
415	ratio indicating higher levels of reactive oxygen species (ROS) production in these organs.
416	This decrease in aconitase enzyme activity was not observed in either the kidneys or the
417	gastrocnemius and there were no effects of sex (See Table 2 and Figure 4 and 5).
418	
419	Insert Table 2 and Figure 4
420	
421	Male and female offspring fitness components
422	For the male offspring fitness components sample sizes were both $n=17$ for the treatments.
423	There was no difference in the concentration of major urinary proteins among the male
424	offspring produced by the novel male treatment females compared to the control offspring
425	(ANOVA: $F_{1,30}$ = 0.76, p= 0.543). The mean protein (mg/mg creatinine) ± s.e for the novel
426	male treatment was $18.44 \pm 3.60$ , and the familiar male treatment males were $20.73 \pm 2.74$ .
427	Experimental block also did not have an effect (ANOVA: $F_{1, 30}$ = 5.8, p= 0.249). The mean ±
428	s.e scent-marking frequency for the novel male treatment offspring was 28.28±10.54 and the
429	familiar male treatment males was 36.96±10.49. A repeated measures ANOVA found that
430	there was no difference in the frequency of scent-marks between the treatments ( $F_{2, 64}$ = 3.00,
431	p=0.57). For the female offspring produced that were bred at adulthood there was no
432	difference in the pup weights that the female offspring gave birth to (ANOVA: $F_{1,40}=0.35$ ,

- 433 p=0.556) or of their pups weaning weights at the end of lactation (F<sub>1,40</sub>=1.002, p=0.323) and
- 434 a repeated measures ANOVA showed there was no difference in growth rate between the
- 435 treatments (Sex x Treatment  $F_{1,41}$ =1.29 p=0.262). The mean birth weights between the
- 436 treatments were  $1.26 \pm 0.04$ g and  $1.355 \pm 0.13$ g for the novel male and familiar male
- 437 treatment females respectively and the mean weaning weights were  $12.75 \pm 2.09$ g and  $12.27 \pm$
- 438 0.13g respectively. Experimental block also had no effect (ANOVA:  $F_{1,40}$ = 0.69, p= 0.837).
- 439

## 440 Discussion

Our results support the prediction that the 'Late Bruce Effect', in which female mice exposed to a novel male late in pregnancy weaned at lower weights (Gale et al. 2013), may be due to a strategic reduction in maternal investment in lactation. Notably, females exposed to a novel male's scent spent less time nursing pups than females exposed to the scent of their mate. Our results support the prediction that females adjust post-partum investment in relation to pups' perceived chances of survival, potentially benefitting maternal lifetime fitness at the expense of the current litter of offspring (Marshall and Uller 2007).

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449 This finding suggests that patterns of post-partum investment can be adjusted by mothers in 450 relation to the risk of infanticide, in ways that mirror the strategic, spontaneous abortion of 451 pups under the Bruce effect (Bruce 1960, 1961; Hrdy 1979; Schwagmeyer 1979; Storey 452 1986), as well as evidence that a mechanism to abort fetuses strategically at the later stages 453 of pregnancy has evolved in other species including the gelada (Roberts et al. 2012) and 454 prairie vole (Clulow and Clarke 1968; Clulow and Langford 1971). As mice are only able to 455 abort in early pregnancy, altering their investment post-partum provides an opportunity to 456 respond to the threat of infanticide.

457

458 Other studies have shown that female mice strategically moderate investment during 459 lactation and that they may alter investment relative to the reproductive value of their 460 offspring. Mashoodh et al., (2012) found that females increased their investment during 461 lactation when mated with a male housed in lifelong socially enriched conditions compared 462 with females mated with males housed in impoverished conditions. This suggests that 463 females may invest more in offspring on the basis of paternal condition. Konig and Markl 464 (1987) showed that despite frequent suckling attempts, offspring do not get more milk than 465 corresponds to the maternal optimum during lactation, suggesting mothers have control over 466 reproductive allocation during lactation.

468 Another potential cause of the reduced weaning weight could be a stress response. In 469 rodents, maternal stress has been previously found to cause depressed growth in offspring 470 (Barlow et al. 1978; Kinsley and Svare 1988; Meek et al. 2000). Previous research on the 471 effects of prenatal stress on offspring is frequently contradictory. The nature, timing and 472 length of the stress inflicted varies from study to study, as do the results. One study, for 473 example that used crowding as a stressor found no difference in time spent nursing between 474 stressed and non-stressed dams (Moore and Power 1986), while another study that used 475 novelty stress found that stressed dams spent significantly more time nursing compared to 476 non-stressed dams (Muir et al. 1985). It is also important to note there is evidence that 477 maternal stress leads to faster (Dantzer et al. 2013) and increased growth of offspring at birth 478 and weaning (Szuran et al. 1991).

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480 We found while females all had similar stress levels before the experiment began, females 481 that were mated and then exposed to either a novel or their mated male had significantly 482 higher FGMs than the females that were not mated in the experiment. Females from the 483 mated treatments both displayed a rise in their FGMs following the exposures, suggesting 484 that the females were equally stressed by encountering the paternal male as they were a novel male. Higher levels of FGM could also result from exposure to a male, as our 485 486 experiment used another female as an unmated control we are unable to distinguish this 487 result properly. Only one other study we could find used a novel male conspecific as a 488 source of stress. Lerch et al. (2016) stressed pregnant or lactating females using unfamiliar 489 male faeces and examined maternal and offspring behaviour to investigate whether early 490 adverse experiences elevate the risk of developing psychiatric disorders. However, unlike 491 our study that used a familiar male as a control, they compared pregnant or lactating females 492 exposed to unfamiliar male faeces with a control group that didn't receive any faeces. The 493 scarcity of research that uses another conspecific as a stressor is surprising as this could be 494 expected to be a more environmentally relevant challenge that other manipulations like 495 forced immobilization challenges, for example. While the over production of maternal 496 glucocorticoids can be harmful (Korgun et al. 2012) levels are known to increase during 497 pregnancy in mice (Barlow et al. 1974; Dalle et al. 1978) as they are essential for fetal 498 development (Korgun et al. 2012). Post -partum, glucocorticoids also have important roles in 499 milk secretion and lactogenesis (Chida et al. 2011). The higher FGM levels that we found in

500 females from both of the mated, compared to the unmated treatment, may, therefore, be just

a normal consequence of pregnancy.

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503 As in our previous study (Gale et al. 2013), the offspring of the females that were exposed to 504 the novel male were smaller at the end of lactation, but they caught up in size by maturity. The catch-up growth exhibited by the offspring coincides with the time after lactation when 505 506 offspring begin to feed themselves on solid food. Offspring can acquire the resources to 507 accelerate their growth rate themselves. For compensation or catch-up growth to occur, the 508 benefits must outweigh the costs of not accelerating growth. While there may be a positive 509 association between size and fitness (Roff, 1992) and accelerated growth may increase 510 overall reproductive success, many studies have found that compensatory growth inflicts 511 various costs over different time-scales (reviewed in Metcalfe and Monaghan (2001)). Costs 512 that have been documented in rats (*Rattus norvegicus*) include deficiencies in protein 513 maintenance (Samuels and Baracos 1995), telomere abrasion rate (Jennings et al. 1999) 514 insulin regulation (Ozanne and Hales 1999), adult obesity (Waterland and Garza 1999), and, 515 perhaps most importantly, reduced lifespan (Jennings et al. 1999; Rollo 2002). While 516 compensatory growth may reduce lifespan it could still be adaptive if it increases overall 517 reproductive success (Metcalfe and Monaghan 2001) or short-term survival chances (Arendt 518 1997). In mice, reproductive allocation in adulthood is influenced by size, and so 519 compensatory growth may allow individuals to attain a normal reproductive rate in 520 adulthood, at least early in life. Female offspring that exhibited catch-up growth produced 521 litters of the same size and weight as those female offspring from the control group; male 522 offspring showed equivalent scent marking abilities in our assays. Thus, catch up growth 523 seems to allow offspring to attain a similar reproductive output early in life that is 524 comparable with the steadier growth in the familiar male group, although costs may be paid 525 for this in terms of late life reproduction or lifespan.

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527 The existence of compensatory and catch up growth shows that organisms do not grow at 528 their maximal rate, but rather at a rate influenced by, and potentially optimized to, their 529 circumstances. Hector and Nakagawa (2012) distinguish these two terms by defining 530 compensatory growth as a faster than usual growth rate and catch-up growth an attainment of 531 control size. Mangel & Munch (2005) propose that growth leads to an accumulation of 532 damage at the cellular level that is expressed at the level of the organism and is an important 533 cost of compensatory growth. We tested for damage on a cellular level in form of oxidative damage and oxidative stress. We found evidence for oxidative damage in the livers and hearts of offspring from the novel male exposure treatment, highlighting that an olfactory change in a pregnant mother's environment can elicit a variety of maternal and offspring responses, ultimately influencing offspring physiological condition in adulthood. This may be a consequence of catch-growth, but could also be a consequence of odour exposure itself, or changes in maternal allocation in response to this. In future studies it may be of interest to limit an offspring's ability to show compensatory growth (through a nutritional or genetic manipulation) and test whether oxidative damage in offspring still occurs in adulthood.

Wild mice are highly territorial and turnover of the dominant male is a common occurrence in wild populations (Oakeshott 1974; Bronson 1979). For the females in the territory that means that they will be exposed to novel males which present the threat of infanticide (Ebensperger 1998). Our experiment was designed with the rationale to mimic this turnover by exposing the females to a novel male to see the effects it would have on reproductive allocation and offspring fitness. Our results suggest that the very prospect of male territorial turnover can have physiological consequences for the offspring and potentially alter their life history. We also suggest that females are capable of strategically modulating their investment relative to their current contextual conditions, which may offer significant fitness benefits in the wild where offspring survival is much more variable and infanticide is a common threat.

- 569 We would like to thank Heather Try for all of her help with animal maintenance and Rebecca
- 570 Hobbs at the Wildlife Reproductive Centre (WRC) at Taronga Western Plains Zoo for
- 571 providing training on how to conduct an assay (using methods as described by Palme &
- 572 Möstl, 1997) for glucocorticoids metabolites in feces.
- 573

# 574 Competing interests

- 575 There are no competing interests to declare.
- 576

#### 577 Author contributions

- 578 TG, MG and RB conceived the ideas and designed methodology;
- 579 TG collected the data;
- 580 TG, MG and RB analysed the data;
- 581 TG led the writing of the manuscript.
- 582 All authors, TG, MG and RB contributed critically to the drafts and gave final approval for
- 583 publication.
- 584
- 585 Funding
- 586 This study was funded by an Australian Research Council (ARC) Discovery Grant and
- 587 Fellowship [grant number: DP150100676] awarded to R.C. Brooks.

#### 588 Data accessibility

- 589 We have archived our data on Dryad. A data set for this experiment is available from the
- 590 Dryad Digital Repository: doi:10.5061/dryad.762k24f
- 591 592
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862	Table 1: Maternal behaviour of Novel male treatment and Familiar male treatment females
863	over lactation

Comparison of the number of times maternal behaviours were recorded during lactation

Mated Control

Novel Male

		F	d.f	Р	Mean	SE	Mean	SE			
_	In nest	3.77	1,41	0.059	114.34	4.15	103.47	3.54			
	Nursing	11.56	1.41	0.002	15.52	1.34	8.4	1.53			
	Licking Pups	1.79	1,41	0.188	1.82	0.63	0.80	0.34			
	Nest Building	0.12	1,41	0.730	0.65	0.27	0.80	0.31			
	Eating/ drinking	0.14	1,41	0.707	5.82	0.62	5.52	0.52			
	Grooming	0.14	1,41	0.705	9.73	0.68	10.14	0.82			
	Active	0.78	1,41	0.382	131.34	6.21	139.76	7.45			
0.6.4	Resting	1.15	1,41	0.288	168.73	5.00	179	7.96			
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885	Table 2: Mar	kers of oxid	lative stress	in offspring	of both treat	ments at adu	ulthood. Sho	wing			
886	protein thiol of	concentratio	n (µmol g-1	protein) and	l aconitase er	nzyme activ	vity/citrate sy	nthase			
887	(units/mg protein). Models were also fitted with plate number and block as a random factor										
888	to control for variation between assay plates.										

#### Oxidative stress results for protein thiol concentration and aconitase enzyme activity

	Liver						Kidne	у						
•	Protein thiol concentration			Aconitase enzyme activity		Protein thiol concentration				Aconitase enzyme activity			_	
	F	d.f	Р	F	d.f	Р	F	d.f	Р		F	d.f	Р	
Treatment	60.65	1,80	<0.001	14.1	1,81	<0.001	1.44	1,80	0.23	4	1.47	1,80	0.228	3
Sex	0.130	1,80	0.719	0.20	1,81	0.656	0.01	1,80	0.99	7	0.32	1,80	0.569	)
Sex <sup>x</sup> treatment	1.43	1,80	0.234	1.59	1,81	0.210	0.61	1,80	0.43	6	1.85	1,80	0.177	,
	Heart						Gastrocnemius							
	Protein	n thiol		Aconi	tase ei	nzyme activity	Pr	otein th	iol con	centratio	on	Aconita	se enzyr	ne
	concer	itration										activity		
	F	d.f	Р	F	d.f	Р	F		d.f	Р		F	d.f	Р
Treatment	10.34	1,80	< 0.001	20.42	1,8	1 <0.001	0.1	17	1,80	0.681		1.76	1,80	0.188
Sex	8.49	1,80	0.005	0.98	1,8	1 0.755	2.8	38	1,80	0.093		0.78	1,80	0.379
Sex <sup>x</sup> treatment	1.80	1,80	0.183	0.41	1,8	1 0.523	0.0	001	1,80	0.971		2.11	1,80	0.150

Mean litter Mass (g) Figure 1: Average litter growth rate over lactation (4 weeks). Data are displayed as means  $\pm$ s.e.m. 

Weight (g) Figure 2: Comparison of individual offspring birth, weaning and adult weight between the two treatments. Data are displayed as means  $\pm 3$  s.e.m. Auth

Mean GC metabolites (ng/g faeces)

950Figure 3: Concentration of fecal glucocortocoid metabolites in females taken before the951experiment started, on the second after exposure to a male/female and then mid lactation in952the mated treatments. Females were either exposed to a novel male (novel male treatment)953their mated male (familiar male) or another female (unmated control). Data are displayed as954means  $\pm$  s.e.m.

# Mean protein thiol concentration per organ (µmol g<sup>-1</sup> protein)

**Figure 4:** Mean protein thiol concentrations. Data is presented as estimated marginal means 975  $\pm 1$  s.e.m. for each measure from general linear mixed models for each tissue sample.

# Mean aconitase activity per organ (units/mg protein)

992

**Figure 5:** Ratio of mean aconitase enzyme activity to citrate synthase. Data is presented as estimated marginal means  $\pm 1$  s.e.m. for each measure from general linear mixed models for each tissue sample.

Author **N**