Do Candidate Genes Mediating Conspecific Sperm Precedence Affect Sperm Competitive Ability Within Species? A Test Case in *Drosophila*

Alberto Civetta¹ and Scott Finn

Department of Biology, University of Winnipeg, Winnipeg, MB, R3B 2E9 Canada

ABSTRACT When females mate to multiple males, the last male to mate fathers the majority of progeny. When males of different species inseminate a female, the sperm of the male conspecific to the female is favored in fertilization in a process known as conspecific sperm precedence (CSP). A large number of studies in Drosophila have assayed the genetic basis of sperm competition, with a main focus on D. melanogaster and accessory gland protein genes. Only a few studies have attempted to disentangle the genetic basis of CSP between related species of Drosophila. Although there is no a priori reason to believe that genes influencing intraspecific sperm competitive ability might also mediate conspecific sperm precedence, no study has addressed the question. Here, we test a group of candidate CSP genes between D. simulans and D. mauritiana for their effect on sperm competition in D. melanogaster. The use of P-element insertion lines identified CG14891 gene disruption as the only one causing a significant decrease in second male paternity success relative to wild-type and ebony tester males. The gene disruption affected both sperm displacement and the sperm fertilizing ability. Out of five genes tested using RNA interference, only gene knockdown of CG6864 (Mst89B) significantly reduced the male's ability to father progeny when second to mate. Our results suggest that CG14891 and CG6864 might have been co-opted from an intraspecies gene function (i.e., sperm competition) into an interspecies avoidance phenotype (i.e., CSP). Alternatively, the dual role of these genes could be a consequence of their pleiotropic roles.

KEYWORDS

sperm competition conspecific sperm precedence gene co-option pleiotropy

The phenomenon of sperm competition in *Drosophila melanogaster* has been extensively studied using mating laboratory assays as well as surveys of parenthood from wild-caught females. Overall, studies show that when *Drosophila* females mate with two or more males, the last male to mate has the greatest paternity success (Clark 2002). Second male paternity advantage holds also in heterotypic crosses involving *D. melanogaster* cosmopolitan and Zimbabwe races (Dixon *et al.* 2003).

Several genes have been tested for their role in competitive success during multiple mating using both gene manipulation assays and tests

that associate gene sequence polymorphism with variation in sperm competitive success measured from paternity scores. The focus has been mainly on male accessory gland proteins (Acps). For example, the deletion of Acp62F increases the male's ability to place sperm in storage, and significant associations have been found between polymorphisms at Acp62F and sperm competitive ability (Fiumera et al. 2007; Mueller et al. 2008). Acp29AB and Acp36DE are needed for sperm storage and they also show significant associations between sequence polymorphism and variation in sperm competitive ability (Clark et al. 1995; Fiumera et al. 2005; Wong et al. 2008; Avila and Wolfner 2009). Other Acps have been found to be required for the proper utilization of stored sperm for fertilization (Avila et al. 2011). There is also both indirect and direct evidence of non-Acp genes influencing sperm competitive ability. Associations have been found between four genes within the X chromosome, a chromosome with a scarcity of genes for seminal proteins, and differences in sperm competitive ability in D. melanogaster (Greenspan and Clark 2011). Recently, deletion of the X-linked genes of the sperm dynein intermediate chain (Sdic) multigene family in D. melanogaster has been shown to affect sperm competitive ability without disrupting sperm motility or male fertility (Yeh et al. 2012).

Copyright © 2014 Civetta and Finn

doi: 10.1534/g3.114.012476

Manuscript received June 3, 2014; accepted for publication July 5, 2014; published Early Online July 16, 2014.

This is an open-access article distributed under the terms of the Creative Commons Attribution Unported License (http://creativecommons.org/licenses/ by/3.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Supporting information is available online at http://www.g3journal.org/lookup/ suppl/doi:10.1534/g3.114.012476/-/DC1

¹Corresponding author: Department of Biology, University of Winnipeg, Winnipeg, MB, R3B 2E9 Canada. E-mail: a.civetta@uwinnipeg.ca

Table 1 List of Drosophila melanogaster strains used

| Strain | Source | Genotype | Construct | Gene affected |
|--------|----------|---------------------------|-----------|-------------------|
| 6550 | Ward | Wild-type | NA | NA |
| 1658 | BDSC | e/e | NA | ebony |
| 18320 | BDSC | +/TM6Be, Tb | PBac | CG7478(Actin 79B) |
| 18237 | BDSC | +/TM6Be, Tb | PBac | CG31542 |
| 18348 | BDSC | TM6Be, Tb | PBac | CG7362 |
| 24853 | BDSC | e/TM3e, Sb | Р | CG3158(spindle-E) |
| 26029 | BDSC | Wild-type | Mi | CG14891 |
| F01790 | Exelixis | Wild-type | PBac | CG6255 |
| 19276 | BDSC | +/TM6Be, Tb | Р | CG31287 |
| 102118 | VDRC | Wild-type | RNAi | CG14891 |
| 102666 | VDRC | Wild-type | RNAi | CG6864 (Mst89B) |
| 103437 | VDRC | Wild-type | RNAi | CG31287 |
| 40461 | VDRC | Wild-type | RNAi | CG3610 |
| 109692 | VDRC | Wild-type | RNAi | CG4836 |
| 25708 | BDSC | UAS-Dcr2; Act5C-GAL4/ CyO | Driver | NA |

BDSC, Bloomington Drosophila Stock Center; VDRC, Vienna Drosophila RNAi Center. Transposable element inserts are: PBac, PiggyBac; P, Pelement; Mi, Minos; NA, not applicable.

We have recently gained a large amount of knowledge regarding the mechanisms of sperm utilization and competition by using males with fluorescently labeled sperm. For example, the use of transgenic males of *D. melanogaster* with fluorescently labeled sperm showed an important role played by second male displacement of resident sperm driven by sperm numbers (Civetta 1999; Manier *et al.* 2010). More recent studies have uncovered a common set of sperm precedence mechanisms shared among closely related species of *Drosophila*, yet a complex scenario of rapid diversification. For example, male sperm characteristics such as differences in sperm length can contribute to sperm advantage, but females can influence sperm utilization through ejection or by controlling the use of sperm for fertilization (Lupold *et al.* 2013; Manier *et al.* 2013a,b,c).

The observation of conspecific over heterospecific male advantage to father progeny when in competition has proven to act as a postmating prezygotic reproductive isolation barrier between species of Drosophila as well as other insects (Price 1997; Howard et al. 1998; Rugman-Jones and Eady 2007; Matsubayashi and Katakura 2009; Larson et al. 2012; Tyler et al. 2013). Among species of the Drosophila simulans clade, double mating of females to conspecific and heterospecific males have shown that the majority of progeny is fathered by the conspecific male regardless of the mating order, with the breakdown in competitive success of the heterospecific males being attributed to poor sperm displacement ability (Price et al. 2000, 2001). More recently, sperm dumping by females, or simply poor sperm storage, has been reported to influence the fate of heterospecific sperm in other species of Drosophila as females rapidly lose sperm from males of related species (Matute and Coyne 2010; Sagga and Civetta 2011). A recent study directly addressed the occurrence and causes of CSP between D. simulans and D. mauritiana by competing males with differentially labeled sperm. Despite differences between species, the study revealed many commonalities between previously known intraspecific mechanisms of sperm precedence and CSP, including a major role of sperm displacement and female control of sperm utilization through sperm ejection and fertilization biases (Manier et al. 2013c).

Only a few studies have attempted to map the genetics of CSP in *Drosophila* and other insects (Civetta *et al.* 2002; Britch *et al.* 2007; Levesque *et al.* 2010). The lack of attempts has probably been a consequence of not having a clear understanding of mechanisms; however, it is feasible to map loci that contribute to the phenotypic manifestation of CSP, *i.e.*, conspecific advantage to father progeny.

We have used D. simulans lines with mapped genetic introgressions from D. mauritiana to test the effect of different heterospecific third chromosome introgressions on D. simulans second male paternity success. We identified a group of candidate CSP genes based on mapped loci with combined analysis of adaptive gene coding sequence diversification and male reproductive gene expression (Levesque et al. 2010). Given the commonalities in mechanisms of sperm utilization and precedence between species, we used a combination of D. melanogaster P-element insertions and RNAi constructs to test the effects of 10 previously identified CSP candidate genes on second male paternity success in D. melanogaster. P-element insertions were used to disrupt genes that have previously shown an effect on CSP based on coding sequence diversification, whereas RNAi knockdowns were used to target CSP candidate genes on the basis of differential gene expression. We found evidence in support of two CSP genes, CG14891 and CG6864, as influencing sperm competition within D. melanogaster. The results indicate the possibility of co-option of gene function from intraspecific male fitness to a role in interspecies hybridization avoidance. It is alternatively possible that both genes might



Figure 1 Schematic diagram of design for sperm competition assays (A) and creation of tester males for gene knock down assays (B). TG, target gene.

Table 2 Viability correction factors (prop wt)

| P-strain | Gene | Ν | Prop wt | RNAi strain | Gene | N (Cy; +) | Prop wt (Cy; +) |
|----------|---------|----|---------|-------------|---------|-----------|-----------------|
| 18237 | CG31542 | 23 | 0.53 | 102118 | CG14891 | 14; 34 | 0.59; 0.56 |
| 18320 | CG7478 | 18 | 0.54 | 103437 | CG31287 | 19; 33 | 0.58; 0.54 |
| 18348 | CG7362 | 22 | 0.53 | 102666 | CG6864 | 10; 24 | 0.55; 0.55 |
| 24853 | CG3158 | 10 | 0.50 | 40461 | CG3610 | 10; 10 | 0.53; 0.55 |
| 26029 | CG14891 | 12 | 0.51 | 109692 | CG4836 | 21; 10 | 0.51; 0.51 |
| F01790 | CG6255 | 20 | 0.55 | | | | |
| 19276 | CG31287 | 22 | 0.60 | | | | |

N= Number of females assayed.

have come to influence both intraspecies and interspecies phenotypes as a consequence of gene pleiotropic effects.

MATERIALS AND METHODS

Phenotypic test of candidate genes in D. melanogaster

Drosophila stocks carrying *P*-element insertions at genes of interest were obtained from the Bloomington or Exelixis stock centers. Transgenics carrying an inducible UAS-RNAi construct for genes of interest were purchased from the Vienna Drosophila RNAi Center (VDRC) and a Dicer (Dcr) GAL4 driver stock (25708) from the Bloomington Drosophila Stock Center (BDSC) (Table 1). Flies were reared in cylindrical polypropylene bottles containing cornmeal-molasses-yeastagar (CMYA) medium at 24°C in a 12-hr light-dark cycle. For controlled crosses, virgin female flies were collected and maintained in polypropylene vials containing fresh CMYA medium, with no more than 20 copies per vial.

Males from lines with P-element gene insertions were used as tester second males to mate to females homozygous for an ebony (e/e) mutation that had already been singly mated to same-aged ebony males (male 1). Similarly, F1 males from the cross between the UAS-RNAi construct stocks and the Dcr-GAL4 driver stock (25708) were used as second tester males (Figure 1). All matings were observed every 15 min for a total period of 8 hr to avoid multiple mating to any one male. After the first mating, females were stored in a vial for 2 d (vial 1), at which point they were paired for a second mating with the tester males (male 2). Four days after the second mating, females were transferred to a fresh vial (vial 3). Progeny from vials 1, 2, and 3 were counted on the 20th day after the beginning of oviposition and scored based on phenotypic body coloration, with ebony progeny being fathered by the first male and nonebony progeny being fathered by the second male. Some gene inserts were over balancers that segregate ebony (Table 1), scores were corrected for segregation bias by monitoring the Tubby phenotype (e.g., +/e and e/e vs. e/e, Tb). For stock 24853, which is phenotypically ebony (Table 1), males from a wild-type stock (6550 from Ward's Natural Science) were used as first to mate. We also tested males from the wild-type and the ebony (1658 BDSC) stocks as second to mate. Females that did not produce progeny from male 1 in vial 1 were discarded from further analysis (i.e., no first mating). The fraction of progeny in vial 2 and vial 3 sired by the tester male was designated as P2.

We corrected P2 scores for differences in egg-to-adult viability of competing developing genotypes in a vial (Gilchrist and Partridge 1997). Viability was scored by crossing ebony females to tested males of wild-type phenotype or the +/TM6Be, Tb stock (Table 1). In tests involving wild-type males, the F1 females were backcrossed to ebony color males, and deviations from 1:1 segregation of wild-type to ebony body color phenotypes were scored. For the test crosses with +/TM6Be Tb males, e/+ F1 females were crossed to e/TM6Be Tb F1

males, and the segregation of wild-type, ebony color, and tubby size with ebony color provided counts to correct viabilities. Finally, for the tester males from stock 24853 (e/TM3e, Sb) (Table 1), ebony females were crossed to wild-type males (stock 6550 from Ward's Natural Science) and F1 females (e/+) were crossed to males from stock 24853. The segregation of wild-type to ebony color gave the counts used for corrections. The viability correction factor is described elsewhere (Clark *et al.* 1999), but it briefly involves dividing scores in the sperm competition trials by twice the ratio of wild-type offspring sired by females in the noncompetitive backcross mating assays described above.

Gene expression data analysis

The effectiveness of the Dcr-GAL4 driver to trigger downregulation of the responder gene construct via RNAi induction (Figure 1) was measured using quantitative reverse-transcriptase PCR (qRT-PCR). We collected and aged males for 4–6 days and dissected single abdomens from each line. Single fly abdomen samples were stored in lysis buffer at -70° C until RNA extraction. RNA was extracted using the Qiagen RNeasy plus mini kit following the manufacturer protocol. cDNA was synthesized using an iScript select cDNA synthesis kit (BioRad) and cDNA was quantified using a nanophotometer so that equal amounts of total cDNA were used in each qRT-PCR reaction. qRT-PCR was performed using the IQ SYBR Green Supermix kit from BioRad and performed in a MiniOpticon PCR System (BioRad). We ran PCR reactions from at least three biological replicates (different RNA samples) for each *Drosophila* line tested.

Primers for qRT-PCR were designed using Primer3 (http://frodo.wi. mit.edu/) to span an exon/intron boundary when possible, because it helped further control for possible DNA contamination. The efficiency of all primers pairs was tested by creating a standard curve using the threshold cycles generated from a dilution series of a template (amplified by qRT-PCR) and plotting those threshold cycle values against the log of the template dilution used in each reaction. The slope of the standard curve was used to assess primer efficiency (Supporting Information, Table S1). Rpl32 was used to calibrate expression, eliminating error from differences in RNA concentration between samples. After amplification, we tested for the presence of single amplification products by using a melting step using 0.5°C increments between 55° and 99°C and 1-sec hold reads. The Δ CT of the tested sample was calculated by subtracting the CT of the reference gene (RpL32) from the CT of the target gene. For each gene, relative expression of the knockdown was normalized by setting the average Δ CT of the F1 Curly males (no knockdown) (Figure 1) as calibrator.

Data analysis

Phenotypic data were analyzed using one-way ANOVA with strains as the main factor followed by *a posteriori* Scheffe's test. Pair-wise



Figure 2 Average second male paternity of males carrying P-element insertions at different candidate genes. Error bars represent \pm 1 SEM. Shared letters above columns indicate that the averages are not statistically different (*post hoc* Scheffe's test). wt, wild-type.

comparisons regarding the effect of gene knockdowns on both gene expression and phenotypic scores were performed using two-sample *t*-test.

RESULTS

Comparison of second male paternity success among transposable element insert lines

We tested second male paternity success for seven strains of *D. mel-anogaster* carrying transposable elements inserts for genes of interest, as well as an ebony mutant and a wild-type strain (File S1). The setting involved scoring phenotypes for 20,411 progeny produced by 290 females. We found slight viability deviations (Table 2) and P2 values were corrected for differences in viability of the genotypes being scored (see *Materials and Methods*). We detected significant differences among strains in second male paternity success (F_{8, 230}= 15.86; *P* < 0.001). Not surprisingly, the average sperm competitive ability of





Figure 3 Average of the difference between second male paternity in vial 3 (P2v3) and in vial 2 (P2v2) for males carrying P-element insertions at different candidate genes.



Figure 4 Average relative gene expression of males with a knockdown of candidate genes (light gray bars) relative to controls (average at zero). Error bars represent \pm 1 SEM.

and ebony flies. Three strains carrying gene disruptions of CG14891, CG31287, and CG6255 show no increments in average second male paternity success from the second to third vial (Figure 3). Overall, only gene disruption of CG14891 had an effective and significant consequence in decreasing the overall competitive ability of males when second to mate, with disruption of other genes having only temporary effects.

Comparison of gene knockdown effects on second male paternity success

We have previously identified eight candidate genes on the basis of mapping differences in CSP using introgression lines and significant differences in male reproductive tract gene expression between genes within the mapped locations. Here, we tested the effectiveness of knockdowns for five (CG6864, CG14891, CG31287, CG3610, and CG4836) of the original eight candidate genes and the effect of the knockdown on second male paternity success (File S2 and File S3). The design allowed us to produce F1 progeny flies with both a Dicer-GAL4 driver and a UAS-RNAi responder that produced a ds-hairpin RNA of the gene to be targeted for knockdown (non-Curly flies) as well as Curly wing flies without the GAL4 driver that were used as controls (Figure 1). Comparisons of gene expression showed a reduction of gene expression in all gene knockdowns. The knockdown for CG4836 showed no significant downregulation compared with Curly flies carrying only the responder (t_6 = 0.80; P = 0.231) and CG14891 was only marginally significant (t_7 = 1.77; P = 0.083), with all other gene knockdowns showing significant reductions in gene expression relative to controls (Figure 4). None of the knockdowns significantly reduced the average fecundity of males and CG3610 showed an increase, although not significant, in average fecundity of the knockdown males (Figure 5A). Surprisingly, most knockdowns showed a slight, but not significant, increase in second male paternity success, with the effect being the largest for CG3610 (Figure 5B). Only the

knockdown of CG6864 (Mst89B) had a significantly lower average P2 than the control (t_{52} = 1.59; P = 0.041) (Figure 5B). The effect was significant immediately after second mating (vial 2; t_{51} = 2.12; P = 0.014), but not as a delay effect (vial 3; t_{49} = 1.12; *P* = 0.115) (Figure 6).

DISCUSSION

It has been shown in Drosophila melanogaster that the amount of sperm stored by the second male to mate and the proportion of progeny he fathers are significantly correlated, suggesting that paternity success of the last male to mate is related to his ability to place sperm in storage and displace resident sperm (Civetta 1999; Manier et al. 2010). An actual displacement from the seminal receptacle of resident sperm by incoming sperm had been more accurately described using fluorescent transgenic sperm flies as occurring rather



Figure 5 Average fecundity (A) and average second male paternity success (B) of control males (dark gray bar) and gene knockdowns (light gray bars) (A). Error bars represent \pm 1 SEM.



Figure 6 Difference in second male paternity success (P2) between gene knockdown and control males in vial 2 (dark gray bar) and in vial 3 (light gray bar).

rapidly, with an apparent establishment of an approximately 2-to-1 advantage of the incoming sperm over resident sperm within 72 hr (Manier et al. 2010). Given that females spent the first days after second mating in vial 2 before been transferred to vial 3, poor second male paternity success in vial 2 might reflect the incoming sperm inability to properly store and displace resident sperm. The effects of P-element gene insert disruptions on sperm storage and displacement ability were only marginal because they reduced average P2 relative to wild-type males but not ebony tester males. Poor second male paternity scores over vial 3 might be linked to deficiencies in sperm physiology and fertilization ability once in storage. Several gene disruptions significantly affected the male's sperm fertilization ability once in storage. However, some of the fertilization effect (vial 3) could be driven by the marginal effects on sperm storage and displacement (vial 2). For example, males with CG7478 inserts showed significantly lower fertilization ability, but there was an effective increase in their paternity success from vial 2 to vial 3. Similarly, the disruption in fertilization ability of CG31287 was not sufficient to affect the overall second male paternity success of the males. CG6255 gene disruptions did not affect overall second male paternity success, which we attribute to the fact that despite its poor fertilization ability (vial 3 effect) such males did well enough at sperm placement in storage and displacement of resident sperm (vial 2 average P2 = 0.6) to compensate possible deficiencies. Thus, only the disruption of CG14891 affected the overall second male paternity success of its carriers, with the effect of the disruption being particularly strong in vial 3 (fertilization effect). CG14891 is expressed at high levels in the testes but not much is known about its function, except for the suggestion that it might be an F-box protein. F-box genes have gene regulatory roles in a wide variety of functions, including signal transduction, cell proliferation, and growth (Ou et al. 2003; Rogers et al. 2009).

Most gene knockdowns did not affect fecundity or second male paternity success. We interpret this result in a conservative manner as evidence that such genes do not influence such phenotypes, although there are a few caveats: (1) we cannot rule out that the effect of this single gene knockdown is being rescued by partner genes and (2) the controls we used, although genotypically more equivalent to the knockdown flies than wild-types, were carriers of the hairpin constructs, so they could have been leaky effects even in the absence of the GAL4 driver. The only significant effect of the gene knockdowns was for CG6864 (Mst89B) on second male paternity success. The significant effect of this gene knockdown was evident on vial 2, suggesting that the gene defect affects the ability of males to store or effectively displace resident sperm. This result is particularly interesting given that yeast two-hybrid data have suggested interactions between Mst89B and Cdlc2, a microtubule motor activity protein expressed in the sperm, and Acp62F, an accessory gland protein that increases a male's ability to place sperm in storage (Giot et al. 2003; Mueller et al. 2008). It is possible to speculate that the interaction among these proteins, and perhaps others, might exert an effect on the sperm ability to reach and properly function in storage. Although the knockdown of Mst89B significantly diminished second male paternity, it did not cause a breakdown (i.e., P2 < 0.5) of second male paternity success, indicating a polygenic basis underlying changes in sperm competitive ability and that Mst89B might not function as a major gene.

Our results can be interpreted as providing support for the hypothesis that genes can be co-opted from a role in providing an advantage in competitive paternity success within species to underlying the genetics of an interspecies hybridization avoidance phenotype (CSP). The fact that intraspecific mechanisms of sperm competition appear as contributors to CSP (Manier *et al.* 2013c) further supports the possibility of gene co-option. Thus, gene co-option might have facilitated the diversification of a reproductive-related phenotype over a relatively short evolutionary time scale in a manner similar to how it has facilitated the origin of evolutionary novelties during the diversification of other phenotypes in a wide variety of species (Moczek and Rose 2009; Martin *et al.* 2014). However, it is alternatively possible that both genes might have diverged to play roles in CSP as a consequence of pleiotropic effects. Future studies should attempt gene silencing experiments in species other than *D. melanogaster*.

Finally, the localization of both CG14891 and CG6864 (*Mst89B*) in the chromosome region 89B is of interest. Although it is unclear whether the 89B location harbors other potentially important genes, there are some interesting candidates that are localized nearby *Mst89B* (3R:12,092,734.0.12,094,155), such as spermatogenesis genes *asunder* (3R:12,089,993.0.12,092,613) and *gish* (3R:12,098,174.0.12,130,490), as well as *CG14879* (3R:12,171,547.0.12,174,948), a functionally unknown but highly expressed male accessory gland gene (http://flybase.org/). Such genes and others in the regions should be targeted for functional annotation, in relation to male paternity success, in future studies.

ACKNOWLEDGMENTS

This work was supported by an NSERC Individual Discovery grant (to A.C.). S.F. was partially funded by a Manitoba Graduate Scholarship (MGS). We thank Reid Miller for help with the collection of data.

LITERATURE CITED

- Avila, F. W., and M. F. Wolfner, 2009 Acp36DE is required for uterine conformational changes in mated *Drosophila* females. Proc. Natl. Acad. Sci. USA 106: 15796–15800.
- Avila, F. W., L. K. Sirot, B. A. LaFlamme, C. D. Rubinstein, and M. F. Wolfner, 2011 Insect seminal fluid proteins: identification and function. Annu. Rev. Entomol. 56: 21–40.

- Britch, S. C., E. J. Swartout, D. D. Hampton, M. L. Draney, J. Chu et al., 2007 Genetic architecture of conspecific sperm precedence in Allonemobius fasciatus and A. socius. Genetics 176: 1209–1222.
- Civetta, A., 1999 Direct visualization of sperm competition and sperm storage in *Drosophila*. Curr. Biol. 9: 841–844.
- Civetta, A., H. M. Waldrip-Dail, and A. G. Clark, 2002 An introgression approach to mapping differences in mating success and sperm competitive ability in *Drosophila simulans* and *D. sechellia*. Genet. Res. 79: 65–74.
- Clark, A. G., 2002 Sperm competition and the maintenance of polymorphism. Heredity 88: 148–153.
- Clark, A. G., M. Aguadé, T. Prout, L. G. Harshman, and C. H. Langley, 1995 Variation in sperm displacement and its association with accessory gland protein loci in *Drosophila melanogaster*. Genetics 139: 189– 201.
- Clark, A. G., D. J. Begun, and T. Prout, 1999 Female x male interactions in Drosophila sperm competition. Science 283: 217–220.
- Dixon, S. M., J. A. Coyne, and M. A. Noor, 2003 The evolution of conspecific sperm precedence in *Drosophila*. Mol. Ecol. 12: 1179–1184.
- Fiumera, A. C., B. L. Dumont, and A. G. Clark, 2005 Sperm competitive ability in *Drosophila melanogaster* associated with variation in male reproductive proteins. Genetics 169: 243–257.
- Fiumera, A. C., B. L. Dumont, and A. G. Clark, 2007 Associations between sperm competition and natural variation in male reproductive genes on the third chromosome of *Drosophila melanogaster*. Genetics 176: 1245– 1260.
- Gilchrist, A. S., and L. Partridge, 1997 Heritability of pre-adult viability differences can explain apparent heritability of sperm displacement ability in *Drosophila melanogaster*. Proc. Biol. Sci. 264: 1271–1275.
- Giot, L., J. S. Bader, C. Brouwer, A. Chaudhuri, B. Kuang et al., 2003 A protein interaction map of Drosophila melanogaster. Science 302: 1727–1736.
- Greenspan, L., and A. G. Clark, 2011 Associations between variation in X chromosome male reproductive genes and sperm competitive ability in *Drosophila melanogaster*. Int. J. Evol. Biol. 2011: 214280.
- Howard, D. J., P. Gregory, J. Chu, and M. Cain, 1998 Conspecific sperm precedence in an effective barrier to hybridization between closely related species. Evolution 52: 511–516.
- Larson, E. L., G. L. Hume, J. A. Andrés, and R. G. Harrison, 2012 Postmating prezygotic barriers to gene exchange between hybridizing field crickets. J. Evol. Biol. 25: 174–186.
- Levesque, L., B. Brouwers, V. Sundararajan, and A. Civetta, 2010 Third chromosome candidate genes for conspecific sperm precedence between *D. simulans* and *D. mauritiana*. BMC Genet. 11: 21.
- Lüpold, S., S. Pitnick, K. S. Berben, C. S. Blengini, J. M. Belote *et al.*, 2013 Female mediation of competitive fertilization success in *Dro-sophila melanogaster*. Proc. Natl. Acad. Sci. USA 110: 10693–10698.
- Manier, M. K., J. M. Belote, K. S. Berben, D. Novikov, and W. T. Stuart, et al., 2010 Resolving mechanisms of competitive fertilization success in Drosophila melanogaster. Science 328: 354–357.
- Manier, M. K., J. M. Belote, K. S. Berben, S. Lüpold, O. Ala-Honkola *et al.*, 2013a Rapid diversification of sperm precedence traits and processes among three sibling *Drosophila* species. Evolution 67: 2348–2362.

- Manier, M. K., S. Lüpold, S. Pitnick, and W. T. Starmer, 2013b An analytical framework for estimating fertilization bias and the fertilization set from multiple sperm-storage organs. Am. Nat. 182: 552–561.
- Manier, M. K., S. Lüpold, J. M. Belote, W. T. Starmer, K. S. Berben *et al.*, 2013c Postcopulatory sexual selection generates speciation phenotypes in *Drosophila*. Curr. Biol. 23: 1853–1862.
- Martin, A., K. J. McCulloch, N. H. Patel, A. D. Briscoe, L. E. Gilbert *et al.*, 2014 Multiple recent co-options of Optix associated with novel traits in adaptive butterfly wing radiations. Evodevo 5: 7.
- Matsubayashi, K. W., and H. Katakura, 2009 Contribution of multiple isolating barriers to reproductive isolation between a pair of phytophagous ladybird beetles. Evolution 63: 2563–2580.
- Matute, D. R., and J. A. Coyne, 2010 Intrinsic reproductive isolation between two sister species of *Drosophila*. Evolution 64: 903–920.
- Moczek, A. P., and D. J. Rose, 2009 Differential recruitment of limb patterning genes during development and diversification of beetle horns. Proc. Natl. Acad. Sci. USA 106: 8992–8997.
- Mueller, J. L., J. R. Linklater, K. Ravi Ram, T. Chapman, and M. F. Wolfner, 2008 Targeted gene deletion and phenotypic analysis of the *Drosophila melanogaster* seminal fluid protease inhibitor Acp62F. Genetics 178: 1605–1614.
- Ou, C. Y., H. Pi, and C. T. Chien, 2003 Control of protein degradation by E3 ubiquitin ligases in *Drosophila* eye development. Trends Genet. 19: 382–389.
- Price, C. S., 1997 Conspecific sperm precedence in *Drosophila*. Nature 388: 663–666.
- Price, C. S., C. H. Kim, J. Posluszny, and J. A. Coyne, 2000 Mechanisms of conspecific sperm precedence in *Drosophila*. Evolution 54: 2028–2037.
- Price, C. S. C., C. H. Kim, C. J. Gronlund, and J. A. Coyne, 2001 Cryptic reproductive isolation in the *Drosophila simulans* species complex. Evolution 55: 81–92.
- Rogers, G. C., N. M. Rusan, D. M. Roberts, M. Peifer, and S. L. Rogers, 2009 The SCF Slimb ubiquitin ligase regulates Plk4/Sak levels to block centriole reduplication. J. Cell Biol. 184: 225–239.
- Rugman-Jones, P. F., and P. E. Eady, 2007 Conspecific sperm precedence in *Callosobruchus subinnotatus* (Coleoptera: Bruchidae): mechanisms and consequences. Proc. Biol. Sci. 274: 983–988.
- Sagga, N., and A. Civetta, 2011 Male-female interactions and the evolution of postmating prezygotic reproductive isolation among species of the *Virilis* subgroup. Int. J. Evol. Biol. 2011: 485460.
- Tyler, F., X. A. Harrison, A. Bretman, T. Veen, R. Rodríguez-Muñoz et al., 2013 Multiple post-mating barriers to hybridization in field crickets. Mol. Ecol. 22: 1640–1649.
- Wong, A., S. N. Albright, J. D. Giebel, K. R. Ram, S. Ji et al., 2008 A role for Acp29AB, a predicted seminal fluid lectin, in female sperm storage in Drosophila melanogaster. Genetics 180: 921–931.
- Yeh, S. D., T. Do, M. Abbassi, and J. M. Ranz, 2012 Functional relevance of the newly evolved sperm dynein intermediate chain multigene family in *Drosophila melanogaster* males. Commun. Integr. Biol. 5: 462–465.

Communicating editor: R. Kulathinal