- 1 Title: Testing for a role of *Dnmt2* in paternal trans-generational immune priming
- 2 Authors: Nora K E Schulz^{1,2,*}, Maike F Diddens-de Buhr¹ & Joachim Kurtz¹

3

4 ¹ Institute for Evolution and Biodiversity

- 5 University of Münster
- 6 Hüfferstr. 1
- 7 48149 Münster
- 8 Germany
- 9 ² Current address
- 10 Vanderbilt University
- 11 465 21st Avenue South
- 12 Nashville, TN37232
- 13 USA

14 * Corresponding author: Nora Schulz, Email: nora.schulz@vanderbilt.edu

Keywords: epigenetics, tRNA methyltransferase, tRNA methylation, *Tribolium castaneum*, immune
 priming, paternal effects

17 Abstract

Trans-generational effects from fathers to offspring are increasingly reported from diverse organisms, 18 19 but the underlying mechanisms are often unknown. Paternal trans-generational immune priming 20 (TGIP) was demonstrated in the red flour beetle Tribolium castaneum: non-infectious bacterial 21 exposure (priming) of fathers protects their offspring against an infectious challenge. Here we studied 22 a potential role of the *Dnmt2* (now also called *Trdnmt1*) gene, which encodes a highly conserved enzyme that provides CpG methylation to a set of tRNAs and has previously been reported to be 23 24 involved in transgenerational epigenetic inheritance in mice. We first studied gene expression and 25 found that Dnmt2 was expressed throughout life, with high expression in testes. Knockdown of Dnmt2 26 in fathers slowed down offspring larval development and increased mortality of the adult offspring 27 upon bacterial infection. However, the observed effects were independent of the paternal priming 28 treatment. In conclusion, our results point towards a role of Dnmt2 for paternal effects, while 29 elucidation of the mechanisms behind paternal TGIP needs further studies.

30

31 **1 Introduction**

Phenotypic plasticity is often enabled by epigenetic mechanisms ^{1,2}. An important epigenetic 32 33 modification of nucleic acids is the covalent binding of a methyl group to a cytosine followed by a 34 guanine, *i.e.* CpG methylation ³. In insects, CpG methylation of DNA has been extensively studied ^{4,5} and is involved in many processes, such as caste determination and phase polyphenism $^{6-10}$. However, 35 36 this modification not only occurs on DNA, but also on a variety of RNAs¹¹. CpG methylation is 37 facilitated by a conserved family of enzymes called DNA methyltransferases, which are found in most but not all animals ^{3,12}. *Dnmt2* is the evolutionary most conserved member of this gene family. It can 38 39 be found in many fungi, plant and animal species, sometimes occurring in the absence of any functional 40 DNA methylation machinery¹³. While Dnmt1 and Dnmt3 are responsible for modifications of DNA, Dnmt2 is involved in methylating three types of tRNA at the C38 position ¹⁴⁻¹⁶. Because of this 41 42 function, the human Dnmt2 gene has been renamed to TRDMT1 (tRNA aspartic acid methyltransferase 43 1). However, the gene is phylogenetically related to *Dnmt* genes, and it was thus suggested to stick to its original name³The methylation mark provided by *Dnmt2* protects the tRNA molecule against 44 cleavage, which can be induced by different stressors ¹⁵. It has been shown that tRNA-derived small 45 46 RNAs (tsRNAs) regulate mRNAs and therefore differences in tRNA cleavage could lead to altered 47 phenotypes. In mice dietary stress can cause increased fragmentation of tRNAs and the resulting 48 metabolic phenotype is paternally transmitted to the offspring through the altered levels of tsRNAs ^{17,18}. This paternal transmission is dependent on *Dnmt2*, which demonstrates the importance of this 49 50 gene in non-genetic inheritance ^{19,20}.

The function of *Dnmt2* has also been studied in *Drosophila melanogaster*. Mutants lacking *Dnmt2* were less protected against a variety of stressors, as increased rearing temperatures led to a reduced lifespan and herbicides caused higher mortality compared to wildtype and control flies ¹⁵. Furthermore, heat shock treatment of flies lacking *Dnmt2* led to the accumulation of transposable elements and changed gene expression ²¹. Also, other studies have demonstrated that *Dnmt2* plays a crucial role in managing endogenous and exogenous RNA stress, by silencing retrotransposons and inhibiting RNA

virus replication ^{22,23}. It has thus been proposed that the enzyme is involved in adaptive immunity and 57 aids in defending against or adapting to pathogens ¹³. 58

59 Phenotypic plasticity, both within and across generations, is widespread in immune defence. A wealth of studies in invertebrates now provide evidence for immune priming, i.e. enhanced survival upon a 60 secondary encounter with a pathogen $^{24-26}$. In some species it has been shown that the immune priming 61 can also be transferred to the offspring ^{25,27,28}. While maternal transfer appears to be a relatively 62 63 common phenomenon, reports about paternal transgenerational immune priming (TGIP) are scarce ^{25,29}. The red flour beetle, *Tribolium castaneum* is the first example where paternal TGIP against a 64 variety of bacterial pathogens has been demonstrated ^{29–31}. However, the mechanisms underlying TGIP 65 66 remain elusive. The paternal route of priming narrows down the possibilities by which effectors or 67 information could be transferred from father to offspring and thus makes the involvement of epigenetic 68 modifications, such as methylation of sperm RNA, especially likely ²⁶. Finally, in another beetle, Tenebrio molitor priming of adults and larvae decreased overall RNA methylation within the 69 generation, hinting at a possible involvement of *Dnmt2*³². 70

T. castaneum possesses two Dnmt genes: one Dnmt1 and one Dnmt2 homolog ³³. The beetle seems to 71 lack any functional levels of CpG DNA methylation ^{4,34,35}, but *Dnmt1* is nevertheless expressed across 72 all life stages and is needed for proper embryonic development ³⁵. To our knowledge no research has 73 74 been dedicated yet to study the role and function of Dnmt2 in T. castaneum. We therefore used gene 75 expression analysis and RNAi to investigate the role of this gene for paternal trans-generational effects. 76 We combined a paternal knockdown with immune priming, to investigate whether *Dnmt2* is involved 77 in and possibly provides the epigenetic mechanism behind paternal TGIP.

78 2

Materials and methods

2.1 79 Model organism

80 T. castaneum has become a well-established model organism in many fields of biology including evolutionary ecology. The availability of a fully sequenced genome 33 and molecular tools, *e.g.* easily 81 established systemic RNAi and genome editing ^{36–39} allow for genetic analyses of ecologically relevant 82 83 phenomena. For this study, the T. castaneum line Cro 1 was used, which was established from 165

pairs of wild caught beetles collected in Croatia in June 2010 ⁴⁰. Beetles were maintained in plastic
breeding boxes with foam stoppers to ensure air circulation. Standard breeding conditions were 30°C
and 70% humidity with a 12-hour light/dark cycle. As food source 250g of heat sterilized (75°C)
organic wheat flour (type550) containing 5% brewer's yeast were given.

88 2.2 Gene expression of *Dnmt2*

To assess the expression of *Dnmt2* throughout the life cycle of the beetle, the four distinct life stages 89 90 were sampled: eggs (n=4 pools of 500-1000 eggs, 24h-48h post oviposition), larvae (n=7 pools of 10 91 larvae, 14-19 days post oviposition (dpo)) pupae (n=8 pools of 6 individuals), virgin adults (n=8 pools 92 of 6 individuals, one week after eclosion). For pupae and adults, half of the pooled samples contained 93 females and the other half males in order to test also for differential expression between the sexes. 94 Furthermore, gonads were dissected from unmated adult males. All samples were shock frozen in 95 liquid nitrogen. Total RNA was extracted, and genomic DNA digested by combining Trizol (Ambion 96 RNA by Life Technologies GmbH, Darmstadt, Germany) and chloroform treatment with the use of the Total RNA extraction kit (Promega GmbH, Mannheim, Germany) as described in Eggert et al.³⁰. 97

Extracted RNA was reverse transcribed to cDNA with the RevertAid First Strand cDNA kit (Thermo 98 Fisher Scientific, Waltham, MA USA) using provided oligo-dTs. In the following RT qPCR with a 99 100 Light-Cycler480 (Roche) and Kapa SYBR Fast (Kapa Biosystems, Sigma-Aldrich), each sample was used in two technical replicates. Further analysis was conducted as described in Eggert et al. ³⁰ and 101 102 replicates were used in further analysis if the standard deviation between their crossing point values 103 was below 0.5, otherwise the reaction was repeated. Previously, high primer efficiency had been 104 confirmed and where possible it was made sure that primers crossed exon-intron boundaries (Table 105 S1). The housekeeping genes ribosomal proteins rp49 and rpl13a were used for normalization of the 106 expression of the target genes.

107 2.3 Paternal *Dnmt2* knockdown and TGIP

We aimed to downregulate *Dnmt2* through paternal RNAi and to investigate whether this knockdown
would affect paternal TGIP (Figure 1). For this, around 2000 adult beetles (one week old) were allowed

to lay eggs for 24h. Two weeks later, larvae were collected and put into individual wells of a 96 well
plate, which contained flour and yeast. The oviposition was repeated with two further, independent
cohorts on the two following days, producing three experimental replicates.

113 2.3.1 Paternal RNAi

114 The sex of the beetles was determined in the pupal stage, and male pupae were prepared for RNAi 115 treatment, while females were individualized and kept for mating. For injections of dsRNA, male 116 pupae (22 dpo) were glued with the hindmost segment of the abdomen to a glass slide to immobilize 117 them. One glass slide held between 16 and 20 pupae. Pupae were either injected with dsRNA of the 118 target gene Dnmt2 or for the control of the treatment procedure with dsRNA transcribed from the 119 asparagine synthetase A (asnA) gene found in Escherichia coli (RNAi control), which bears no 120 sequence similarity to any known T. castaneum gene (34, Table S1). The dsRNA construct for the 121 RNAi control was produced in our lab via cloning followed by PCR and *in vitro* transcription using 122 the T7 MEGAscript Kit (Ambion by Life TechnologiesTM GmbH, Darmstadt, Gemany) ³⁸. The 123 Dnmt2 dsRNA construct has been previously used in the ibeetle RNAi scan (33; http://ibeetle-base.uni-124 goettingen.de/details/TC005511) and was obtained from EupheriaBiotech (Dresden, Germany). Injections were carried out with a microliter injector (FemtoJet, Eppendorf AG, Hamburg, Germany) 125 126 and borosilicate glass capillaries (100 mm length, 1.0 mm outside diameter, 0.021 mm wall thickness; 127 Hilgenberg GmbH, Malsfeld, Deutschland) using dsRNA at a concentration of 1000 ng/µl dissolved 128 in ultrapure water. We injected pupae between the second and third lowest segment of their abdomen.

Over the three experimental blocks a total of 583 pupae were injected with *Dnmt2* dsRNA and 585 pupae served as RNAi control and were therefore injected with *asnA* dsRNA. Eclosion and survival of the procedure were recorded daily from three to six days post injection.

132 2.3.2 TGIP

When all surviving males from the RNAi treatment had reached sexual maturity seven days after
eclosion, they were injected for priming with around 37,000 cells of heat-killed *B. thuringiensis* (DSM
no. 2046, obtained from the German Collection of Microorganisms and Cell Cultures, DSMZ)

suspended in phosphate buffered saline (PBS). This treatment has successfully been used in prior TGIP
experiments ^{29,30}. Bacterial cultures were grown overnight as previously described ⁴¹. They were
washed with PBS and heat-killed by exposure to 95°C for 30 minutes. Control groups were either
injected with PBS (injection control) containing no bacterial cells or were left naïve. Injections were
performed using the nanolitre injector Nanoject II (Drummond Scientific Company, Broomall, PA,
USA) and individuals were injected between head and thorax. Beetles were kept individually before
and after the injections. Survival of the priming procedure was recorded 24h later.

143 2.3.3 Gene expression after RNAi and priming treatment

144 Twenty-four hours post priming, a subgroup of males was used for gene expression analysis to confirm 145 the knockdown of *Dnmt2*. In addition to the expression of *Dnmt2*, the expression of three immunity 146 and stress-related genes (hsp83, nimB and PGRP; Table S1) was analyzed, based on a previous study showing effects of paternal priming on gene expression ³⁰. For each RNAi*priming treatment 147 148 combination and block five samples were taken consisting of a pool of two to five individuals. RNA 149 extraction, cDNA reverse transcription and RT qPCR were performed as described above (2). Finally, 150 we also analyzed the expression of seven transposable elements (TEs) (Table S1), because the absence of *Dnmt2* can cause the activation of TEs²¹. Because of the lack of polyadenylation of some TE 151 152 transcripts, we used random hexamer primers for cDNA reverse transcription in this case (Thermo 153 Fisher Scientific, Waltham, MA USA).

154 2.3.4 Production and development of offspring generation

One day after the priming procedure, single pair matings were carried out for 24h with virgin females from the same population (n=12-50 mating pairs per treatment combination and experimental replicate). Twelve days after the oviposition to produce the F1 offspring generation, larvae from each pair were counted and up to six individuals were individualized and kept for further analyses. Additionally, one larva from each mating pair that produced offspring was used for developmental checks until it died or eclosed as an adult. The development was monitored daily from 21 to 23 dpo to check for pupation and at 26 dpo we recorded the proportion of eclosed adults. 162 2.3.5 Gene expression in the offspring generation

One week after the majority of the offspring generation had eclosed, five pools per RNAi and priming treatment combination and experimental replicate were sampled for gene expression analysis. Each sample consisted of five adult beetles of unknown sex. To avoid pseudo-replication only one beetle per family was used. Again, the expression of *Dnmt2* and three potential TGIP marker genes (*hsp83*, *nimB*, *PGRP*; 26) was analyzed as described above (3).

168 2.3.6 Bacterial challenge of adult offspring

169 One week after their eclosion, adults of the F1 offspring generation were submitted to a potentially 170 lethal bacterial injection (challenge). For this, bacteria from the same B. thuringiensis stock as for the priming were used. Again, an overnight culture from a glycerol stock was grown in liquid medium and 171 172 washed in PBS. The injection procedure was similar to priming and again included an injection control and a naïve group. The dose was adjusted to around 370 live bacterial cells per animal. From every 173 174 family one sibling each was used for the treatment and the controls. Again, beetles were kept 175 individually before and after injection to avoid any cross contaminations. Survival of the challenge 176 was recorded one day and four days post injection.

177 2.4 Statistics

All gene expression data were analyzed with the REST2009 software ⁴² as described in Eggert *et al.*³⁰. All other analyses were performed in RStudio version 0.99.467 ⁴³ under R version 3.3.3 ⁴⁴ using
additional packages lme4 ⁴⁵ and MASS ⁴⁶.

Survival of injections for RNAi and priming in the parental generation, the fertility of the treated males as well as the development of the offspring (proportion of pupae 21-23 dpo and proportion of adults 26 dpo) and their survival after bacterial challenge were analyzed in generalized linear mixed effect models (GLMMs) with the according error distributions and experimental replicate as a random factor.

185 **3 Results**

186 *3.1* Expression of *Dnmt2*

187 Before investigating a possible function of *Dnmt2* in *T. castaneum*, we monitored expression of this 188 gene throughout the life cycle of the beetle (i.e. in eggs, larvae, pupae and adults). The levels of 189 Dnmt2 transcripts (relative to two housekeeping genes) in eggs and pupae closely resembled those in 190 adults (eggs: relative expression=0.932, n=4, p=0.76; pupae: relative expression=0.989, n=8, p=0.94; 191 Figure 2a). Larvae expressed detectable amounts of *Dnmt2*, but significantly less than adults (relative 192 expression=0.352, n=7, p<0.001; Figure 2a). Additionally, *Dnmt2* appears to serve functions in both 193 males and females as its expression did not differ significantly between the sexes for pupae (female: 194 relative expression=0.784, n=4, p=0.23) or adults (female: relative expression=0.709, n=4, p=0.14). 195 Furthermore, we analyzed the expression of *Dnmt2* in the reproductive tissue of the male beetles and 196 compared it to whole body samples of the same sex, because expression in the testes could hint to a 197 possible relevance of the protein in male reproduction or even an involvement in the transfer of 198 information from father to offspring as possibly needed for TGIP. Dnmt2 mRNA levels in the testes 199 were significantly higher than in whole-body samples (relative expression=2.497, n=6, p=0.001; 200 Figure 2b).

201 3.2 Paternal *Dnmt2* knockdown and TGIP

To determine whether *Dnmt2* is involved in the paternal transfer of immunity, we combined a
knockdown with paternal TGIP treatment and exposed the offspring to a bacterial challenge (Figure
1).

205 3.2.1 Survival of RNAi and priming injections

The RNAi treatment with *Dnmt2* dsRNA did not increase mortality of the treated pupae (Figure S1). Injections of dsRNA in male pupae did not significantly alter survival rates neither directly following the RNAi treatment (GLMM, df=1, X^2 =0.04, p=0.84, Figure S1a) nor after the priming treatment in the mature adults ten days later (GLMM, df=1, X^2 =0.16, p=0.69; Figure S1b). The priming procedure itself led to a significantly increased mortality. However, the effect size was small and independent of whether the beetles were injected with heat killed bacteria or the PBS treatment control, which can be attributed to the wounding during these injections as none of the naïve individuals died (GLMM, df=2,
X²=15.89, p<0.001; Figure S1b).

214 3.2.2 Knockdown of Dnmt2

One day after the priming procedure, we confirmed the knockdown of *Dnmt2* after pupal RNAi in a subgroup of the adults. *Dnmt2* was significantly downregulated compared to RNAi control regardless of the received priming treatment (Table 1). As expected, *Dnmt2* mRNA levels had returned to normal in the adult offspring and there were no significant differences between the RNAi treatments detectable (Table 1). Additionally, the paternal priming procedure did not affect *Dnmt2* expression in the adult offspring (Table 1).

221 3.2.3 Knockdown of *Dnmt2* and adult priming did not affect male fertility

Neither the knockdown of *Dnmt2* nor the bacterial priming appear to affect the fitness of the treated individuals, as neither treatment significantly altered male fertility. The number of live offspring obtained from a 24 h single pair mating period did not differ significantly for either of the treatments (GLMM: RNAi, df=1, X^2 =2.11, p=0.15; priming, df=2, X^2 =0.44, p=0.80; Figure S2).

226 3.2.4 Paternal knockdown of *Dnmt2* but not priming affected offspring development

227 We monitored offspring development by measuring the proportion of pupae over three consecutive 228 days and the proportion of eclosed adults 26 days post oviposition (dpo). Animals from all six 229 treatment combinations of RNAi and priming showed similar pupation rates 21 and 22 dpo (Figure 3a, 230 Figure S3a). However, at 23 dpo significantly less larvae had reached pupation in the Dnmt2 paternal 231 knockdown group than in the RNAi control, independent of paternal priming treatment (GLMM: RNAi, df=1, X^2 =3.9, p<0.05; priming, df=2, X^2 =0.19, p=0.91; Figure 3a). The proportion of eclosed 232 233 adults 26 dpo was not significantly affected by any paternal treatment (GLMM: RNAi, df=1, X^2 =0.04, 234 p=0.84; priming, df=2, X²=0.48, p=0.79; Figure S3b).

235 3.2.5 Expression of TGIP marker genes and TEs is not affected by *Dnmt2* knockdown or priming

In fathers and offspring alike, we measured the expression of three genes, which are related to stress or immune responses and were previously shown to be upregulated in the adult offspring of primed fathers (26). By measuring the expression in the fathers, we intended to see whether these genes would already be affected within the treated generation. None of the three candidate genes (hsp83, nimB and PGRP) showed any significant differential expression neither in the paternal nor in the adult offspring generation (Table S2).

For the same animals from the paternal generation we also measured the expression of seven TEs. Genenncher *et al.* ²¹ observed that the absence of *Dnmt2* and the exposure to heat stress led to the activation and accumulation of certain TEs in *D. melanogaster*. Here, we could not observe any significant upregulation in the expression of TEs after wounding or priming in the knockdown or control treatment (Table S3).

247 3.2.6 Paternal Dnmt2 knockdown reduces offspring survival after bacterial challenge

248 Finally, we injected adult beetles from the offspring generation with a potentially lethal dose of B. thuringiensis to see whether the immune priming was transmitted to the offspring and if this was 249 250 affected by the downregulation of *Dnmt2* in the fathers. Paternal priming treatment did not affect offspring survival after bacterial challenge (GLMM, df=2, X²= 0.17, p=0.92; Figure S4), which can 251 252 possibly be explained by the unavoidable additional wounding that all fathers were subjected to because of the RNAi injection treatment. However, offspring of individuals that had received a 253 254 knockdown were significantly less likely to survive the bacterial challenge (GLMM, df=2, X²=7.78, 255 p<0.01; Figure 3), demonstrating that *Dnmt2* is involved in paternal effects and that its reduction can 256 increase the offspring's susceptibility to pathogens in the beetle.

257 4 Discussion

258 *Dnmt2* can be found in almost every eukaryote species and is the most conserved member of the Dnmt 259 family ⁴⁷. It also has a function in some organisms lacking one or both other Dnmts and which are 260 often devoid of any functional DNA methylation system ¹³. This also appears to be the case in 261 *T. castaneum*, which lacks *Dnmt3* and does not have any functional CpG DNA methylation ^{4,33–35}, but 262 still expresses Dnmt2. Although, we know today that Dnmt2 methylates tRNA and not DNA, it remains unclear what the role of this epigenetic mechanism is. We observed that Dnmt2 mRNA transcripts are 263 264 present at relatively low but consistent levels in all life stages and in both sexes of the beetle, therefore 265 the enzyme likely functions in both males and females and throughout the entire life cycle. In fruit flies, mice and humans, *Dnmt2* canonically methylates a small set of tRNAs ^{14,15,48}, which are highly 266 abundant in sperm ⁴⁹ and have been shown to be involved in paternal transmission of metabolic 267 phenotypes in mice ^{17,18}. The significantly higher expression of *Dnmt2* that we observed in testes of 268 269 T. castaneum could hint at the involvement of Dnmt2 in paternal epigenetic inheritance also in this 270 beetle.

271 We used systemic paternal RNAi, to investigate how a Dnmt2 knockdown would affect offspring phenotypes and to discover a potential role of this gene in intergenerational epigenetic inheritance, 272 273 specifically from father to offspring. The offspring of Dnmt2 RNAi-treated fathers needed longer to 274 reach pupation and were less well dealing with a *B. thuringiensis* infection, which may point towards a generally higher stress sensitivity. In recent years, it has become clear that biological functions of 275 Dnmt2 are more easily detected under stress conditions ¹³. Increased sensitivity to thermal and 276 277 oxidative stress has been observed in D. melanogaster Dnmt2 mutants ¹⁵, while overexpression of the same gene led to increased stress tolerance 50. During the stress response, *Dnmt2* appears to control 278 the fragmentation of tRNA and can be located at cellular stress compartments ^{15,51}. More recently, it 279 280 has also been demonstrated that the knockout of Dnmt2 leads to a decline of immune function with ageing in adult flies ⁵². Finally, its absence disrupts the small interfering RNA pathway by inhibiting 281 dsRNA degradation by *Dicer*⁵¹. However, all these effects occur in the treated generation and have 282 283 not been tested for an offspring generation. In our case the developmental and immunological effects 284 were observed in the offspring of treated fathers, even though offspring themselves exhibited normal 285 Dnmt2 expression, which suggests transgenerational effects. A role in transgenerational paternal 286 effects for *Dnmt2* has so far only been established in mice, where the gene is essential for the 287 transmission of an acquired metabolic disorder ¹⁹. Therefore, further studies are needed to clarify the 288 role that *Dnmt2* plays in enabling appropriate stress responses within and across generations.

289 Finally, we combined the knockdown of *Dnmt2* with a TGIP treatment, to determine whether this 290 enzyme is involved in the transfer of priming information from father to offspring. We did not observe 291 any TGIP in the present study, as offspring survival was independent of the paternal priming treatment. 292 Furthermore, we did not observe any upregulation in the previously described marker genes for immune and stress responses ³⁰ or any changes in the expression of transposable elements. This 293 294 surprising lack of any observable difference between offspring of primed and control males might have 295 been caused by the injection of all males for the pupal RNAi treatment that took place in the pupal 296 stage, before the priming treatment in the adult stage (note that we included an RNAi control, but did 297 not have a fully naïve control for the RNAi treatment, Fig. 1). The wounding associated with the 298 dsRNA injection might have activated immune responses, such that all animals could already have 299 entered a 'primed' state. To our knowledge there are no studies that directly address the question how 300 injuries during the pupal phase influence later immune responses. However, some experiments show 301 that wounding in control treatments also to a certain extent may increase survival of a subsequent bacterial challenge ^{29,53}. Alternatively, the pupal RNAi injections might have inhibited priming. Lastly, 302 303 although TGIP in *T. castaneum* is robust and repeatable ^{29,30,53}, it also has become apparent that this phenomenon cannot be observed in every experiment ⁵³ and beetle population ⁵⁴. 304

We did not observe any effects of the Dnmt2 knockdown on expression of the studied TEs, in contrast to *D. melanogaster Dnmt2* mutants that showed increased TE expression 21 . However, further studies would be needed to make any firm conclusions, because the lack of expression differences for a limited set of TEs does not exclude the possibility that *Dnmt2* plays a role in the regulation of TEs in *T. castaneum*.

In plants, flies and mice the absence of *Dnmt2* is not lethal under standard conditions and mutants remain fertile ¹⁴. The same appears to be true in the case of *T. castaneum*, where we did not observe any mortality nor apparent phenotypic changes after a significant downregulation of *Dnmt2*. Additionally, male fertility was not affected by the knockdown under *ad libitum* conditions. Therefore, the maintenance of knockout lines appears feasible, which makes this gene a suitable target for CRISPR/Cas knockout to further study its function without the necessity of repeated RNAi injections for each experiment. Moreover, methylation-sensitive sequencing of sperm RNA could furtherelucidate the underlying molecular processes.

In conclusion, our study for the first time describes paternal effects of *Dnmt2* in an invertebrate. The here observed prolonged development and increased susceptibility to infection in the offspring occurred in the presence of normal *Dnmt2* expression in the offspring themselves. Therefore, tRNA methylation in sperm provides a fascinating possibility of transmitting information from fathers to offspring through paternal epigenetic inheritance.

- 323
- 324 References
- Feinberg, A. P. Phenotypic plasticity and the epigenetics of human disease. *Nature* 447, 433–
 440 (2007).
- Duncan, E. J., Gluckman, P. D. & Dearden, P. K. Epigenetics, plasticity, and evolution: How
 do we link epigenetic change to phenotype? *J. Exp. Zool. Part B Mol. Dev. Evol.* 322, 208–
 220 (2014).
- 330 3. Lyko, F. The DNA methyltransferase family: a versatile toolkit for epigenetic regulation. *Nat.* 331 *Rev. Genet.* 19, 81–92 (2018).
- Bewick, A. J., Vogel, K. J., Moore, A. J. & Schmitz, R. J. Evolution of DNA methylation
 across insects. *Mol. Biol. Evol.* 34, 654–665 (2017).
- 334 5. Provataris, P. *et al.* Signatures of DNA methylation across insects suggest reduced DNA
 335 methylation levels in Holometabola. *Genome Biol. Evol.* 10, 1185–1197 (2018).
- Ernst, U. R. *et al.* Epigenetics and locust life phase transitions. *J. Exp. Biol.* 218, 88–99
 (2015).
- Pasquier, C. *et al.* Environmentally selected aphid variants in clonality context display
 differential patterns of methylation in the genome. *PLoS One* 9, e115022 (2014).
- Falckenhayn, C. *et al.* Characterization of genome methylation patterns in the desert locust
 Schistocerca gregaria. J. Exp. Biol. 216, 1423–9 (2013).
- 342 9. Elango, N., Hunt, B. G., Goodisman, M. a D. & Yi, S. V. DNA methylation is widespread

| 343 | and associated | with differential | gene expression in | n castes of the hone | ybee, Apis mellifera. |
|-----|----------------|-------------------|--------------------|----------------------|-----------------------|
| | | | | | |

- 344 *Proc. Natl. Acad. Sci. U. S. A.* **106**, 11206–11 (2009).
- Kucharski, R., Maleszka, J., Foret, S. & Maleszka, R. Nutritional control of reproductive
 status in honeybees via DNA methylation. *Science*. 319, 1827–1830 (2008).
- Rana, A. K. & Ankri, S. Reviving the RNA world: An insight into the appearance of RNA
 methyltransferases. *Frontiers in Genetics* 7, 99 (2016).
- 349 12. Goll, M. G. & Bestor, T. H. Eukaryotic cytosine methyltransferases. *Annu. Rev. Biochem.* 74, 481–514 (2005).
- 13. Durdevic, Z. & Schaefer, M. *Dnmt2* methyltransferases and immunity: An ancient
- 352 overlooked connection between nucleotide modification and host defense? *BioEssays* 35,

353 1044–1049 (2013).

- 354 14. Goll, M. G. *et al.* Methylation of tRNA Asp by the DNA methyltransferase homolog Dnmt2.
 355 *Science* **311**, 395–398 (2006).
- 356 15. Schaefer, M. *et al.* RNA methylation by *Dnmt2* protects transfer RNAs against stress-induced
 357 cleavage. *Genes Dev.* 24, 1590–1595 (2010).
- 16. Raddatz, G. *et al.* Dnmt2-dependent methylomes lack defined DNA methylation patterns.
- **359** *Proc. Natl. Acad. Sci.* **110**, 8627–31 (2013).
- 360 17. Chen, Q. *et al.* Sperm tsRNAs contribute to intergenerational inheritance of an acquired
 361 metabolic disorder. *Science.* 351, 397–400 (2016).
- 362 18. Sharma, U. *et al.* Biogenesis and function of tRNA fragments during sperm maturation and
 363 fertilization in mammals. *Science*. 351, 391–396 (2016).
- 364 19. Zhang, Y. et al. Dnmt2 mediates intergenerational transmission of paternally acquired
- 365 metabolic disorders through sperm small non-coding RNAs. *Nat. Cell Biol.* 20, 535–540
 366 (2018).
- 20. Zhang, Y., Shi, J., Rassoulzadegan, M., Tuorto, F. & Chen, Q. Sperm RNA code programmes
 the metabolic health of offspring. *Nat. Rev. Endocrinol.* 15, 489–498 (2019).
- 369 21. Genenncher, B. et al. Mutations in Cytosine-5 tRNA methyltransferases impact mobile
- element expression and genome stability at specific DNA repeats. *Cell Rep.* 22, 1861–1874

| 371 | | (2018). |
|-----|-----|--|
| 372 | 22. | Phalke, S. et al. Retrotransposon silencing and telomere integrity in somatic cells of |
| 373 | | Drosophila depends on the cytosine-5 methyltransferase DNMT2. Nat. Genet. 41, 696–702 |
| 374 | | (2009). |
| 375 | 23. | Durdevic, Z. et al. Efficient RNA virus control in Drosophila requires the RNA |
| 376 | | methyltransferase Dnmt2. EMBO Rep. 14, 269–75 (2013). |
| 377 | 24. | Schmid-Hempel, P. Natural insect host-parasite systems show immune priming and |
| 378 | | specificity: puzzles to be solved. BioEssays 27, 1026–1034 (2005). |
| 379 | 25. | Milutinović, B. & Kurtz, J. Immune memory in invertebrates. Semin. Immunol. 28, 328-342 |
| 380 | | (2016). |
| 381 | 26. | Roth, O., Beemelmanns, A., Barribeau, S. M. & Sadd, B. M. Recent advances in vertebrate |
| 382 | | and invertebrate transgenerational immunity in the light of ecology and evolution. Heredity. |
| 383 | | 121 , 225–238 (2018). |
| 384 | 27. | Kurtz, J. & Armitage, S. A. O. Dissecting the dynamics of trans-generational immune |
| 385 | | priming. Mol. Ecol. 26, 3857–3859 (2017). |
| 386 | 28. | Zanchi, C., Troussard, JP., Moreau, J. & Moret, Y. Relationship between maternal transfer |
| 387 | | of immunity and mother fecundity in an insect. Proc. R. Soc. B Biol. Sci. 279, 3223–3230 |
| 388 | | (2012). |
| 389 | 29. | Roth, O. et al. Paternally derived immune priming for offspring in the red flour beetle, |
| 390 | | Tribolium castaneum. J. Anim. Ecol. 79, 403–413 (2010). |
| 391 | 30. | Eggert, H., Kurtz, J. & Diddens-de Buhr, M. F. Different effects of paternal trans- |
| 392 | | generational immune priming on survival and immunity in step and genetic offspring. Proc. |
| 393 | | <i>R. Soc. B Biol. Sci.</i> 281 , (2014). |
| 394 | 31. | Schulz, N. K. E., Sell, M. P., Ferro, K., Kleinhölting, N. & Kurtz, J. Transgenerational |
| 395 | | developmental effects of immune priming in the red flour beetle Tribolium castaneum. Front. |
| 396 | | Physiol. 10, (2019). |
| 397 | 32. | Castro-Vargas, C. et al. Methylation on RNA: A potential mechanism related to immune |
| 398 | | priming within but not across generations. Front. Microbiol. 8, 1–11 (2017). |

- 399 33. Richards, S. *et al.* The genome of the model beetle and pest *Tribolium castaneum*. *Nature*400 452, 949–55 (2008).
- 401 34. Zemach, A., Mcdaniel, I. E., Silva, P. & Zilberman, D. Genome-wide evolutionary analysis
 402 of eukaryotic DNA methylation. *Science* 328, 916–9 (2010).
- 403 35. Schulz, N. K. E. et al. Dnmt1 has an essential function despite the absence of CpG DNA
- 404 methylation in the red flour beetle *Tribolium castaneum*. Sci. Rep. **8**, 16462 (2018).
- 405 36. Bucher, G., Scholten, J. & Klingler, M. Parental RNAi in *Tribolium* (Coleoptera). *Curr. Biol.*406 12, R85-6 (2002).
- 407 37. Schmitt-Engel, C. *et al.* The iBeetle large-scale RNAi screen reveals gene functions for insect
 408 development and physiology. *Nat. Commun.* 6, 7822 (2015).
- 409 38. Peuß, R. et al. Down syndrome cell adhesion molecule 1 : testing for a role in insect
- 410 immunity, behaviour and reproduction. *R. Soc. Open Sci.* **3**, 160138 (2016).
- Gilles, A. F. & Averof, M. Functional genetics for all: Engineered nucleases, CRISPR and the
 gene editing revolution. *Evodevo* 5, 1–13 (2014).
- 413 40. Milutinović, B., Stolpe, C., Peuß, R., Armitage, S. A. O. & Kurtz, J. The red flour beetle as a
 414 model for bacterial oral infections. *PLoS One* 8, (2013).
- 415 41. Roth, O. & Kurtz, J. Phagocytosis mediates specificity in the immune defence of an
- 416 invertebrate, the woodlouse *Porcellio scaber* (Crustacea: Isopoda). *Dev. Comp. Immunol.* 33,
 417 1151–5 (2009).
- 418 42. Pfaffl, M. W., Horgan, G. W. & Dempfle, L. Relative expression software tool (REST) for
- group-wise comparison and statistical analysis of relative expression results in real-time PCR. *Nucleic Acids Res.* 30, e36 (2002).
- 421 43. RStudio Team. R-Studio: integrated development for R. (2015).
- 422 44. R Development Core Team. R: A language and environment for statistical computing.423 (2008).
- 424 45. Bates, D., Mächler, M., Bolker, B. & Walker, S. Fitting Linear Mixed-Effects Models Using
 425 lme4. *J. Stat. Softw.* 67, 1–48 (2015).
- 426 46. Venables, W. N. & Ripley, B. D. Modern Applied Statistics with S. (Springer New York,

| 427 | 2002). doi:10.1007/978-0-387-21706-2 |
|-----|--------------------------------------|
|-----|--------------------------------------|

- 428 47. Schaefer, M. & Lyko, F. Solving the *Dnmt2* enigma. *Chromosoma* **119**, 35–40 (2010).
- 429 48. Jurkowski, T. P. *et al.* Human DNMT2 methylates tRNA(Asp) molecules using a DNA

430 methyltransferase-like catalytic mechanism. *RNA* 14, 1663–70 (2008).

- 431 49. Peng, H. *et al.* A novel class of tRNA-derived small RNAs extremely enriched in mature
 432 mouse sperm. *Cell Res.* 22, 1609–12 (2012).
- 433 50. Lin, M.-J., Tang, L.-Y., Reddy, M. N. & Shen, C.-K. J. DNA methyltransferase gene *dDnmt2*434 and longevity of *Drosophila*. *J. Biol. Chem.* 280, 861–4 (2005).
- 435 51. Durdevic, Z., Mobin, M. B., Hanna, K., Lyko, F. & Schaefer, M. The RNA methyltransferase

Dnmt2 is required for efficient Dicer-2-dependent siRNA pathway activity in Drosophila. *Cell Rep.* 4, 931–937 (2013).

- 438 52. Abhyankar, V., Kaduskar, B., Kamat, S. S., Deobagkar, D. & Ratnaparkhi, G. S. Drosophila
- 439 DNA/RNA methyltransferase contributes to robust host defense in ageing animals by
- regulating sphingolipid metabolism. J. Exp. Biol. jeb.187989 (2018). doi:10.1242/jeb.187989
- 441 53. Tate, A. T., Andolfatto, P., Demuth, J. P. & Graham, A. L. The within-host dynamics of
- infection in trans-generationally primed flour beetles. *Mol. Ecol.* **26**, 3794–3807 (2017).
- Khan, I., Prakash, A. & Agashe, D. Divergent immune priming responses across flour beetle
 life stages and populations. *Ecol. Evol.* 6, 7847–7855 (2016).

445

446 Contributions

All authors conceived and designed the experiments. NS conducted the experiments, analyzed the data,and wrote the manuscripts with comments from all authors.

449 Acknowledgements

We thank Barbara Hasert and Kathrin Brüggemann for help with lab work, Jürgen Schmitz for support
regarding the expression analysis of TEs and Sina Flügge for providing us with drawings of *T. castaneum*. This work was supported in part by the Volkswagen Stiftung, project number I/84 794.

453 Tables and Figures

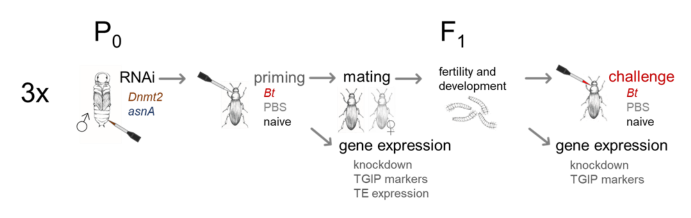
Table 1 Dnmt2 expression after paternal RNAi and priming in the treated males and their adult offspring. Given
 is the relative expression compared to RNAi control, per priming control group, for the knockdown target gene
 Dnmt2 normalized over the expression of two housekeeping genes. Per treatment combination, generation

436 *Dimite* normalized over the expression of two housekeeping genes. Per treatment comb 457 and three experimental replicates five samples comprising of 2-5 individuals were used.

| Gene | Treatment | | Р | | | F ₁ | | |
|-------|-----------|-----------|-----------------|-------------|---------|-----------------|-------------|---------|
| | RNAi | Priming | rel. expression | 95% C.I. | p value | rel. expression | 95% C.I. | p value |
| Dnmt2 | Dnmt2 | bacterial | 0.177 | 0.03 - 1.73 | <0.001 | 1.116 | 0.54 - 2.04 | 0.284 |
| | | control | 0.088 | 0.04 - 0.48 | <0.001 | 1.061 | 0.53 - 1.97 | 0.530 |
| | | naive | 0.112 | 0.03 - 0.73 | <0.001 | 0.955 | 0.5 - 1.76 | 0.620 |
| | control | bacterial | 0.897 | 0.27 - 5.90 | 0.571 | 1.06 | 0.52 - 1.94 | 0.519 |
| | | naive | 1.175 | 0.26 - 7.21 | 0.384 | 1.092 | 0.53 - 3.63 | 0.480 |

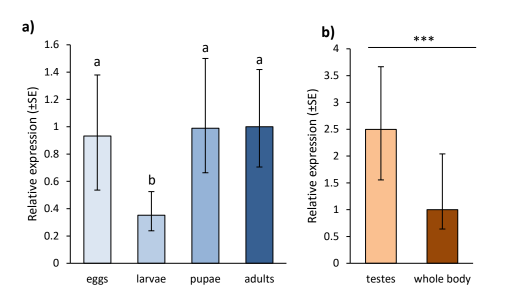
458

459



- 460 Figure 1 Overview of the experimental design combining paternal RNAi to knockdown Dnmt2 with paternal
- 461 TGIP. The entire experiment was conducted in three replicates (asnA=RNAi control; PBS=injection control;
- 462 naïve=handling control).

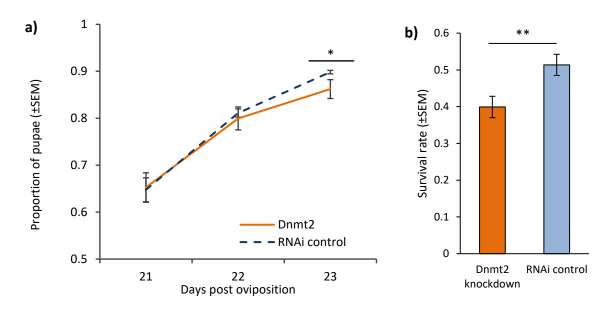




464

Figure 2 Dnmt2 gene expression a) Relative expression in four distinct life stages of *T. castaneum* compared
to adult samples (eggs: n=4 pools of 500-1000 eggs, 24h-48h post oviposition; larvae: n=7 pools of 10 larvae,
14-19 dpo; pupae: n=8 pools of 6 individuals; adults: n=8 pools of 6 individuals, one week after eclosion). b)
Relative expression in testes and male whole-body samples (n=6). Different lettering and asterisks indicate
significant differences (p≤0.001).





471

Figure 3 Effects of paternal *Dnmt2* knock-down on offspring **a**) Pupation rate of F_1 generation 21-23 dpo (±SEM for three experimental replicates) **b**) Survival of the F_1 generation after bacterial challenge according to paternal RNAi treatment. Shown are the proportions of adults that were alive four days post injection with a of *B. thuringiensis* (±SEM for three experimental replicates). Asterisks indicate significant differences (*=p<0.05, **=p<0.01).