

ORIGINAL ARTICLE

Quantitative genetic variation in the control of ovarian apoptosis under different environments

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Fertility loss in otherwise healthy individuals can be an evolutionary conundrum. Most studies on the evolution of post-reproductive lifespan focus on the fitness effects of survival past the age of last reproduction. A complementary approach, which has been largely neglected, is to develop an understanding of the nature of variation in the mechanism underlying loss of fertility, ovarian apoptosis. Variation in the genetics underlying the regulation of ovarian apoptosis could hold the key to understanding the evolution of midlife fertility loss. We estimated quantitative genetic variation in the regulation of ovarian apoptosis in females of the cockroach *Nauphoeta cinerea*, an insect with reproductive cycles. We have earlier shown that delaying reproduction incites loss of fertility. Here, we forced

females to delay reproduction under conditions of excess or limited food and examined apoptosis under both conditions. We found substantial additive genetic variation in levels of apoptosis when females experienced a limited period of starvation during sexual maturation but not when females had unlimited access to food. Hence, selection could act on the regulation of ovarian apoptosis to change the rate of fertility loss with age at least under some environmental circumstances. Our results suggest that an understanding of how loss of fertility evolves requires an understanding of the interaction between genes involved in the regulation of apoptosis and environmental factors such as diet. *Heredity* (2009) **103**, 217–222; doi:10.1038/hdy.2009.44; published online 29 April 2009

Keywords: loss of fertility; oocyte apoptosis; quantitative genetics

Introduction

Female fertility declines with age. In many species this loss of fertility is associated with senescence (Partridge and Fowler, 1992; Kirkwood, 2002). In some species, however, females lose fertility relatively early in their lifespan rather than late in life. This midlife fertility loss can present an evolutionary puzzle; why should otherwise healthy individuals lose reproductive capacity? One possibility is that females have an extended post-reproductive lifespan, and contribute to fitness in ways other than personal reproduction. Another possibility is oosorption, the reallocation of reproductive resources to somatic maintenance under conditions of environmental stress. Studies addressing the evolution of loss of fertility have focused on the selective pressures that may be acting on the cessation of reproduction before senescence (Peccei, 2001; Reznick *et al.*, 2006) or on oosorption (Boggs 2009). A complementary approach, which has been largely ignored so far, is developing an understanding of the nature of variation in the target of selection and how this may contribute to its evolution. Here we examine the genetic variation underlying one such target, the control of oocyte apoptosis, the process by which females with a healthy somatic body lose fertility.

Ovarian apoptosis has been identified as a universal mechanism of loss of fertility in species ranging from humans (Morita and Tilly, 1999) to insects (Terashima and Bownes, 2004; Moore and Sharma, 2005). Precise control of this evolutionarily conserved mechanism of programmed cell death is essential to the normal functioning of all tissues in all multicellular organisms (Vaux and Strasser, 1996; Reed, 2000; Adams, 2003). Imprecise regulation has disastrous consequences including aberrant embryogenesis, autoimmunity, early loss of fertility and cancer. Thus, we might expect genetic variation in the regulation of apoptosis to be low, as it should be under strong direct selection. However, the decline in midlife fertility is variable (Reznick *et al.*, 2006; Broekmans *et al.*, 2007; Moore *et al.*, 2007) and has the potential to evolve (Moore *et al.*, 2007). This variation may also depend on the specific environmental conditions for an organism. In guppies, for example, the presence of predators results in a delay in the decline in reproductive potential compared with females in populations where predators are absent (Reznick *et al.*, 2006). In humans, phenotypic variation in age at natural menopause also appears to be affected by factors such as smoking and diet (Nagata *et al.*, 2000; Gold *et al.*, 2001; de Bruin *et al.*, 2001; Kok *et al.*, 2005). The variation underlying rates of fertility loss are not only environmental. The age of onset of natural menopause in humans can be predicted by family history, with heritability estimates ranging from 0.30 to 0.85 (Kirk *et al.*, 2001; de Bruin *et al.*, 2001; Kok *et al.*, 2005; Pettay *et al.*, 2005; Broekmans *et al.*, 2007). Linkage analysis of age at natural menopause has identified two significant quantitative trait loci influencing age at menopause

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Received 11 September 2008; revised 29 January 2009; accepted 19 March 2009; published online 29 April 2009

(van Asselt *et al.*, 2004). Thus, genetic variation does exist in populations in the age at which females lose fertility. The variation in age of onset of post-reproductive lifespan suggests that there must be variation in the loss of oocytes through apoptosis. However, it is unclear where variation might occur in such a highly conserved biochemical pathway.

Age is not the only factor that influences loss of fertility. We expect animals to have evolved mechanisms by which energy saved by a reduction in reproduction can be used to increase survivorship (Bell and Bohm 1975). Oosorption, the ability to resorb eggs, is thought to be an adaptive mechanism by which nutrients can be reallocated into survival when opportunities for reproduction are limited (Bell and Bohm 1975; Papaj 2000; Boggs, 2009). For example, in butterflies a reduction in food leads to a reduction in fecundity, an increase in oocyte resorption, but no reduction in lifespan (Boggs and Ross 1993). This suggests a trade-off between reproduction and lifespan mediated by the recycling of nutrients invested in oocytes that cannot be used. Thus, alleles that may be beneficial in one context, reducing oocyte death and extending relative reproductive lifespan, could be detrimental under other contexts, limiting the ability of individuals to adjust to resource limitation. However, the underlying physiological and genetic control of this trade-off has not been explored. Insight into patterns of resource allocation in varied environments is essential to developing a predictive theory of life history (Boggs 2009), including the relationship between early reproductive history and later life fecundity.

We hypothesize that variation in oocyte and fertility loss arises through variation in the control of apoptosis, through both environmental and genetic factors, rather than in the cell death machinery itself. Thus, a quantitative genetic approach should be informative (Roff, 2007). We tested this hypothesis in the cockroach *Nauphoeta cinerea*. In *N. cinerea* we can manipulate the rate of fertility loss through both age at first reproduction (Moore and Moore, 2001) and diet (Barrett *et al.*, 2007). *N. cinerea* is ovoviviparous: females carry the embryos in a brood pouch until the offspring are born as first-instar nymphs (Roth, 1964a, b). This species differs from most other invertebrate laboratory systems in that it shows reproductive cycles, with bouts of oocyte maturation followed by pregnancy. Females lose reproductive potential through loss of viable oocytes through apoptosis (Moore and Sharma, 2005) and there is additive genetic variation in fecundity following mating late relative to sexual maturation (Moore *et al.*, 2007).

We investigated the quantitative genetic variation in the levels of ovarian apoptosis in female *N. cinerea* that were forced to delay reproduction by preventing access to males under either *ad libitum* food or food-restricted conditions by undertaking a paternal half-sib breeding design with broods split between fully fed and food-restricted conditions during the period between adult eclosion and mating. These treatments were used because diet has been shown to alter rates of apoptosis in *N. cinerea* (Barrett *et al.*, 2007) and to affect ovarian apoptosis in *Drosophila melanogaster*; starvation triggers apoptosis in the ovaries resulting in fewer eggs being laid (Terashima and Bownes, 2004). In our study, we found substantial genetic variation in levels of ovarian

apoptosis under food-restricted conditions but not under *ad libitum* food.

Materials and methods

Animal husbandry

We obtained the males used as sires and the females used as dams in this study from eight out-bred mixed-sex, mass rearing colonies. The colonies have been established for over 50 years, contain several thousand individuals each, with enforced gene flow among them. The population is maintained under the standard rearing conditions of dry rat chow and water at 28 °C under a 12:12 h light/dark photoperiod. Assessment of allozyme variation has shown that the population is not inbred (Corley *et al.*, 2001). Last instar nymphs were selected at random from the mass colonies, sexed and placed in single sex colonies in 30 × 18 × 12-cm plastic boxes under the standard rearing conditions. When adults emerged they were isolated in individual 11 × 11 × 3-cm plastic containers with *ad libitum* rat chow and water until they were mated. Males were returned to their containers after each mating.

Breeding design and rearing of offspring

We staged all mating trials under red light in a constant temperature room, kept at 28 °C, during the dark phase of the 12:12 h light/dark photoperiod. Sires ($n = 52$) were mated sequentially to five virgin females each (a minimum of four were included in the analysis), with 3 days (72 ± 1 h) between matings. All males were virgin at the time of their first mating and had eclosed as adults 10 days earlier. Dams were also virgin and 10 days post-eclosion at the time of mating, the optimal time of mating for females (Moore and Moore, 2001).

We placed mated females in individual 17 × 12 × 6.5-cm plastic boxes under the standard rearing conditions. Females were removed from these boxes as soon as the first clutch of offspring appeared. We monitored the boxes and made sure that the developing offspring had access to *ad libitum* food and water.

Manipulation of experimental females

We isolated experimental females from each clutch by moving females from the family box into individual 11 × 11 × 3-cm plastic boxes on the day they emerged as adults. These females were either provided with *ad libitum* food and water (three daughters per dam) or were food-restricted, being provided with water only (three daughters per dam). Females were mated at 18 days. Females that mate at sexual maturity (6–9 days post-adult eclosion) have low levels of ovarian apoptosis and high fecundity, whereas females that mate between 9 and 18 days of age show increasing levels of ovarian apoptosis and decreasing lifetime fecundity (Moore and Moore 2001; Moore and Sharma 2005). The reduction in fecundity is seen in both the first clutch produced, but also all clutches produced over a female's lifespan of approximately 300 days.

Apoptosis trials

After 18 days, we dissected the experimental females and collected 10 mature oocytes from each female. We stained the oocytes for apoptosis using the nucleic acid dye

YO-PRO-1 and propidium iodide. To confirm that the YO-PRO-1 stain, which is fast and economical (thus allowing for the large numbers of sample necessary for a quantitative genetic study to be stained), represents apoptosis as opposed to other forms of programmed cell death, we stained the oocytes of 18 day post-eclosion females with the stain Casp 3/7 Mito Casp (Bachem, Torrance, CA, USA). This stain is a simultaneous assay for membrane potential and active caspase 3/7. Staining patterns for active caspase 3/7 are identical to those seen with the YO-PRO-1 stain in the ovaries of both *ad libitum* food and food-restricted 18-day-old females (Figure 1), indicating that YO-PRO-1 does reveal apoptosis in the oocytes under study here. We scored each oocyte depending on the proportion of this epithelium that was stained as follows: (1) <5% stained, (2) 5–40% stained, (3) 40–70% stained, (4) 70–100% stained, (5) 100% stained—the follicular epithelium was completely apoptotic. The score for each female was then calculated by adding the scores of all 10 oocytes.

Statistical methods

Our experimental design involved a half-sib/full-sib, split brood design with 52 sires and 217 dams. Full-sib families were reared together until the last instar stage, at which time females were separated into individual containers and assigned to either *ad libitum* food conditions or food-restricted condition. Each family contributed a minimum of two offspring into each environment. Thus, there were 635 individual female offspring assessed for the *ad libitum* environment and 645 individual offspring for the food-restricted environment.

Our first analysis examined the entire data set with a fully random effects analysis of anova model. Environment was considered a random effect because, similar to family, the specific level of food restriction or abundance was randomly selected from an (nearly) infinite range of possible levels of food. In such a model, the denominator degrees of freedom vary depending on the main effect examined. We used a SAS model III (Ayres and Thomas,

1990; Fry, 1992) where the denominator degrees of freedom for the test of significance for the environment treatment were the sire by environment mean squares; for the test of the effect of sire the denominator mean square was a combination of mean squares (the mean square term for the dam nested within sire + mean square of the sire by environment—mean square for dam nested within sire by environment with 37 degrees of freedom (d.f.)); the test for the dam nested within sire was the dam nested within sire by environment mean squares; the test of the sire by environment effect also used the dam nested within sire by environment means squares; the dam nested within sire by environment effect used the error mean squares in the denominator of the F-test.

Within environments, we calculated quantitative genetic parameters from our half-sib/full-sib design using restricted maximum likelihood method to generate variance estimates. The level of imbalance in our data was minor, with little difference between restricted maximum likelihood and least square estimates, and we therefore calculate significance from least-squares estimates of expected means squares in a nested analysis of variance (Lynch and Walsh, 1998).

Results

The levels of oocyte apoptosis differed under the two environmental conditions (Figure 2). Controlling for genetic influences by analyzing the data with a fully random effects analysis of variance model (with environment, sire, dam nested under sire and their interactions), there was a highly significant difference in the apoptosis scores in the *ad libitum* food environment compared with the food-restricted environment (Table 1). The average apoptosis score in the *ad libitum* environment was 12.8 (s.d. = 3.2), whereas the average apoptosis score in the food-restricted environment was 18.3 (s.d. = 12.0). Although there was no significant effect of sire or dam in the overall model, both terms significantly

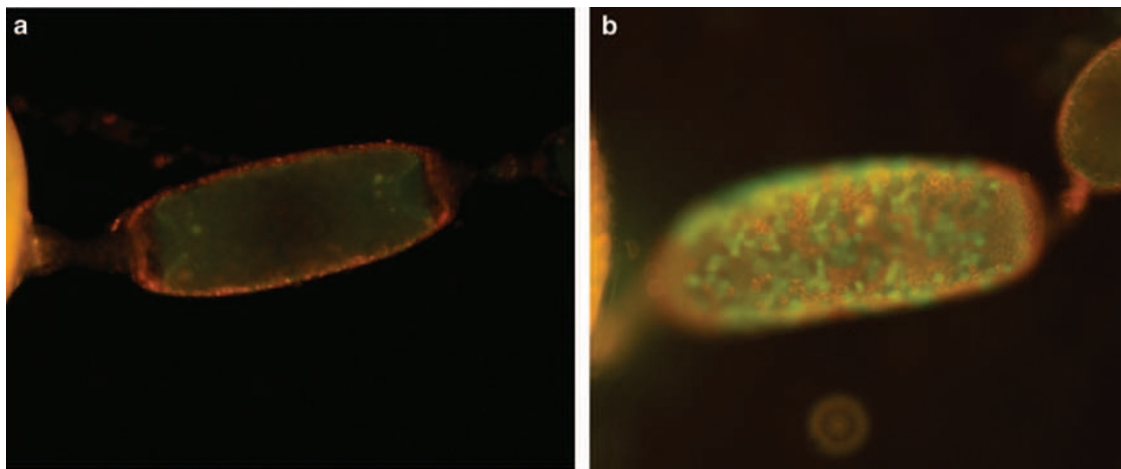


Figure 1 Apoptosis occurs in the ovaries of females that delay mating relative to sexual maturation. Ovaries from *ad libitum* food females 18 days post-adult eclosion are stained with a simultaneous assay for membrane potential (red fluorescence; Casp 3/7 Mito Casp, Bachem) and active caspase (green fluorescence; carbocyanine-labeled fluoromethyl ketone peptide inhibitor). Apoptosis occurs within oocytes (a; apoptotic oocyte surrounded by healthy follicle cells) and also in the follicle cells that act as nurse cells to the developing oocytes (b; a mixture of healthy and apoptotic follicle cells surrounding the immature oocyte). Similar staining patterns are seen in ovaries of food-restricted females.

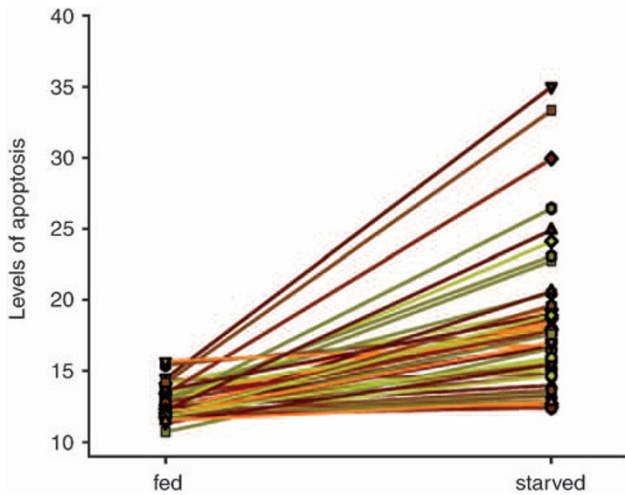


Figure 2 The levels of apoptosis expressed as sire family means of scores from mature oocytes of 18-day-old females. Each line connects the mean score of the daughters of one sire in the two treatments. The level of apoptosis was scored in 10 mature oocytes from a female offspring of each dam and the mean was then calculated for each dam.

Table 1 Partially nested two-way analysis of variance from SAS GLM investigating factors that influence oocyte apoptosis in *N. cinerea*

| Factor | d.f. | MS | F-value | P-value |
|-------------------------|------|---------|---------|---------|
| Food level | 1 | 9132.07 | 66.71 | <0.001 |
| Sire | 51 | 188.43 | 1.43 | 0.13 |
| Dam (sire) | 165 | 89.17 | 0.94 | 0.66 |
| Sire × food level | 51 | 137.62 | 1.45 | 0.04 |
| Dam (sire) × food level | 165 | 95.18 | 1.56 | <0.001 |
| Error | 846 | 60.91 | | |

Table 2 Separate REML estimates of components of variation underlying oocyte apoptosis for females reared under two conditions

| Treatment | V_A | V_C | V_E | V_P | $h^2 \pm s.e.$ | N_{sire} | N_{dam} | N_{offsp} |
|------------------------|--------|-------|-------|--------|-----------------|------------|-----------|-------------|
| <i>Ad libitum</i> food | 0.0049 | 2.12 | 8.00 | 10.12 | 0.00 ± 0.11 | 52 | 218 | 636 |
| Food-restricted | 47.93 | 1.20 | 95.39 | 144.52 | 0.33 ± 0.15 | 52 | 218 | 646 |

V_A is the additive genetic variance; V_C is the variance attributable to dominance, epistasis and common environment; V_E is the environmental variance; V_P is the phenotypic variance; h^2 is the narrow sense heritability.

interacted with the environment (Table 1). Thus, the expression and control of apoptosis in the two environments were clearly different.

Our design allowed us to estimate genetic influences on the levels of apoptosis within each environment (Table 2). In the *ad libitum* food environment, there was no significant effect of sire ($F=0.93$, d.f. = 51,165, $P=0.62$) but a significant effect of dam nested within sire ($F=1.65$, d.f. = 165,418, $P<0.001$). In contrast, in the food-restricted environment we found both a significant sire effect ($F=1.82$, d.f. = 51,165, $P=0.002$) and a significant dam nested within sire effect ($F=1.54$, d.f. = 165,428, $P<0.001$). In food-restricted females, the

environmental variance increased relative to that seen in fed females (Table 2, V_E), as would be expected under stressful conditions (Hoffmann and Parsons, 1991), but this increase alone did not explain the increase of phenotypic variance observed. Both conditions exhibited significant non-additive and common environment effects (Table 2, V_C) but there was relatively little difference in the non-additive genetic and common environment variance contribution between fed and food-restricted females. In contrast, in the *ad libitum* food environment, there was a lack of additive genetic variation compared with a significant level of additive genetic variation in the food-restricted environment (Table 2, V_A). Over 30% of the variation in levels of apoptosis observed in the ovaries of females under nutritional stress during sexual maturation can be attributed to additive genetic differences among families, whereas this is nearly zero in the *ad-libitum* food environment.

Discussion

We have provided the first demonstration of quantitative genetic variation in the levels of ovarian apoptosis and, indeed, of the regulation of apoptosis in any tissue. Our results show that ovarian apoptosis is turned on in response to nutritional stress, but to what extent depends on the genotype of the individual female. Therefore, in our population of cockroaches, selection could act on the regulation of ovarian apoptosis to change the rate of fertility loss with age under some environmental circumstances.

Although no other studies have examined quantitative genetic variation in the control of apoptosis, there is reason to believe that the complex interaction between environment and genetic variation we have uncovered in *N. cinerea* mirrors the control of ovarian apoptosis in other species. Ovarian apoptosis is involved in maintaining optimal functioning of the mammalian ovary (Amsterdam *et al.*, 1999), where fertility is lost with age through the loss of ovarian follicles through apoptosis (Morita and Tilly, 1999). Female mice in which the gene for Bax, a key regulator of apoptosis, has been knocked out have a prolonged ovarian lifespan (Perez *et al.*, 1999). In addition, in a linkage analysis of age at menopause, one of the two quantitative trait loci that explained variation in age of onset of menopause in humans was a region on chromosome 9 that contains a gene from the BCL2 family, an important protein in the apoptotic pathway (van Asselt *et al.*, 2004). Therefore, the potential exists for similar genetic variation in control of apoptosis over time in humans.

Apoptosis is critical to a large number of processes, which may mean that the genes controlling apoptosis often are under conflicting selection pressures and face trade-offs between different functions. However, our results highlight the fact that natural selection is unlikely to have removed all allelic variation in control of apoptosis for every function, be it apoptosis of oocytes or suppression of cancer, under all environmental circumstances. It is well documented that the heritability of a trait can change under different environmental conditions (Sgrò and Hoffmann, 1998; Hoffmann and Merilä, 1999). Heritability can decrease under stress due to an increased environmental variance or change because the history of selection differs between environ-

ments (Hoffmann and Parsons, 1991). Selection in a stable environment should decrease additive genetic variance, as alleles with favorable effects become fixed in the population. More additive genetic variance may remain for the expression of traits in environments that are encountered only rarely and where selection is therefore expected to be relatively weak.

In this study, the heritability of levels of apoptosis increased under nutritional stress because of additive genetic contributions to the phenotypic variation. Ovarian apoptosis in females that delay reproduction under conditions of adequate nutritional resource may reflect a developmental pathway to remove oocytes from the ovary at the end of an unproductive reproductive cycle to clear the system for the next cycle (Tyndale-Biscoe and Watson, 1977). The lack of additive genetic variance in this trait under benign conditions indicates that there has been strong directional or stabilizing selection on the levels of apoptosis in the oocytes as females pass sexual maturation without obtaining a mate. The induction of apoptosis in the ovaries of food-restricted females, however, is not simply a developmental response. These females are also responding to the environmental insult of nutritional stress. The observed increase in the additive genetic variance reflects the contribution of genes that control the onset of apoptosis only under nutritional stress. Because families were split between environments, we can show that the increase in additive genetic variance occurs even within families. However, because there is no heritability within one of the environments, this is not a genotype by environment interaction in which we would observe the change in expression of one set of genes over two environments. Rather, what we have measured is the action of genes controlling the initiation of the apoptotic pathway under nutritional stress, but not when females are fully fed. These genes are under different selection than the selection experienced by the genes controlling apoptosis during normal developmental progression. Interestingly, females that mate late relative to sexual maturation in the absence of food have higher fecundity and increased longevity compared with females that delay mating in the presence of food (Barrett *et al.*, 2009). Thus, the difference in genetic architecture of ovarian apoptosis in starved females that delay mating is reflected in variation in fitness consequences of timing of reproduction in the two nutritional environments.

The role of nutritional stress on ovarian function and apoptosis has been best studied in *D. melanogaster*. Starved female *D. melanogaster* have significantly fewer egg chambers at stage 8/9 due to an increase in apoptosis (Terashima and Bownes, 2004). *Broad-Complex (BR-C)* is a key gene for selecting between apoptosis and oocyte development, with differential expression of BR-C isoforms under nutritional stress, leading to the induction of apoptosis rather than yolk protein synthesis and oocyte development. Individual variation in expression of the BR-C isoforms has not been examined, but it is likely that such variation exists. Substantial genetic variation in starvation resistance in the natural populations has been documented (Harbison *et al.*, 2004) and an increase in starvation resistance is often negatively correlated with early fecundity (Service and Rose, 1985). The physiological mechanisms underlying the mediation of trade-offs between starvation resistance and

fecundity are largely unknown (Wayne *et al.*, 2006). Apoptosis of oocytes, however, would seem to play an important role in this process.

The evolution of ovarian apoptosis is complex. A full understanding of why reproduction may end early in a female's lifespan requires an integration of how genetics and selection might influence the physiological mechanisms that result in a loss of fertility, as well as these influences in various environments, including variation in access to resources. The results of this study provide insight into the complexity of the individual variation upon which selection could be acting, and this additional complexity needs to be incorporated into our considerations of the evolution of fertility loss.

Acknowledgements

We thank F Champion de Crespigny for practical help during the experiment. F Champion de Crespigny, D Hosken and T Tregenza provided constructive comments on an earlier draft of this paper. This research was supported by a Leverhulme grant to PJM and AJM. JH was supported by a National Environmental Research Council Fellowship.

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