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Molecular evolution of non-fertilizing sperm in Lepidoptera suggests minimal direct involvement in sperm competition — Source link \square

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27 Abstract

28 Sperm are among the most variable cells in nature. Some of this variation results from non-adaptive 29 errors in spermatogenesis, but many species consistently produce multiple sperm morphs, the adaptive 30 significance of which remains unknown. Here, we investigate the evolution of dimorphic sperm in 31 Lepidoptera, the butterflies and moths. Males of this order produce both fertilizing sperm and a 32 secondary, non-fertilizing type that lacks DNA. Previous organismal studies suggested a role for non-33 fertilizing sperm in sperm competition, but this hypothesis has never been evaluated from a molecular 34 framework. We combined published datasets with new sequencing in two species, the monandrous 35 Carolina sphinx moth and the highly polyandrous monarch butterfly. Based on population genetic 36 analyses, we see evidence for increased adaptive evolution in fertilizing sperm, but only in the 37 polyandrous species. This signal comes primarily from a decrease in non-synonymous polymorphism in 38 sperm proteins compared to the rest of the genome, suggesting stronger purifying selection, consistent 39 with selection via sperm competition. Non-fertilizing sperm proteins, in contrast, do not show an effect of mating system and do not appear to evolve differently from the background genome in either species, 40 41 arguing against the involvement of non-fertilizing sperm in direct sperm competition. Based on our 42 results and previous work, we suggest that non-fertilizing sperm may be used to delay female remating 43 in these insects and decrease the risk of sperm competition rather than directly affect its outcome.

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53 Introduction

54 Sperm cells display remarkable diversity throughout the animal kingdom (Pitnick, Hosken, & Birkhead, 55 2009), from small and plentiful to gigantic and few (Pizzari, 2006) to super-structure-forming (Higginson, 56 Miller, Segraves, & Pitnick, 2012). This variation exists at every level, from fixed differences between 57 species to variability within individual males (John Buckland-Nicks, 1998; Marks, Biermann, Eanes, & 58 Kryvi, 2008; Sasakawa, 2009; Swallow & Wilkinson, 2002; Tavares-Bastos, Teixeira, Colli, & Báo, 2002). In 59 many independently evolved cases, males consistently produce two different sperm types, a 60 phenomenon known as sperm dimorphism. In all cases examined, only one of the two sperm morphs is capable of fertilization (Bressac et al., 1991; Carcupino, Baldacci, Fausto, Scapigliati, & Mazzini, 1999; 61 62 Eckelbarger, Young, & Cameron, 1989; Sasakawa, 2009; Wilms, 1986). The evolutionary causes and 63 consequences of variation in sperm morphology, both within and between morphs, are immediately 64 intriguing. As gametes, these cells are the final step in the long chain of events leading to reproductive 65 success or failure. Why should such important components of fitness be so variable?

66 Much of this morphological diversity within morphs can be attributed to deleterious variation, *e.g.* 67 genetic defects (Chenoweth, 2005) or age-related decline in sperm quality (Preston, Saint Jalme, Hingrat, 68 Lacroix, & Sorci, 2015). This deleterious variation has been shown to be inversely correlated with rates 69 of sperm competition between species; taxa that experience more sperm competition tend to have less 70 morphologically variable sperm at both population and individual levels (Kleven, Laskemoen, Fossøy, 71 Robertson, & Lifjeld, 2008). In other words, sperm often vary in spite of constraint imposed by their 72 reproductive importance. In species with high rates of polyandry, postcopulatory selection through 73 sperm competition and cryptic female choice weeds out the suboptimal sperm variants, at least for 74 fertilizing sperm (Birkhead, 1998; Immler, Calhim, & Birkhead, 2008).

75 Production of multiple sperm morphs, conversely, is often posited to be adaptive in some way. The very 76 fact that sperm dimorphism has repeatedly evolved suggests that it has some fitness benefit. Most 77 commonly, non-fertilizing sperm in dimorphic systems are proposed to be specialized agents of male-78 male competition, acting as final combatants in the struggle for reproductive success (J Buckland-Nicks, 79 Bryson, Hart, & Partridge, 2010; John Buckland-Nicks, 1998; Swallow & Wilkinson, 2002). Indeed, some 80 have suggested that sperm dimorphism allows specialization in the non-fertilizing sperm for a 81 competitor-inhibiting function, sometimes called "kamikaze sperm" (Baker & Bellis, 1989). Although this 82 hypothesis has fallen out of favor, it was proposed and mainly evaluated in the context of mammalian 83 sperm (A. Harcourt, 1991; A. H. Harcourt, 1989; Moore, Martin, & Birkhead, 1999), where non-fertilizing 84 sperm are not usually differentiated from fertilizing sperm in a sophisticated way.

85 One of the most extreme cases of sperm dimorphism occurs in butterflies and moths (Lepidoptera). In 86 nearly all species of this order, males produce both fertilizing (eupyrene) sperm and a second type 87 (apyrene) that lacks a nucleus and nuclear DNA (Meves, 1902). The function of apyrene sperm is poorly 88 understood, but because it lacks DNA, it is clearly incapable of fertilizing eggs. Nevertheless, it does not 89 appear to be the result of errors in spermatogenesis; apyrene sperm production is hormonally regulated 90 and occurs in a developmentally predictable way, implying a novel gain of function in these insects 91 (Friedlander, 1997). Organismal studies have demonstrated that males can control the ratio of the two 92 sperm types in their ejaculate and typically transfer to females 10 to 20 times as much apyrene sperm as 93 eupyrene sperm, depending in part on the female's past mating history (Oberhauser, 1988). These 94 observations have led some to suggest that apyrene sperm play a specialized role in sperm competition 95 (Silberglied, Shepherd, & Dickinson, 1984), yet there remain several other competing hypotheses for 96 apyrene sperm function that have not been resolved through organismal observations and experiments 97 (Swallow & Wilkinson, 2002).

Recently, characterizations of the proteins found in lepidopteran sperm has opened a new avenue to 98 99 assess their evolution and function (Whittington et al., 2017; Whittington, Zhao, Borziak, Walters, & 100 Dorus, 2015). Proteomic studies have revealed distinct protein profiles for these two cell types 101 (Whittington, Karr, Mongue, Walters, & Dorus, in press). In both morphs, these proteins are retained 102 through maturation, and, in the case of apyrene sperm, the discarding of the nucleus. Because distinct 103 cellular functions are ultimately the product of their expressed protein complement, the class of 104 proteins uniquely found in apyrene sperm make logical targets for understanding the function of these 105 cells from a molecular perspective.

106 At the molecular level, sperm and other reproductive proteins are often observed to evolve rapidly 107 (Civetta & Singh, 1995; Dorus, Evans, Wyckoff, Sun, & Lahn, 2004; Willie J. Swanson & Vacquier, 2002). 108 For certain reproductive proteins, like sperm-egg interaction pairs, there is compelling evidence that 109 adaptive co-evolution drives this accelerated change (Herberg, Gert, Schleiffer, & Pauli, 2018; W J 110 Swanson & Vacquier, 1998). Yet there are also many instances of reproductive proteins that diverge 111 quickly because of relaxed purifying selection owing to expression in a single sex instead of the whole 112 population (Barker, Demuth, & Wade, 2005; Wade, Priest, & Cruickshank, 2008). Many other factors, 113 including number of protein-protein interactions or importance of reproductive role, can also act to 114 shape the intensity of positive or purifying selection on reproductive proteins (Schumacher, Rosenkranz, 115 & Herlyn, 2014; Schumacher, Zischler, & Herlyn, 2017). Recent theoretical work has formalized the

prediction that strong purifying selection on sperm proteins should depend on high rates of polyandry to generate sperm competition (Dapper & Wade, 2016). Thus, with the appropriate datasets, the degree of each sperm morph's role in sperm competition can be assessed via molecular tests of evolution.

119 In this study, we report the first molecular evolutionary analyses of dimorphic sperm. We assessed 120 patterns of both polymorphism and divergence among sperm proteins from both eupyrene and apyrene 121 sperm using proteomic datasets of two species: the monarch butterfly, Danaus plexippus, and the 122 Carolina sphinx moth, Manduca sexta (Whittington et al., in press). North American monarchs spend 123 time at incredibly high density in overwintering colonies in Mexico and California (Urguhart, 1976) and, 124 owing to these unique population dynamics, have some of the highest female remating rates observed 125 in Lepidoptera. Female monarchs mate an average of 2.6 times (and up to 14 times) in overwintering 126 colonies in the wild (Hill Jr., Wenner, & Wells, 1976; Smith, 1984), creating ample opportunity for sperm 127 competition. In contrast, Carolina sphinx moths are typically monandrous (Snow et al., 1974), making 128 sperm competition rarely relevant as a selective force. Taking advantage of this contrast, we investigate 129 the differences in patterns of selection between the two sperm morphs in each species to assess the 130 role of apyrene sperm in sperm competition. If apyrene sperm are involved in sperm competition, their 131 proteins should show evidence of stronger purifying selection in the monarch butterfly. To complete 132 these analyses, we have generated the first published set of whole-genome resequencing data for 133 Manduca sexta from a wild population. To test the general predictions for relaxed selection in sex-134 limited proteins, we used RNA-seq gene expression datasets from previously published data for Carolina 135 sphinx moths (Cao & Jiang, 2017) and newly generated data for the monarch butterfly.

136 Materials and Methods

137 Sources of data

138 We used gene sets from the published genomes of each species (Kanost et al., 2016; Zhan & Reppert, 139 2013) with sperm genes identified from their respective proteomes (Whittington, Karr, Mongue, Walters, 140 & Dorus, in press). We inferred selection from patterns of polymorphism and divergence from 141 congeners using whole genome Illumina resequencing data for both species: a previously published 142 dataset for North American monarch butterflies (Zhan et al., 2014) and a new dataset of North 143 Carolinian sphinx moths. Focal moths were collected with a mercury vapor light trap in July of 2017 in 144 Rocky Mount, North Carolina (see supplemental table S1 for sequencing summary statistics and 145 accessions). Divergences were called by comparison to the queen butterfly (Danaus gilippus, previously

146 published in Zhan et al. (2014)) for monarchs, and the five-spotted hawkmoth (Manduca

147 *quinquemaculata*, sequenced for this project) for the Carolina sphinx moth.

148 In both focal species, we used twelve wild-caught individuals for sampling of polymorphism. In the case 149 of Carolina sphinx moths, these were twelve males caught over the course of three nights. The sex-150 biased sampling reflects a sex bias in dispersal and collection at the light trap. In the case of monarchs, 151 samples were selected based on depth of sequencing coverage in the published dataset and included 8 152 females and 4 males from the panmictic North American migratory population. This mixed-sex sampling 153 added the complication of unequal sampling between the autosomes (n = 24) and Z sex chromosome (n = 24)154 = 16). Despite the male-biased gene accumulation on the Z chromosome, the vast majority of sperm 155 genes (92% in the Carolina sphinx, 90% in the monarch) are autosomal in both species (Mongue & 156 Walters, 2017). Due to the sampling complication and limited inference to be gained from Z-linked 157 genes, we focused on the autosomal genes in both species in subsequent analyses.

158 SNP-based methods

- 159 We aligned sequenced reads with bowtie2 (Langmead & Salzberg, 2012) for conspecifics to their
- reference genome or with stampy (Lunter & Goodson, 2011) with an increased allowance for
- substitution for heterospecific alignments. Alignments were taken through GATK's best practices
- pipeline (McKenna et al., 2010), including hard filtering, to yield a set of high quality variants both within
- 163 and between species. Effect-class of each variable site (synonymous, non-synonymous, intergenic, etc.)
- 164 was determined using custom databases for the two species created with SnpEff (Cingolani et al., 2012).
- 165 Annotated SNPs were curated to remove false divergences (ancestral polymorphism) and then
- differences in adaptive evolution were calculated using an estimator of the neutrality index to calculate
- 167 α, the proportion of substitutions driven by adaptive evolution (Stoletzki & Eyre-Walker, 2011). This
- 168 form of α corrects the inherent bias in a ratio of ratios while also allowing summation across multiple
- 169 genes to reduce noise associated with small numbers in count data. For any set of *i* genes with non-zero
- 170 counts of synonymous (s) polymorphism (P) and divergence (D):

$$\alpha = 1 - \frac{\sum (Dsi * Pni)/(Psi + Dsi)}{\sum (Dni * Psi)/(Psi + Dsi)}$$

171 This statistic was calculated with custom scripts in R (R Core Team, 2017).

172 Assessment of adaptive evolution and statistical significance

173 In each analysis, we calculated α for a biologically meaningful set of genes, *e.q.* the sperm proteome and 174 the background genome, and generated a test statistic from the absolute difference of the two point-175 estimates. To determine significance, we combined the two sets and randomly assigned genes into two 176 new sets of sizes equal to the originals. The difference of these two datasets was determined and the 177 process was repeated for 50,000 permutations to build a distribution of differences between the point 178 estimates of two gene sets of these relative sizes. The p-value was taken as the proportion of times a 179 greater absolute difference was observed between the two random data sets than between the original 180 sets.

181 We used this permutation approach to make within-species comparisons of α for several different 182 groupings of genes. We first examined differences between the whole sperm proteome and background 183 genome (i.e. all autosomal non-sperm proteins). Next, we considered differences between sperm 184 homologs and sperm proteins unique to one species to assess how selection acted on the same genes in 185 different species. We identified sperm homologs as predicted orthologs that are present in the sperm of 186 both species, with orthology predicted via the proteinOrtho pipeline, as previously reported in Mongue 187 & Walters (2017). Unique sperm proteins may or may not have an ortholog in the other species but are 188 present in the sperm of only one species. Finally, we compared among proteins grouped by their 189 presence in apyrene versus eupyrene sperm. To do so, we classified sperm proteins into three subsets: 190 specific to eupyrene sperm, specific to apyrene sperm, or shared in both types. Pairwise comparisons 191 were made between each subset. For these analyses, we did not consider orthology status owing to the 192 reduction in power that would accompany multiple layers of subdivision of the dataset. For the whole 193 proteome and morph subset comparisons, we further assessed the relative contributions of 194 synonymous and non-synonymous polymorphism and divergence to the α calculation, using a Wilcoxon-195 Mann-Whitney test to assess significant differences.

196 Site-frequency-based methods

We also investigated molecular evolution by leveraging site-frequency-spectrum-based approaches as complimentary evidence. Owing to the redundancy in results, we have included these analyses in the supplement rather the main text. In brief, we used the population genetics software suite ANGSD (Korneliussen, Albrechtsen, & Nielsen, 2014) to generate site frequency spectra at putatively neutral (four-fold degenerate) and selected (zero-fold-degenerate) sites in the genome. We unfolded site frequency spectra and analyzed these spectra with the software polyDFE (Tataru, Mollion, Glémin, & Bataillon, 2017) to examine rates of adaptive evolution in the whole sperm proteomes and background

genomes with a more complex likelihood model that corrects for effects of demography and potentialmisattribution of ancestral state.

206 Investigation of sex-limited and tissue-specific expression

207 Next, we used RNA-seq data to assess whether or not differences in tissue specificity of expression 208 impacted our results from the sperm proteomes in these taxa. For *Manduca sexta*, there already existed 209 a wealth of tissue-specific data at multiple developmental timepoints (Cao & Jiang, 2017). Because we 210 were primarily interested in sperm involvement, we focused on data from adult males, specifically RNA 211 from the testes, head, thorax, and gut. Expression (measured as fragments per kilobase of transcript per 212 million mapped reads, FPKM) was averaged across biological replicates where available in this species. 213 Monarchs had no comparable published data, so we generated separate RNA-seq data sets from the 214 head, thorax, gut, testes, and accessory gland of three adult males (summarized in Table S2 with 215 accessions).

216 We quantified tissue-specificity of expression using the *specificity metric* (SPM) statistic, a ratio ranging 217 from 0 to 1 indicating the proportion of gene expression occurring in a given focal tissue (Kryuchkova-218 Mostacci & Robinson-Rechavi, 2017). For instance, a gene with SPM = 0.8 for the testes shows 80% of its 219 total expression across all sampled tissues in the testes. This same gene would have a much lower SPM 220 value in head, thorax, or other tissues. We observed a bimodal distribution of tissue specificities, which 221 allowed us to bin genes into one of two classes: those that displayed low levels of specificity (SPM < 0.5) 222 and those that displayed high levels (SPM > 0.5). After separating genes by specificity, we calculated α 223 for three classes of genes in these two specificity bins.

224 We had two goals with these analyses: (1) to determine if patterns of adaptive evolution between 225 classes remained the same at both low- and high-specificities and (2) if α increased within a class of 226 genes at higher specificity compared to low. First, we considered background genome genes (i.e. non-227 sperm genes) ranked by maximum specificity observed in the head, thorax, or gut for each of these 228 genes. Next, we considered only genes identified in the sperm proteome and ranked them by SPM in the 229 testes. Finally, for putatively male-limited non-sperm genes, we excluded sperm proteome genes and 230 considered again those ranked by specificity in the testes (or testes and accessory glands for monarchs). 231 As with our other α calculations, we used non-parametric bootstrapping to generate 95% confidence 232 intervals. For cases in which confidence intervals overlapped, we assessed significance with permutation 233 testing. These analyses were completed with custom R scripts.

234 Demographic estimates

Finally, to contextualize the previous analyses and take full advantage of our newly-generated data, we characterized present and historical population sizes of our study species from genomic data. Using folded four-fold degenerate site frequency spectra, we estimated neutral coalescence patterns with Stairway Plot (Liu & Fu, 2015). For estimated generation time, we used four generations per year for monarchs and three for the Carolina sphinx moth. For mutation rate, we chose the estimate 2.9*10⁻⁹ from the butterfly *Heliconius melpomene*, the closest relative with a spontaneous mutation rate estimate (Keightley et al., 2015).

242 Results

243 Differences Between Sperm Proteins and the Background Genome

244 First, we considered the sperm proteome as a whole (*i.e.* all apyrene, shared, and eupyrene proteins) 245 and compared adaptive evolution of genes found in sperm to those in the background genome, defined 246 as all autosomal protein coding genes not present in the sperm proteome. Z-linked genes were excluded 247 from the analysis. We counted and classified synonymous and non-synonymous single nucleotide 248 polymorphisms within species and divergences to a congener (Danaus gilippus for the monarch, and 249 Manduca quinquemaculata for the Carolina sphinx). These quantities were used to generate an estimate 250 of the proportion of adaptive substitutions (α) per gene-class for both the sperm proteome and the 251 background genome. We found no difference in α between the sperm proteome and the rest of the 252 genome in the Carolina sphinx (p = 0.40892 by permutation testing, Figure 1A, left); for monarchs, 253 however, the sperm proteome showed a significantly greater proportion of adaptive substitutions than 254 the rest of the genome (p = 0.00006, Figure 1A, right). Note that in the strict sense, negative α values are 255 not biologically meaningful and likely point to an abundance of weakly deleterious variants within 256 populations or complex demographic histories (Eyre-Walker & Keightley, 2009); nevertheless, these 257 confounding variables should not differentially affect genes within species, so our observed differences 258 point to true differences in selection in gene sets.

To better understand the relative roles of polymorphism and divergence in sperm and background
genes, we investigated the individual components of α: counts of non-synonymous polymorphism (Pn),
synonymous polymorphism (Ps), non-synonymous divergence (Dn), and synonymous divergence (Ds).
We compared the scaled estimates of each (*e.g.* non-synonymous polymorphisms per non-synonymous
site) to the background genome within each species using a Wilcoxon-Mann-Whitney test (Figure 1B).

We found no differences between sperm and the background for any class of variants in *M. sexta* (Pn: W = 3014100, p = 0.5964; Ps: W = 2879300, p = 0.1830; Dn: W = 3068300, p = 0.2009; Ds: W = 2895700, p = 0.2686). The signal for elevated α in monarch sperm primarily reflects non-synonymous polymorphism, which was greatly depressed (W = 3062400; p = 3.224 * 10⁻¹¹), as would be expected under strong purifying selection, while other classes were comparable between sperm and the background genome (Ps: W = 2684200, p = 0.2720; Dn: W = 2506400, p = 0.1300; Ds: W = 2544400, p = 0.3437).

270 Next, we leveraged orthology, as established by Whittington et al. (2017), to test for differences in 271 mating system while controlling for the effects of sperm proteome content. Substantial numbers of 272 orthologous proteins are found in the sperm proteomes of both species, which we hereafter referred to 273 as sperm homologs. Sperm homologs offer the opportunity to directly assess the selective pressures 274 experienced by the same genes with putatively conserved function but found in species with different 275 levels of postcopulatory selection. Nearly half of the monarch sperm proteome (~42%, 216 genes, Figure 276 2A) shares an ortholog in the sperm proteome of *M. sexta*; reciprocally, there are 236 genes (37%) in the 277 Carolina sphinx sperm proteome that share an ortholog in the monarch sperm proteome; these 278 numbers are not equal due to lineage-specific duplications among sperm homologs creating a few cases 279 of one-to-many orthology. We tested for differences in adaptive evolution between sperm homologs 280 and sperm proteins unique to one species (orthology outside of sperm or no detectable orthology). In 281 Carolina sphinx moths, genes of these two classes did not differ in the proportion of adaptive 282 substitutions with permutation testing (p = 0.6174, Figure 2B). In monarchs, we detected an increased 283 proportion of adaptive substitution in the sperm homologs compared to unique proteins (p = 0.0372, 284 Figure 2B). Comparing between species, sperm homologs had much higher α values in monarchs than in 285 Carolina sphinx moths (p = 0.00008), while genes with unique expression in either species did not show 286 differences between species (p = 0.5922). Thus, the same sperm proteins appear to be evolving more 287 adaptively in the polyandrous species.

288 Site-frequency based methods

We also took a likelihood approach to modeling adaptive evolution using site frequency spectra
generated from the same samples we used for SNP-counting. These results are detailed in the
supplement. In short though, we found a shift in the predicted distribution of fitness effects of new
mutations in monarch sperm proteins compared to the background consistent with stronger purifying
selection (Figure S1) and drastically higher α in sperm genes in monarchs alone (Figure S2).

294 Patterns of Adaptive Evolution in Sex-Specific Tissues

295 Next, we used RNA-seq data to examine the effect of tissue-specificity on selection in these insects. 296 With these data, we calculated the tissue specificity metric, SPM (Kryuchkova-Mostacci & Robinson-297 Rechavi, 2017), which ranges from ubiquitous expression (near 0) to single-tissue specific (1). Although 298 the sperm proteomes of both of our species were enriched for gene products specifically expressed in 299 testes, they also contained broadly expressed gene products (Figure 3B). To assess the effect of these 300 broadly expressed genes on our inference of selection, we recalculated the α statistic for two bins of 301 genes (Figure 3C): those with broad expression (SPM < 0.5) and those with high tissue-specificity (SPM > 302 0.5).

303 In Carolina sphinx moths, there were no significant changes in α between low- and high-specificity genes 304 in any part of the genome (background genes: p = 0.3868, sperm proteome genes: p = 0.3248, male-305 limited genes: p = 0.5579; Figure 3C, left), nor did any of the gene classes differ from each other within a 306 specificity bin. In monarchs, however, both sperm proteome genes (p = 0.0242) and testes genes (p =307 0.0137) showed higher α in the high-specificity group than the low-specificity group, though somatically 308 expressed genes did not (p = 0.6831). Additionally, we found that sperm genes showed much greater α 309 than the background genome or other genes expressed in the testes at both low- and high-specificities 310 (Figure 3C, right). This result indicates that our initial results (considering the whole sperm proteome) 311 are not dependent on the underlying specificity of sperm genes.

312 Molecular evolution in dimorphic sperm

313 Having verified the patterns of evolution in the whole sperm proteomes with several approaches, we 314 turned to our primary question, assessing apyrene sperm function through analysis of molecular 315 evolution. We considered the different subsets of the sperm proteomes based on the two sperm types. 316 The two datasets consisted of three classes of sperm proteins: unique to eupyrene sperm, unique to 317 apyrene sperm, or found in both cell types (henceforth "shared", Figure 4A). We assessed differences in 318 selective pressures between the sperm morphs with another series of permutation tests, both 319 comparing parts of the sperm proteome to the background genome and comparing parts of the 320 proteome to each other.

- 321 As expected based on the whole-proteome results from Carolina sphinx moth, neither eupyrene-specific
- 322 (p = 0.55912), shared (p = 0.4647), nor apyrene-specific proteins (p = 0.96496) differed from the
- background genome (Figure 4B). α did not vary between apyrene-specific and eupyrene-specific

proteins (p = 0.7271), between apyrene-specific and shared (p = 0.7176) or eupyrene-specific and shared proteins (p = 0.9979). In monarchs, both eupyrene-specific proteins (p = 0.00018) and shared proteins (p = 0.01038) showed elevated α , but apyrene-specific proteins did not evolve differently from the background genome (p = 0.55934). Neither apyrene nor eupyrene sperm differed significantly from the shared set in monarchs (p = 0.6332 & p = 0.6234, respectively), but there was a trend towards significantly increased α in eupyrene-specific proteins compared to apyrene-specific proteins (p = 0.0986).

- As with the whole sperm proteome, we investigated which classes of variants contributed to our
- observed differences in α (Figure 4C). Consistent with the results above, none of the variant classes
- 333 significantly differed from the genome background in the sphinx moth eupyrene-specific proteins (Pn: W
- 334 = 995190, p = 0.0857; Ps: W = 943550, p = 0.6782; Dn: W = 966630, p = 0.3183; Ds: W = 963410, p =
- 0.3596). Shared proteins also showed the same level of variation as the background across all variants
- 336 (Pn: W = 1470300, p = 0.3277; Ps: W = 1444400, p = 0.1369; Dn: W = 1540700, p = 0.6883; Ds: W =
- 337 1437100, p = 0.1030). And finally, apyrene-specific proteins were not significantly different either (Pn: W
- 338 = 548570, p = 0.4974; Ps: W = 491410, p = 0.2149; Dn: W = 560910, p = 0.2741; Ds: W = 495180, p =
- 0.2653). In summary, there was no evidence for stronger selection on either sperm morph in Carolina
- 340 sphinx moths.

341 For monarchs, we found that the elevated α in the eupyrene-specific and shared subsets was driven 342 primarily by a decrease in non-synonymous polymorphism compared to the background genome (W = 1291700, p = 0.0003 for eupyrene; W = 1486100, $p = 1.167*10^{-8}$ for shared). Apyrene-specific proteins 343 344 did not show a reduction in non-synonymous polymorphism (W = 284620, p = 0.1684). Synonymous 345 polymorphism did not significantly differ from the background in any subset of the sperm proteome (eupyrene: W = 1164200, p = 0.4492; shared: W = 1249900, p = 0.5570, apyrene: W = 270160, p = 346 347 0.4927). Nor did synonymous divergence (eupyrene: W = 1056000, p = 0.0928; shared: W = 1209000, p = 0.0928; shared: W = 0.0928348 0.7665, apyrene: W = 279420, p = 0.2594). Intriguingly, non-synonymous divergence was elevated

- compared to the background in eupyrene-specific proteins (W = 1021800, p = 0.0151), but not the
- shared (W = 121800, p = 0.9185) or apyrene-specific portions of the proteome (W = 266580, p = 0.6042).
- 351 This suggests periodic sweeps of positively selected variants in fertilizing sperm proteins.

We did not examine orthology within dimorphic sperm owing to small gene counts giving reduced statistical power. Nor could we could examine tissue specificity here because apyrene and eupyrene sperm are produced at different developmental timepoints and we did not have suitable expression data in both species. Nonetheless, the consistency of results in the whole proteome datasets gives us no
 reason to expect that within-proteome results would be idiosyncratic to our methodology.

357 *Demographic estimates*

358 Finally, to contextualize our results with population dynamics, we estimated population size history 359 using site frequency from 4-fold degenerate sites in the two species' genomes (Figure S3). Both have 360 effective population sizes near 2,000,000, as expected of herbivorous invertebrates with high dispersal 361 potential, numerous host plants, and a large range over North America. We also recovered a population 362 size increase in monarch butterflies in the recent past, which has been previously reported with genomic 363 data (Zhan et al., 2014). We note that our inferred timing of this event differs from that of the previous 364 authors, who used mutation rate estimates from Drosophila melanogaster. Such input parameter 365 differences affect the estimated time of events, but not the trajectories.

366 Discussion

367 We investigated the molecular evolution of eupyrene (fertilizing) and apyrene (non-fertilizing) sperm, 368 the ubiquitous lepidopteran cell-type of unknown functional significance. These sperm have long been 369 posited to interfere with competitors' sperm, in part because their quantity varies with levels of male-370 male competition (Silberglied et al., 1984; Solensky & Oberhauser, 2009; Swallow & Wilkinson, 2002). In 371 contrast to these organismal observations, the results of our molecular analyses cast doubt on this 372 hypothesis. If apyrene sperm played an active role in sperm competition, we would expect evidence for 373 stronger selection in apyrene sperm compared to the background genome in monarchs. We found a 374 signal for elevated adaptive evolution (α) in the sperm proteome compared to the background genome 375 in these polyandrous butterflies, but this signal did not include apyrene-sperm-specific proteins. Instead, 376 genes encoding apyrene sperm proteins evolve similarly to the background genome in both monarchs 377 and Carolina sphinx moths. This result is unlikely to have arisen from a lack of power in our 378 methodologies, as eupyrene-specific and shared sperm proteins showed patterns in line with 379 expectations for a role of sperm competition in molecular evolution in monarchs.

380 Selection consistent with sperm competition, but only in fertilizing sperm

The source of the apparently elevated α in the monarch sperm proteome came mainly from a dearth of non-synonymous polymorphisms in sperm proteins compared to the background genome, indicating the action of purifying selection to remove many variants before fixation in monarchs. Strong purifying

selection has been similarly observed in genes expressed in pollen, the main male-male competitors in
flowering plants (Arunkumar, Josephs, Williamson, & Wright, 2013). A similar pattern can also be
observed in passerine birds, in which species with higher rates of sperm competition show less
intraspecific and intra-male variation in sperm length compared to sperm of less polyandrous species
(Immler et al., 2008; Kleven et al., 2008).

389 Moreover, the elevated α in sperm homologs in monarchs suggests that genes that have had conserved 390 sperm function since the divergence of the two species some 100 million years ago (Heikkila, Kaila, 391 Mutanen, Pena, & Wahlberg, 2012) are under stronger purifying selection in the polyandrous species. 392 According to recent gene ontology analyses, such genes are enriched for core traits in sperm, such as 393 mitochondrial function, respiration, and flagellar structure. Similarly, proteins shared between the two 394 sperm types and those unique to eupyrene sperm show an elevated α compared to the background 395 genome in monarchs. Sperm proteins shared between morphs are enriched for structural proteins that 396 give rise to the sperm tail and thus impact motility (Whittington et al., in press), while those expressed 397 only in eupyrene sperm doubtless include important mediators of fertilization. At the cellular level, 398 variation in sperm traits like swimming ability, longevity, and overall viability affects sperm competition 399 outcomes (Burness, Casselman, Schulte-Hostedde, Moyes, & Montgomerie, 2004; Kim et al., 2017) and 400 has a polygenic basis in other taxa (Hering, Olenski, & Kaminski, 2014). For traits like longevity and 401 motility there is a threshold below which fertilization becomes significantly impaired, but in the absence 402 of competitor alleles, there is a larger range of effectively-neutral trait-values, allowing for more 403 variation to be maintained in the population. In the presence of competitor alleles, however, marginal 404 differences in fertilization success come under selection, leading to the removal of deleterious variants 405 through sperm competition.

406 Stronger selection from competition may include even the event of fertilization itself. Lepidopteran eggs 407 are known to possess multiple micropyle openings for sperm (Kumar, Kariappa, Babu, & Dandin, 2007) 408 and eupyrene sperm possess structures resembling an acrosome (while their apyrene counterparts do 409 not) (Friedlander, 1997). This rare combination of male and female gamete structures is also found in 410 sturgeon, in which the multiple micropyles give several sperm potential access to the egg nucleus and 411 there is competition among sperm to initiate karyogamy via the acrosome reaction (Psenicka, Rodina, & 412 Linhart, 2010). Consistent with micropyle-mediated competition, it has been shown that more 413 polyandrous species of Lepidoptera tend to have more micropyles on their egg surfaces than 414 monandrous species (lossa, Gage, & Eady, 2016). If this truly does extend the opportunity for male-male

415 competition and cryptic choice, then acrosomal proteins in eupyrene sperm would be likely targets for416 selection in polyandrous systems.

Whatever the mechanics of fertilization are, paternity outcomes in polyandrous species are often bimodally distributed (Simmons & Siva-Jothy, 1998; Wedell & Cook, 1998), including in monarch butterflies (Mongue, Ahmed, Tsai, & De Roode, 2015). For females that mate twice, one of the two males typically fathers most, if not all, of the observed offspring produced by the female, but there is little consistency in whether it is the first or second male. With these dynamics, fitness differences between winning and losing sperm phenotypes are large and selection can reliably remove less successful genotypes.

424 Evidence of this can be seen in the estimated distribution of fitness effects of new mutations in monarch 425 sperm proteins. Compared to the background genome, we see a decrease in the proportion of 426 effectively neutral and weakly deleterious mutations and an increase in both strongly deleterious and 427 beneficial mutations. In the absence of competition, not only are mildly suboptimal variants effectively 428 neutral, but novel, more efficient competitors should have no selective advantage in monandrous 429 species unless they also markedly increase fitness in a single mating. This reasoning is supported by the 430 estimated distribution of fitness effect for the complimentary gene sets in the Carolina sphinx moth; in 431 this species, we see little variation in the DFE between the background genome and the sperm 432 proteome. Moreover, there is no decrease (and indeed) an increase in non-synonymous divergence of 433 eupyrene sperm proteins in monarchs compared to the rest of the genome. This pattern suggests that in 434 addition to strong purifying selection there must be periodic sweeps of beneficial alleles. Without a 435 broader, phylogenetically controlled study, these results between a single pair of species are not 436 conclusive, but they fit well with the prediction that sperm protein evolution depends on the rates of 437 polyandry in a species (Dapper & Wade, 2016).

438 Evolution of tissue-specific and male-limited genes

Other studies have demonstrated that tissue specificity of expression can strongly influence the
molecular evolution of reproductive proteins (Schumacher & Herlyn, 2018), in some cases more than
mating system (Carnahan-Craig & Jensen-Seaman, 2014). Because our proteomic data did not contain
information on tissue specificity on their own, we examined this dynamic with RNA-seq data. We found
increased adaptive evolution in monarch sperm genes with higher specificity compared to lowspecificity sperm genes. Furthermore, while not significantly different from background genome, α for

non-sperm genes expressed in the testes increased with greater specificity in monarchs, suggesting that
they too may be subject to stronger sexual selection in this polyandrous species. Neither of these
patterns held for Carolina sphinx moths, which showed no differences based on tissue specificity. This
consistency further suggests the difference in mating system as an explanation for differences in
intensity of selection.

450 Finally, we did not observe relaxed constraint in reproductive proteins predicted due to the smaller 451 effective population size of males or females compared to the population as a whole, as predicted by 452 theory (Dapper & Wade, 2016; Wade et al., 2008). Specifically, we did not observe a difference in the 453 adaptive evolution of genes with testes-specific expression, our proxy for sex-limited expression, 454 compared to the background genome. To explain this discrepancy between theory and observation, we 455 turn to Nearly Neutral Theory. Large populations have more efficient selection than small populations 456 and a smaller range of slightly deleterious mutations that behave neutrally (Ohta, 1992). Mutations with 457 a selective effect less than $1/N_e$ are expected to behave neutrally. For instance, one commonly cited 458 estimate for human population size is Ne \approx 10,000 over evolutionary history (Zhao et al., 2000). Based 459 on this, mutations with selective effects less than 0.0001 should behave neutrally for alleles expressed in 460 both sexes, while those with effects of 0.0002 are effectively neutral for alleles only expressed in one 461 sex. And indeed, there is evidence that genes expressed only in men have a higher mutational load than 462 those expressed in both sexes (Gershoni & Pietrokovski, 2014). Chimpanzees, another species with a 463 similar effective population size (Won & Hey, 2005), also show increased non-synonymous divergence in 464 reproductive proteins (Wong, 2010). Broadly, male reproductive protein evolution appears to depend 465 more on effective population sizes than intensity of sperm competition in the great apes in general 466 (Good et al., 2013), as one would expect for species with relatively small effective population sizes.

467 In contrast to mammals, the effective population sizes of most insect species are orders of magnitude 468 higher. Using neutral site frequency spectra, we estimated effective populations near 2,000,000 for both 469 North American monarchs and Carolina sphinx moths. Selection is much more effective in these massive populations; mutations with effects above 5*10⁻⁷ should be subject to selection in both sexes and those 470 471 above $1*10^{-6}$ should be subject to selection if expression is sex-limited. Thus, even selection on alleles 472 with sex-limited expression in these insects should be 100 times stronger than selection on the entire 473 human population. Even if there is a relative two-fold difference in selection, the absolute magnitude of 474 the difference should be miniscule, and the effects of mating system more apparent.

475 Advancing understanding of apyrene sperm

476 Previous morphological work found that eupyrene sperm traits (like sperm length) but not apyrene 477 sperm traits, varied with risk of sperm competition in other butterflies (Gage, 1994). Similarly, from a 478 molecular perspective, none of the patterns of increased purifying and positive selection that we 479 observed for monarch sperm proteins applied to the apyrene-specific proteins. That we also do not see 480 evidence for the action of sperm competition on apyrene-specific protein evolution is itself informative, 481 however. Research to-date has proposed four main hypotheses for apyrene sperm (Swallow & 482 Wilkinson, 2002): active sperm competition agents, passive competition agents, nutrient nuptial gifts, or 483 necessary facilitators of fertilization. Our molecular analyses argue against apyrene sperm as active 484 agents of sperm competition, but it is worth considering predictions for molecular evolution of apyrene 485 sperm under the other hypotheses.

486 Indeed, apyrene sperm may still have adaptive significance without specialized molecular function, 487 especially under the filler hypothesis. This proposed function also relates to sperm competition, but 488 posits that apyrene sperm are employed proactively, to fill the female's sperm storage organ and delay remating, thus decreasing the risk of sperm competition, rather than impacting its outcome (Swallow & 489 490 Wilkinson, 2002). Both in monarchs and the butterfly Pieris napi, female time to remating increases with 491 the number of apyrene sperm received from males (Cook & Wedell, 1999; Oberhauser, 1988). Such 492 observations are somewhat confounded by the size of the spermatophore nuptial gift that males 493 provide during mating, but apyrene sperm themselves have been proposed as a form of nutritional 494 nuptial gift (He, Tanaka, & Miyata, 1995; Lamunyon, 2000). Under both the nutrient and filler 495 hypotheses, the actual sequence of apyrene sperm proteins should be less important than their physical 496 presence and abundance, so factors affecting the rate of apyrene sperm production would be more 497 likely targets for selection in polyandrous species than the proteins sequences themselves.

498 Finally, apyrene sperm appear to capacitate fertilization in *Bombyx mori* (Takemura, Sahara, Mochida, & 499 Ohnuma, 2006); the mechanism here is unclear and the phenomenon is untested in other taxa, but it 500 could conceivably involve proteins that modulate female reproductive physiology to make conditions 501 more favorable for eupyrene sperm or induce oviposition. In such a case, these proteins would behave 502 more akin to the broader class of reproductive proteins and evolve independently of rates of polyandry 503 in a species. If there is an evolutionarily conserved capacitation effector in our study taxa, it is possible 504 that this function is governed by a small subset of apyrene-specific proteins. Because our methods 505 aggregate signal for selection across multiple genes or sites to counteract high variance in variant counts

within genes (Stoletzki & Eyre-Walker, 2011), the importance of one or a few genes could be lost in the
heterogeneous selection on different proteins.

508 Conclusions

509 Variation in reproductive traits has long been studied at the morphological and molecular level, 510 generally. Yet sperm dimorphism, one of the most striking and enigmatic reproductive traits, has not 511 previously been assessed using population genetic analyses. Our investigation of the sperm proteome in 512 two Lepidoptera demonstrates a pattern of stronger purifying selection on fertilizing-sperm genes in a 513 species with higher rates of sperm competition. In this polyandrous species, these genes experience a 514 strikingly different selective environment than the rest of the genome, with strong purifying selection 515 reducing variation in sperm genes. In contrast, fertilizing-sperm genes in the monandrous species hold 516 as much deleterious variation as other parts of their genome. Our new molecular findings fit well with 517 established studies on sperm morphology which show that sperm competition results in decreased 518 variation in sperm traits.

- 519 The evolution of non-fertilizing sperm, however, does not show a strong influence of sperm competition.
- 520 This lack of pattern itself argues against apyrene sperm as active agents of sperm competition, one of
- 521 the long-held hypotheses for non-fertilizing sperm function. Instead, apyrene sperm may play a passive
- role in reducing the risk of competition by delaying female remating. The method by which apyrene
- 523 sperm capacitate fertilization in some species remains unclear based solely on genomic approaches and
- will likely require functional experiments to completely understand.

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experiment design and manuscript editing.
Data accessibility: Manduca sexta whole genome resequencing data can be found on NCBI's Sequence
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Figure 1. A. In the Carolina sphinx moth (*M. sexta*), there is no difference between the sperm proteome

and the rest of the genome (left); conversely, genes in the sperm proteome of monarch butterflies (D.

plexippus) show a significantly higher proportion of adaptive substitutions (α) than the rest of the

767 genome (right). P-values come from permutation tests. Error bars represent 95% bootstrapped

confidence intervals from the point estimates. **B.** Decomposing α into its components: Pn, Ps, Pn, and Ds

and comparing the sperm proteome (filled boxes) to the background genome (open boxes). There were
 no strong differences between sperm genes and the background genome in Carolina sphinx moths. In

771 monarch butterflies, the signal for increased adaptive substitution comes from a marginal increase in

772 non-synonymous divergence (bottom left) combined with a great reduction in non-synonymous

polymorphism in sperm genes compared to the rest of the genome (top left). P-values reflect Wilcoxon-

774 Mann-Whitney tests, with * < 0.05, ** < 0.005, *** < 0.0005, etc.





B Adaptive evolution accounting for orthology in the sperm proteins

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Figure 2. A. Composition of the portion of the sperm proteomes analyzed in this study. Numbers
indicate counts of proteins unique to one species' sperm or with an ortholog in the other species' sperm
(sperm homologs). Note that the overlap number varies between species due to the presence of a few
one-to-many-orthologs. B. Sperm homologs show evidence for a greater proportion of adaptive
substitutions (α) in monarch butterflies, but not in Carolina sphinx moths. P-values are based on
permutation tests comparing the difference between two sets of genes randomly assigned from the
sperm proteome in each species; error bars are 95% bootstrap confidence intervals.

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787 Figure 3. Investigating how tissue specificity of gene expression impacts adaptive evolution in both 788 Carolina sphinx moths (left column) and monarchs (right column). A. Maximum specificity of all genes 789 across all studied tissues using the RNA-seq data considered in these analyses. **B.** Observed distribution 790 of specificity of sperm proteome genes expressed in the testes. Based on these distributions, we 791 separated genes into one of two categories, |ow-specificity (SPM < 0.5) or high-specificity (SPM > 0.5), 792 divided by the dashed line. C. Inferred proportion of adaptive substitutions (α) in background genes 793 (squares), sperm proteome genes (circles), and male-limited genes (as defined by testes expression). Bars represent 95% confidence intervals from non-parametric bootstrapping. Non-overlapping 794 795 confidence intervals imply significant differences generally, but we have also highlighted two significant 796 differences that are less visibly apparent. Monarchs show evidence for increasing α with increasing 797 tissue specificity in sperm and testes genes, but sphinx moths do not. Moreover, sperm proteome genes 798 evolve more adaptively than background or testes-specific genes in both specificity groups for monarchs 799 but not sphinx moths.



801 Figure 4. A. Composition of the sperm proteome with respect to dimorphic sperm. The majority of 802 identified proteins were shared between the two cell types, followed by the set unique to eupyrene 803 sperm, and finally the smallest set was the proteins found only in apyrene sperm. **B**. None of the sets of 804 sperm proteins evolved either differently from each other or distinctly from the background genome 805 (shaded regions represent 95% confidence intervals of the background) in the Carolina sphinx (left). In 806 the monarch however (right), the signal for elevated α was localized to the eupyrene-specific and shared 807 proteins. There was also a trend for increased α in eupyrene-specific proteins as compared to apyrene-808 specific. Error bars represent 95% confidence intervals from bootstrapping. **C**. Decomposing α into Pn, 809 Ps, Dn, and Ds for dimorphic sperm. Plotting of variation follows the coloring and order in parts A and B; 810 from left to right in each panel: background genome, eupyrene, shared, and apyrene sperm. Asterisks 811 denote significant differences from the background genome based on a Wilcoxon-Mann-Whitney test, 812 with * < 0.05, ** < 0.005, *** < 0.0005, etc.