Complex interactions between sperm viability and female fertility

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Sperm viability is a major male fitness component, with higher sperm viability associated with enhanced sperm competitiveness. While many studies have focussed on sperm viability from the male fitness standpoint, its impact on female fitness is less clear. Here we used a panel of 32 isogenic Drosophila simulans lines to test for genetic variation in sperm viability (percentage of viable cells). We then tested whether sperm viability affected female fitness by mating females to males from low or high sperm viability genotypes. We found significant variation in sperm viability among genotypes, and consistent with this, sperm viability was highly repeatable within genotypes. Additionally, females mated to high sperm viability males laid more eggs in the first seven hours after mating, and produced more offspring in total. However, the early increase in oviposition did not result in more offspring in the 8 hours following mating, suggesting that mating with high sperm-viability genotypes leads to egg wastage for females shortly after copulation. Although mating with high sperm-viability males resulted in higher female fitness in the long term, high quality ejaculates would result in a short-term female fitness penalty, or at least lower realised fitness, potentially generating sexual conflict over optimal sperm viability.

Sperm competition, the competition between sperm from two or more males to fertilize a given set of ova1, is a powerful selective agent shaping the evolution of ejaculates2. Besides almost ubiquitous effects on testis size3–7 and sperm number8–11, adaptations to increase sperm competitiveness have been reported in a number of traits related to the quality of individual sperm themselves, such as morphology12–17, performance18–22, and metabolism23,24. For example, the proportion of viable sperm within an ejaculate (sperm viability) can affect ejaculate competitiveness and influence siring success. This is particularly well established in insects where sperm viability increases with sperm competition risk in comparative studies25, and predicts the outcome of competitive fertilizations intraspecifically26,27. Additionally, males strategically adjust the viability of sperm in an ejaculate to perceived risks of sperm competition28,29 and environmental insults like pesticide exposure can reduce sperm viability30. While the impacts of sperm viability on male fitness are reasonably well established, it is less clear how sperm viability affects female fitness.

Females need sufficient numbers of viable sperm for full fertility (avoidance of sperm limitation) and this is one possible reason for the evolution of polyandry31–33. For example, the number of eggs a female Drosophila melanogaster can produce over a lifetime and the number of sperm stored after a single mating are of similar magnitude34,35, although variation in fertilization efficiency could mean that more than one sperm is required per egg fertilized36. This suggests that, with limited mating opportunities, low sperm viability might result in unfilled reproductive potential for females. Interestingly, ejaculate adaptations that increase male fitness may result in decreased female fitness, thereby creating sexual conflict over ejaculate quality (reviewed by Stockley37 and Edward et al.38).

Sexual conflict is widespread because the sexes are often under contrasting patterns of selection, such that each sex has different optima for a shared trait (intralocus sexual conflict) or different preferred outcomes of an intersexual interaction (interlocus sexual conflict). For example, many traits that improve male fitness because they are beneficial in male-male competition are costly for females39–43. For example, male cockroaches (Nauphoeta cinerea) engage in female guarding behaviour even after becoming sperm depleted, which reduces female fitness via enforced monogamy44. There can also be sexual conflict over seminal proteins and penis morphology, which can increase sperm competitiveness, but reduce female fitness45–47. For example, the D. melanogaster seminal protein sex-peptide increases male sperm competitiveness but reduces female longevity48. Additionally, increased numbers49, size50, and velocity19,20,50 of sperm in an ejaculate can increase sperm competitiveness, but also result...
in an increased risk of polyspermy and subsequent ova loss for females\(^5^1\). Males can also allocate fewer but more competitive sperm to ejaculates, in which case females might face sperm limitation\(^5^2\). Finally, ejaculate quality might trade off with non-ejaculate traits that enhance female fitness such as nuptial gifts or parental care\(^5^3\). So, there are multiple ways that variance in ejaculate quality – including sperm viability - could at least in principle impact female fitness.

Here we assessed potential effects of sperm viability on female fitness in *D. simulans*. Sperm viability is a key determinant of ejaculate quality and sperm competitiveness for insects in general\(^5^4\)\(^5^5\), and *Drosophila* in particular, where males produce highly viability ejaculates in response to increased sperm competition risk\(^5^6\). The present study was conducted using isofemale lines, which allowed us to replicate genotypes and assess sperm viability across them - thereby testing for genetic variation in sperm viability\(^5^7\), while also testing female fitness after receiving an ejaculate from genotypes with different sperm viability. Sexual selection has been extensively studied in *D. simulans*\(^5^8\)\(^5^9\)\(^6^0\). Sperm competitiveness is heritable and positively correlated with male attractiveness\(^6^1\), and while attractiveness is subject to inbreeding depression, sperm viability is not\(^6^2\). We used a panel of *D. simulans* genotypes to test for the presence of genetic variation in sperm viability, and to test how ejaculate quality (measured as sperm viability) affected female fitness by mating females to genotypes that had low- or high-viability sperm.

**Methods**

**Animals and husbandry.** In order to capture a significant portion of the genetic variation of a natural population of *D. simulans*, gravid females collected from Greece in 2010 were used to establish 32 isolines via full-sib mating. Isolines have been maintained by housing groups of males and females in 115 \(\times\) 45 mm vials with non-overlapping generations (approximately 50 adults per vial in two replicates per isolate for >100 generations).

All flies in this study were maintained in incubators at 25°C on a 12/12 light/dark cycle with ad libitum *Drosophila* Jazz-Mix food (Fisher Scientific). Light CO\(_2\) anaesthesia was used for fly collection, movement, and transference, and before fly dissections. No anaesthesia was used on flies within 24 hours of any mating assays as this affects mating behaviour\(^6^3\); instead, flies were individually aspirated into their housing vials.

**Sperm viability assay.** Two groups of 10 females and 5 males per isolate (hereafter, genotype) were established in vials (95 \(\times\) 25 mm) containing an excess of food to produce eggs for subsequent experiments. Flies were transferred into new vials every 24 hours for seven consecutive days (i.e. one vial per day per group). Eggs from these vials were incubated and virgin males were collected from these vials and housed individually (in 95 \(\times\) 25 mm vials). Five males per isolate were collected on each of seven consecutive days (from a different vial each day). These males were held individually until they were five days old and then the sperm viability of one random male per day per isolate (=seven males per genotype) was assessed.

To assay sperm viability, males were anaesthetized using CO\(_2\) and their seminal vesicles (site of mature sperm storage) were separated from the male reproductive tract, placed in 10 \(\mu\)l Beadle solution (128.3 mM NaCl, 4.7 mM KCl, 2.3 mM CaCl\(_2\)\(^6^4\)) on a microscope slide, and pierced using a 0.1 \(\times\) 12 mm insect pin. Sperm were forced out of the vesicles by gently pressing the vesicles with the flat side of the needle. Sperm viability was assessed after incubation in a moist chamber in the dark for 35 minutes using a LIVE/DEAD sperm viability kit (Molecular Probes L-7011), which contains two fluorescence probes that bind to DNA\(^6^5\)\(^6^6\). The first probe (SYBR-14) emits green fluorescence and is actively incorporated by living cells, and the second probe (propidium iodide, PI) emits red fluorescence and can only enter cells with damaged membranes (i.e. non-viable sperm). Briefly, an aqueous staining solution was prepared by diluting 2.4 \(\mu\)l of SYBR-14 stock solution (1 mM in DMSO), and 4.5 \(\mu\)l of PI stock solution (2.4 mM in water) in 93.1 \(\mu\)l of Beadle’s solution (working concentrations: SYBR-14 24 \(\mu\)M, PI 108 \(\mu\)M). Then, 5 \(\mu\)l of the staining solution were added to the drop containing the sperm (final concentrations SYBR-14 8 \(\mu\)M, PI 36 \(\mu\)M), and incubated for 5 minutes in the dark.

Sperm were examined at \(\times\)400 magnification under phase contrast and fluorescence microscopy (BX61 Olympus), the latter employing a mercury excitation beam passed through 480/20 and 535/30 nm filters to assess green and red fluorescence respectively. Two images, one for each fluorescence filter, were captured from each field using a camera (DP70 Olympus) attached to the microscope, and subsequently merged (DP Manager v2.1.1.163 Olympus) to assess sperm status. A mean of 266 (\(\pm\)11.06 SE) sperm were counted (Cell Counter plugin Image J v.1.51j8) in at least 2 randomly chosen fields per sample and scored live (green) or dead (red) (see example in Fig. 1).

Cells that were stained both green and red (less than 1% of sperm) were not counted. From our original sample of 224 males (1 male \(\times\) 7 days \(\times\) 32 isolines), 32 males were excluded because they generated samples with less than 40 visible cells (for a final sample size of 192 males). To test the reproducibility of our method a subsample of 19 males was recounted (10% of the original dataset) and a linear regression of the two samples with less than 40 visible cells (for a final sample size of 192 males). To test the reproducibility of our method a subsample of 19 males was recounted (10% of the original dataset) and a linear regression of the two samples with less than 40 visible cells (for a final sample size of 192 males). To test the reproducibility of our method a subsample of 19 males was recounted (10% of the original dataset) and a linear regression of the two samples with less than 40 visible cells (for a final sample size of 192 males).

**Female fitness: Assay 1.** Once sperm viability for all genotypes (isolines) had been measured, we chose four extreme genotypes (two low sperm viability, two high sperm viability isolines) (Fig. 1), to test whether sperm viability had any effect on the fertility of females from a tester stock population (maintained in the lab with free mating at population size \(>1000\) for \(>10\) years). Thirty randomly chosen females from each of the high and low sperm viability isolines (housed in groups of 5/vial) were allowed to lay eggs for 24 hours. Virgin males (30 per isolate) were collected from these vials and housed individually for five days with excess food. Each male was then mated to a 4-day-old, virgin female collected from the outbred stock-population.

One day before mating, tester females were transferred to a vial (95 \(\times\) 25 mm) containing 8 ml of Jazz mix food individually. The following day, males were introduced to females and pairs were left to interact for 7 hours. Then females were allowed to lay eggs for 8 days – this time period accurately captures *D. simulans* total fitness\(^6^7\). Since...
the mean remating interval for *D. simulans* females is approximately 2.7 days, and remating frequency within 24 h following the initial mating is less than 2% in this species, we assumed that the female reproductive output in the period of assessment corresponded to a single copulation with an experimental male. Initially females laid in egg counting vials (hereafter “initial period”) for seven hours (the time males were present), the number of eggs laid and the number of unhatched 20 h later were counted, and this allowed the proportion of eggs hatching to be estimated. After this initial laying period, females were allowed to lay eggs in three consecutive sets of vials for 2, 3, and 3 days respectively. All sets of vials were monitored twice a day in the following 15 days for signs of larval activity and adult eclosion. After the first adult eclosion, each vial was monitored for 7 days and then frozen at −20°C. The eclosed adults in each vial were counted and the total number of adult offspring per female was calculated as the sum of adults emerging from all four vials (hereafter “total offspring”). Eight females laid no eggs in the first seven hours and produced no offspring after 7 days of monitoring. These females were accounted as having failed to mate and were excluded from analysis, resulting in a final sample size of 112 females.

**Female fitness: Assay 2.** In the previous experiment, we allowed females to copulate and lay eggs for 7 hours without monitoring copulation latency or duration. To test whether males from different sperm-viability group differed in the time taken to copulate or in copulation duration – which could potentially affect initial egg laying time in the previous experiment – we conducted a second series of investigations. Briefly, males were introduced to females and pairs were observed and copulation latency and duration were recorded (pairs not mating within 3 hours were discarded). Subsequently females were transferred to egg counting vials (vial 1) immediately after copulation ceased, where they were housed for 2 h, and then sequentially transferred to a new vial every 2 h (vials 2, 3 and 4), for a total of 4 vials per female and 8 h of egg-laying time (hereafter “initial period”). In addition, we extended the time assigned for hatching evaluation from 20 to 25 h (one hour more than the 100% eclosion time estimated in reference) to ensure that the differences in egg hatching rates were not biased by different developmental times between sperm viability groups. The number of eggs laid in each vial and the number of unhatched (25 h later) were counted, and the proportion of hatched eggs was estimated. To test whether the egg-laying activity of the females at a later phase of the reproductive output differed from the initial period after mating, we compared the numbers of eggs laid and the hatching rate between the “late” assay period (vials 8 and 9) and a 4 h sub-interval of the initial period (vials 3 and 4, hereafter “early assay period”). We chose this combination of vials from the initial period since an extremely low number of females laid eggs in vials 1 and 2 (0 and 10 females respectively).
All sets of vials were monitored twice a day in the following 15 days for signs of larval activity and adult eclosion. After the first adult eclosion, each vial was monitored for 7 days and then frozen at −20°C. The eclosed adults in each vial were counted and the total number of adult offspring per female was calculated as the sum of adults emerging from all eight vials (hereafter "total offspring"). A total of 117 females in two experimental blocks were assessed. Two females that died after mating and one female that mated but laid no eggs in the first seven hours and produced no offspring after 7 days of monitoring were excluded from analysis (final n = 114 females).

The total number of adult offspring that emerged in the 8 days post mating were combined from both fitness assays for statistical analyses. However, eggs laid in the first hours after mating were not combined across assays as we did not analyse egg number for Assay 2. This is because in Assay 2 we continually disturbed egg laying by moving females to new vials every 2 hours. Thus, the number of early eggs laid was not comparable to Assay 1, and not indicative of how many eggs an undisturbed female would lay. Nonetheless, the percentage of eggs hatched is still informative from Assay 2 data.

**Statistical analysis.** All analyses were conducted using R version 3.6.0. 87. The effect of genetic background on sperm viability was analysed with generalized mixed effect models (GLMM), including isolate as a fixed factor (32 levels), and day of analysis (7 levels) and egg-laying group (64 levels: two parental groups per isolate) as random factors. We also estimated the magnitude of genotypic effects upon the variability in sperm viability. Since our isolines have been kept in the lab for > 5 generations, calculating the broad-sense heritabilities (\(h^2\)) would yield inaccurate estimates since inbreeding increases between line variance. Instead, we calculated the repeatability of sperm viability as the intraclass coefficient (ICC), which is an estimator of the upper limit of heritability. To estimate repeatability, we used a model similar to the one previously described for sperm viability, but included isolate as a random effect instead of as a fixed factor. However, since GLMMs do not provide a direct estimate of residual variance, and the interpretation of ICC estimations performed on GLMMs are only possible after transformation to the original data scale, repeatability cannot be calculated for GLMMs using traditional approaches. Instead, we used the method described by Nakagawa et al. 74 to estimate repeatability while accounting for the nonlinear relationships among the link scale and the scale of the data. The repeatability in the observation scale was estimated calculating the observation level variance as

\[
\frac{1}{np(1 - p)}
\]

where \(n\) = number of observations, and \(p\) = mean predicted viability value. This formula takes into account the specific logit link-function used in the model. In addition, we compared sperm viability between four extreme isolines (two low sperm viability, two high sperm viability), by means of a targeted post-hoc test using the same GLMM model than for the original isolate panel, and a subsequent estimated marginal means test (emmeans package).

The effect of sperm viability on mating success (second assay), latency to copulation (second assay), copulation duration (second assay), laid eggs (first assay and second assay separated), unhatched eggs (first assay), proportion of hatched eggs (first assay and second assay separated), offspring in the initial period (first assay), total offspring (first assay and second assay combined), and latency to copulation was analysed with GLMMs including sperm viability category (high or low) as a fixed factor, and isolate (4 levels) and egg-laying group nested within isolate (six parental groups per isolate) as random factors. When analysing data belonging to the second set of fertility experiments, we also included the experimental block (2 levels) as a random factor. When pooling data from the two fertility assays, we included the assay as a random factor and nested the blocks within. To test whether eggs laid and proportion of hatched eggs at the initial period after mating were similar to those on an equivalent time interval at a later phase of the egg-laying period (second assay only), we used a similar model than the ones detailed above with the addition of time and the interaction between time and sperm viability category as fixed effects. Time was included as a factor with two levels: early (vial 3 and 4) and late (vials 8 and 9).

Latency to copulation and copulation duration were analysed using Gamma distributions and “inverse” link functions (function “glm” in the lme4 package). The proportion of viable sperm, hatched eggs, and mating success were analysed using binomial distributions and “logit” link function. In the first two binomial models, the response variable was included in the model as a double vector containing the numbers of positive (i.e. sperm viability assay - viable sperm, fertility assays - hatched eggs) and negative (i.e. sperm viability assay - inviable sperm, fertility assays - unhatched eggs) results to account for the effect of the variable number of observations in the distribution of binomial data. Since these models showed over-dispersed data, we included an observation-level random effect nested within egg-laying group. Numbers of laid eggs, unhatched eggs, and offspring, were analysed using negative binomial distributions and “log” link functions (function “glmfit” in the glmfit package). These variables were tested for overrepresentation of zero values (zero-inflation) by comparing the observed proportion of zeros in the data set with the proportion of zeros expected for simulated distributions with identical parameters. If zero-inflation was present, we applied a zero-inflation term to the corresponding models by including the sperm viability category as a fixed effect predicting zero-inflation. When data from the two fertility assays was pooled, we added the assay as a random predictor of zero-inflation. Significance of the fixed effects for the GLMMs was tested by comparing the fit of two different models that either include or exclude the fixed factor using a likelihood ratio test (LRT).

**Results**

Sperm viability was variable across individual males with mean values ranging from 49–98% (Fig. 1). Male genotype significantly influenced sperm viability (GLMM: LRT \(X^2 = 47.99, p = 0.026\)). This indicates significant genetic variation in sperm viability. Consistent with this, the repeatability of sperm viability, which sets an upper limit on the heritability of sperm viability, was high at 59%.
We chose four extreme male genotypes based on their mean sperm viability to test whether viability affected female fitness - two extremely low sperm viability genotypes (lines 02 and 03 – these had poor sperm viability and little within-genotype variation) - and two with very high sperm viability (lines 31 and 32) (Fig. 1). A targeted post-hoc test revealed significant differences in sperm viability between the two groups (GLMM: LRT $\chi^2 = 16.90$, $p = 0.0007$), the "low viability group" averaging 73% (SE $\pm 2.9\%$) live sperm, and the "high viability" had a mean sperm viability of 90% (SE $\pm 1.4\%$).

In Assay 1 of female fitness, females mated with males from high sperm viability genotypes laid approximately 25% more eggs ($18.67 \pm 1.24$ vs $23.42 \pm 1.57$ low and high sperm viability respectively) in the first seven hours than females mated with males from low sperm viability lines (Table 1, Fig. 2A). Sperm viability did not affect the numbers of females laying no eggs (non-significant zero-inflation factor, Table 1). Both groups had a similar percentage of eggs (~26%) hatching into larvae (Table 1). This occurs because, although females mated to high sperm viability males laid more eggs, they had significantly more unhatched eggs ($13.56 \pm 1.04$ vs $17.14 \pm 1.24$, Table 1, Fig. 2B) (as well as weak non-significant trend of more hatched eggs: $5.11 \pm 0.75$ vs $6.28 \pm 0.69$, GLMM: LRT $\chi^2 = 1.58$, $p = 0.21$). Furthermore, the numbers of adult offspring emerging from eggs laid in the first seven hours after mating (vial 1) were similar between the two sperm viability groups (Table 1, Fig. 2C). This tentatively suggests more death occurred during the larval or pupal stages in the high sperm-viability treatment: they laid more eggs, but had the same number of adult offspring emerge during the first 7 hours after mating as the low sperm viability group.

In Assay 2 of female fitness, there were no significant differences in mating success, copulation latency or copulation duration between high or low sperm viability males (Table 1). Thus, differences described above are unlikely to be associated to differences in the timing of mating events. Additionally, these assays revealed that although the proportion of hatched eggs ($0.45 \pm 0.01$) was higher when assessed 25 h after oviposition (Assay 2) than when assessed after 20 h (Assay 1), again the proportion of hatched eggs did not differ between sperm viability groups (Table 1). This suggests that although assessing egg hatching at 20 h post-laying might underestimate the total proportion of hatched eggs; it does not bias the estimation with respect to sperm viability. This is true shortly after mating (4–8 h after mating) and 5 days after mating and oviposition (Table 2).

Combining data from the two fertility assays to compare female fitness, revealed females mated with high sperm viability genotypes showed a small but significant increase in the numbers of offspring produced over the complete 7-day laying period (low sperm viability: 28.79 ± 1.88, high sperm viability: 31.26 ± 1.93, Table 1, Fig. 2D) and sperm viability did not affect the number of females that produced zero adult offspring (non-significant zero-inflation factor, Table 1).

**Discussion**

Sperm viability varied significantly among *D. simulans* genotypes (isolines), and its repeatability was consistent with the heritability of sperm competitiveness previously reported in this taxon88. Similar results have been reported across other studies too89–91. Perhaps more interestingly, when females were mated with males from high sperm viability genotypes they laid more eggs in the first seven hours after mating, and produced more offspring in total. However, the short-term increase in oviposition (7 hours of laying) did not result in more offspring initially, suggesting females mated to males with high sperm viability wasted eggs shortly after copulation. This is despite the fact that mating with high sperm-viability males ultimately resulted in higher female fitness.

The short-term finding suggests that high sperm viability elevates the initial egg laying response in females. Although virgin *Drosophila* females are able to ovulate and lay (infertile) eggs at a very low rate, mating drastically and rapidly increases ovulation32. Both sperm and seminal-fluid proteins (Sfps) contribute to this effect, which is not as pronounced if one of these components is not present35,36,83–85. Although few studies have investigated how sperm (independently of Sfps) in the female reproductive tract affect ovulation, a possible mechanism explaining our finding has been described in butterflies where sperm activate mechano-receptors in the female tract to stimulate egg laying46. Alternatively, more viable sperm could result in more Sfps being transferred to females because Sfps can be attached to *Drosophila* sperm87. For example, in the closely related *D. melanogaster*, sex-peptide (one key Sfp) is bound to sperm and cleaved and released upon female sperm storage, triggering and sustaining ovulatory responses88,89, and *D. simulans* has a sex-peptide analog90. Either mechanism (binding or mechano-reception) could generate our results since it appears sperm have to actively enter the female sperm storage organs (sperm receptacle and spermatheca: collectively SSOs)90 and thus more viable sperm would result in more sperm storage, and consequently a higher degree of mechanical stimulation and/or higher concentrations of Sfps in the system. Although either of these outcomes could in turn promote a more exaggerated ovulatory response, the exact mechanism involved was not the focus of this study. However, it must be noted that *D. simulans* females only store about 30% of the sperm in an ejaculate and eject the rest before oviposition commences94. Thus, inviable sperm may never reach the SSOs and the sperm population ultimately stored may all be of high viability even if they come from ejaculates with different proportions of viable sperm. This would suggest a pre-storage overstimulation mechanism (mechanical or otherwise), in which the totality of the ejaculate might affect the female physiological state, as the more plausible alternative.

Nonetheless, in the short-term, more viable sperm resulted in more eggs laid, but no increase in offspring. This is possibly because, in many *Drosophila* species, females initiate ovulation and egg release before completing sperm storage85,82. These early-ovulated eggs are less likely to develop than eggs laid at later points89,94, suggesting that they may reach the fertilization site before sperm are ready to fertilize35. Thus a general acceleration of ovulation can decrease the efficiency of fertilization, explaining the increased number of unhatched eggs shortly after matings with high sperm-viability males. This suggests mating to high sperm-viability genotypes results in a short-term resource wastage for females. However, females mated to high sperm viability males also produced more offspring across the whole assay period: they had higher fitness. This is consistent with sperm viability...
determining the fertilizing capacity of a sperm population⁹²,⁹³, and decreased female fertility when viable sperm are depleted from SSOs⁹¹,⁹⁴,⁹⁵.

Associations among different sperm traits are generally not clear, with studies finding strongly positive⁷⁹,⁹⁶, to negative⁸¹,⁹⁷, and no⁹⁸,⁹⁹ correlations between characters thought to determine sperm quality. Moreover, the recruitment, maintenance, and usage of the sperm population in the SSOs of female Drosophila mainly depend on the female nervous system, SSO secretions, and male secreted SfPs⁹¹. This suggests that associations between the viability of ejaculated sperm and the long-term viability of the sperm in the SSOs can be complex. Surprisingly, our results appear to indicate a clear link, and that an ejaculate containing more viable sperm results in the establishment of a more fertile sperm reserve in the longer term.

While high sperm viability ultimately resulted in higher female fitness, the initial egg wastage at least represents “potential fitness” lost, and egg loss cannot be cheap. In fact one reason for the general lack of female secondary sex-traits is the cost of egg production¹⁰⁰–¹⁰². It therefore appears that a trait likely to be beneficial in

**Table 1.** Sperm viability effects on D. simulans female fertility. Values shown here (mean ± standard error) are the outcomes for singly mating females with males from low and high sperm viability genotypes. Statistical parameters for numbers of laid eggs (Wald’s z-value and variance), unhatched eggs, and offspring were estimated using a GLMM with negative binomial distribution and “log” link function, while in the case of proportion of hatching eggs a GLMM with binomial distribution and “logit” link function was used. X² and p values for fixed factors correspond to likelihood ratio tests between the full model and a restricted model excluding the fixed factor. *ZI: zero inflation term for datasets with higher numbers of zero values than predicted by the specified distribution.
sperm competition (sperm viability) has short-term costs for females, and thus there is likely to be some sexual conflict over sperm viability\(^{38}\). This is consistent with conflict being characterised as males and females agreeing on the destination, but not on how to get there\(^{103}\), and some male-female conflict over optimal sperm viability is consistent with the lack of inbreeding depression for it\(^{60}\). Inbreeding depression requires directional dominance, which results from a history of directional selection on a trait\(^{71,72,104}\), and lack of inbreeding depression can be a signal of stabilising selection. Since sexual conflict can act as a form of balancing selection\(^{105}\), our results might be indicative of net-stabilising selection on sperm viability – good for males but bad for females – which offers a tentative explanation for the lack of inbreeding depression in viability previously documented\(^{60,106}\). As noted however, female fitness was ultimately greater with high sperm viability despite the initial egg loss. How this would play out with multiple mating is not clear.

Figure 2. Sperm viability effect on \textit{D. simulans} female fertility. (A) Numbers of eggs laid in the first seven hours post mating (first fertility assay). (B) Numbers of unhatched eggs from those laid in the first seven hours post mating (first fertility assay). (C) Numbers of adult offspring eclosing from the initial oviposition vial (first fertility assay). (D) Numbers of adult offspring eclosing from the total eight days of egg-laying (first + second fertility assays). Values for females mated to males from low and high sperm viability genotypes are represented in red and green colors respectively. Boxes correspond to the interquartile range (IQR), whiskers extend to the largest value no further than 1.5 * IQR from the limit of the box, black dots represent the mean, black bars represent the median, and empty dots represent outlier values (exceeding 3 standard deviations from the mean). Profiles at the right of each panel illustrate the distribution density of each sperm viability category.
Table 2. Effects of sperm viability and time of egg laying after mating on D. simulans female fertility. Values shown here (mean ± standard error) correspond to eggs laid in two 4 h intervals (early: 4 h after mating, late: 5 days after mating) after singly mating virgin females with males from low and high sperm viability genotypes in the second set of mating assays. Statistical parameters (Wald’s z-value and variance) for numbers of eggs laid and offspring were estimated using a GLMM with negative binomial distribution and “log” link function, while in the case of proportion of hatching eggs a GLMM with binomial distribution and “logit” link function was used. X² and p values for fixed factors correspond to likelihood ratio tests between the full model and a restricted model excluding the fixed factor. *ZI: zero inflation term for datasets with higher numbers of zero values than predicted by the specified distribution.

| Dependent variable | Low viability          | High viability         | Independent variable          | z   | Variance | X²  | p     |
|--------------------|------------------------|------------------------|-------------------------------|-----|----------|-----|-------|
|                    | Early                  | Late                   | Early                        | Late|          |     |       |
| Laid eggs          | 2.83 ± 0.51            | 3.22 ± 0.47            | 2.43 ± 0.28                  | 3.10 ± 0.42 | Viability category (Fixed) | 1.438 | 1.00 | 0.317 |
| (Second assay)     |                        |                        | Time of egg laying (Fixed)   | 0.994 | 0.26     | 0.608 |
|                    |                        |                        | Viability * Time (Fixed)     | −0.923 | 0.90     | 0.344 |

Genotype (Random)  <0.001  
Egg-laying group (Random)  <0.001  
Female (Random)  0.006  
Block (random)  0.097  
Viability category (ZI - Fixed)*  1.106 | 2.01 | 0.126 |
Proportional hatch  0.42 ± 0.06 | 0.38 ± 0.05 | 0.49 ± 0.04 | 0.43 ± 0.06 | Viability category (Fixed) | 1.764 | 3.16 | 0.076 |
(Second assay)     |                        |                        | Time of egg laying (Fixed)   | 1.396 | 1.91     | 0.167 |
|                    |                        |                        | Viability * Time (Fixed)     | 0.822 | 0.68     | 0.410 |
Genotype (Random)  <0.001   
Egg-laying group (Random)  0.113  
Female (Random)  0.187  
Block (random)  0.038  

Overall, our study shows significant genetic variation and highly repeatable sperm quality in D. simulans, and that ejaculates with high sperm viability ultimately resulted in more offspring and higher female fitness. How males with higher sperm viability are able to cause females to waste resource and have higher fitness is not clear, but logic suggests there must be a female cost somewhere – it does not seem feasible to us that less efficiency generates higher output in a biological system, unless this is the optimal female response to the male “game” for example. This requires further investigation.

Data availability

The datasets generated and analysed during the current study have been uploaded on Dryad and are available for download: https://doi.org/10.5061/dryad.n8pk0p2qz.

Received: 7 February 2019; Accepted: 3 October 2019;
Published online: 25 October 2019

References

1. Parker, G. A. Sperm competition and its evolutionary consequences in insects. Biol Rev 45, 525–567, https://doi.org/10.1111/j.1469-185X.1970.tb0176.x (1970).
2. Birkhead, T. R., Hosken, D. J. & Pitnick, S. Sperm Biology: An Evolutionary Perspective. (Academic Press, 2009).
3. Gage, M. J. G. Associations between body size, mating pattern, testis size and sperm lengths across butterflies. Proc R Soc B 258, 247–254, https://doi.org/10.1098/rspb.1994.0169 (1994).
4. Stockley, P., Gage, M. J., Parker, G. A. & Møller, A. P. Sperm competition in fishes: The evolution of testis size and ejaculate characteristics. Ann Nat 149, 933–954, https://doi.org/10.1086/286031 (1997).
5. Hosken, D. J. Sperm competition in bats. Proc R Soc B 264, 385–392, https://doi.org/10.1098/rspb.1997.0055 (1997).
6. Hosken, D. J. & Ward, P. I. Experimental evidence for testis size evolution via sperm competition. Ecol Lett 4, 10–13, https://doi.org/10.1046/j.1461-0248.2001.00198.x (2001).
7. Firman, R. C. & Simmons, L. W. The frequency of multiple paternity predicts variation in testes size among island populations of house mice. J Evol Biol 21, 1524–1533, https://doi.org/10.1111/j.1420-9101.2008.01612.x (2008).
8. Immel, S. et al. Resolving variation in the reproductive tradeoff between sperm size and number. Proc Natl Acad Sci USA 108, 5325–5330, https://doi.org/10.1073/pnas.1009059108 (2011).
9. Tourmente, M., Delbarco Trillo, J. & Roldan, E. R. S. No evidence of trade-offs in the evolution of sperm numbers and sperm size in mammals. J Evol Biol 28, 1816–1827, https://doi.org/10.1111/jeb.12698 (2015).
10. Aron, S. et al. Sperm production characteristics vary with level of sperm competition in Cataglyphis desert ants. Funct Ecol 30, 614–624, https://doi.org/10.1111/1365-2435.12333 (2016).
11. Liao, W. B. et al. Ejaculate evolution in external fertilizers: Influenced by sperm competition or sperm limitation? Evolution 72, 4–17, https://doi.org/10.1111/evol.13372 (2018).
12. Johnson, D. D. P. & Briskie, J. V. Sperm competition and sperm length in shorebirds. Condor 101, 848–854 (1999).
13. Byrne, P. G., Simmons, L. W. & Roberts, J. D. Sperm competition and the evolution of gamete morphology in frogs. Proc R Soc Lond B Biol Sci 270, 2079–2086, https://doi.org/10.1098/rspb.2003.2433 (2003).
52. Wedell, N., Gage, M. J. G. & Parker, G. A. Sperm competition, male prudence and sperm-limited females.

46. Civetta, A. & Clark, A. G. Correlated effects of sperm competition and postmating female mortality.

49. Wedell, N. & Cook, P. A. Butterflies tailor their ejaculate in response to sperm competition risk and intensity.

45. Chapman, T., Liddle, L. F., Kalb, J. M., Wolfner, M. F. & Partridge, L. Cost of mating in...
94. Collins, A. M. Functional longevity of honey bee, Apis mellifera, queens inseminated with low viability semen. *J Apic Res* **43**, 167–171, https://doi.org/10.1080/00218839.2004.11101131 (2004).
95. Reinhardt, K. & Ribou, A.-C. Females become infertile as the stored sperm’s oxygen radicals increase. *Scientific reports* **3**, 2888, https://doi.org/10.1038/srep02888 (2013).
96. Gomez Montoto, L. et al. Sperm competition, sperm numbers and sperm quality in muroid rodents. *PLoS One* **6**, e18173, https://doi.org/10.1371/journal.pone.0018173 (2011).
97. Taborsky, M., Schütz, D., Godfray, H. C. O. & Sander van Doorn, G. Alternative male morphs solve sperm performance/longevity trade-off in opposite directions. *Science Advances* **4**, eeeaap8563 (2018).
98. Simmons, L. W. & Kotiaho, J. S. Evolution of ejaculates: patterns of phenotypic and genotypic variation and condition dependence in sperm competition traits. *Evolution* **56**, 1622–1631 (2002).
99. Damiens, D., Bressac, C. & Chevrier, C. The effect of age on sperm stock and egg laying in the parasitoid wasp, Dinarmaeus basalis. *J Insect Sci* **3**, 22 (2003).
100. Wang, Y., Salmon, A. B. & Harshman, L. G. A cost of reproduction: oxidative stress susceptibility is associated with increased egg production in *Drosophila melanogaster*. *Experimental gerontology* **36**, 1349–1359, https://doi.org/10.1016/S0531-5565(01)00095-X (2001).
101. Yang, C. H., Belawat, P., Hafen, E., Jan, L. Y. & Jan, Y. N. *Drosophila* egg-laying site selection as a system to study simple decision-making processes. *Science* **319**, 1679–1683, https://doi.org/10.1126/science.1151842 (2008).
102. Hosken, D. J., Alonzo, S. H. & Wedell, N. Why aren’t signals of female quality more common? *Anim Behav* **114**, 199–201, https://doi.org/10.1016/j.anbehav.2016.02.015 (2016).
103. Hosken, D. J., Archer, C. R. & Mank, J. E. Sexual conflict. *Curr Biol* **29**, R451–R455, https://doi.org/10.1016/j.cub.2019.03.052 (2019).
104. Roff, D. A. *Evolutionary Quantitative Genetics*. (Chapman and Hall, 1997).
105. Mank, J. E. Population genetics of sexual conflict in the genomic era. *Nature Rev Genet* **18**, 721–730, https://doi.org/10.1038/nrg.2017.83 (2017).
106. Ala-Honkola, O. et al. Inbreeding reveals mode of past selection on male reproductive characters in *Drosophila melanogaster*. *Ecology and evolution* **3**, 2089–21002, https://doi.org/10.1002/ece3.625 (2013).
107. McNamara, J. M. & Houston, A. I. Credible threats and promises. *Philos Trans R Soc Lond B Biol Sci* **357**, 1607–1616, https://doi.org/10.1098/rstb.2002.1669 (2002).

**Acknowledgements**

We would like to thank Noam Hosken, Keiko Oku, and Jacqui Glencross for their help with data collection and fly stocks maintenance. This study has received funding from the European Union’s Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie (Grant Agreement No. 749784).

**Author contributions**

M.T., C.R.A. and D.J.H. conceived and designed the experiments. M.T. conducted the experiments on sperm viability and fertility. M.T. analysed the data. All authors wrote and reviewed the manuscript.

**Competing interests**

The authors declare no competing interests.

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