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Honey bee males and queens use glandular secretions to enhance sperm viability before and after storage

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ABSTRACT

Internal fertilization requires live sperm to be transferred from male to female before egg fertilization. Both males and females assist the insemination process by providing sperm with glandular secretions, which have been inferred to contain subsets of proteins that maintain sperm viability. Here we show that in the honeybee (Apis mellifera) secretions of the male accessory glands, the major contributors towards seminal fluid, enhance sperm survival. We further demonstrate that the protein fraction of the male accessory gland secretion is indeed important for achieving the maximal effect on sperm survival. After sperm storage, the queens also provide sperm with secretions from spermathecal glands and we show that these secretions have a comparable positive effect on sperm viability. SDS gels show that the proteomic profiles of accessory gland secretion and spermathecal fluid secretion hardly overlap, which suggests that males and females use different proteins to enhance sperm viability during, respectively, ejaculation and final sperm storage.

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1. Introduction

Producing high quality ejaculates that remain viable during insemination is crucial for males to maximize their reproductive success (García-González and Simmons, 2005; Hunter and Birkhead, 2002), but how males actually influence the viability of their ejaculates remains unclear. In many species, males provide sperm with glandular secretions that are usually referred to as seminal fluid or seminal plasma, but details about the molecular composition of seminal fluid fractions are only available for a few insects such as fruit flies (Ravi Ram and Wolfner, 2007) and honeybees (Baer et al., in press; Collins et al., 2006) and several vertebrates including humans (Fung et al., 2004; Pilch and Mann, 2006). These seminal fluid components can affect both sperm cells and female physiology (reviewed by Chapman and Davies, 2004; Gillott, 2003; Poiani, 2006; Ravi Ram and Wolfner, 2007; Simmons, 2001) and at least some seminal fluid proteins have been predicted to enhance sperm viability and sperm survival (Baer et al., in press; Chapman and Davies, 2004).

Females are also known to provide sperm with glandular secretions. In vertebrates, bovine oviduct secretions have been shown to affect sperm motility and viability (Abe et al., 1995) and to enable sperm capacitation to increase fertilization success (King et al., 1994). Females of invertebrate species often possess specialized sperm storage organs (sometimes referred to as spermathecae) where sperm is kept between mating and egg fertilization (Eberhard, 1996; Simmons, 2001). These storage organs are often accompanied by glands and their secretions have been hypothesized to benefit the survival of stored sperm (Prokupek et al., 2008), although neither the molecular composition of these secretions nor their biological activity have been studied in great detail (but see Klenk et al., 2004; Koeniger, 1970; Lensky and Schindler, 1967).

Investigating the mechanisms by which males and females affect sperm viability is particularly interesting in the eusocial Hymenoptera (the ants and some of the bees and wasps). Copulations and insemination are restricted to a single brief mating episode early in a queen’s life (Boomsma et al., 2005; Boomsma and Ratnieks, 1996). Males die during or shortly after copulating while queens store large amounts of sperm that remain viable over prolonged periods of time, sometimes for several decades (Keller, 1998; Pamilo, 1991). Reproductive success of males and queens is therefore likely to be correlated with both the quantity and the quality of the sperm cells that females are able to acquire and store (Cole, 1983). Male seminal fluid may thus be particularly important for sperm viability during the provisional storage of ejaculates in the female sexual tract prior to final storage. After transfer to the spermatheca, sperm might have to
survive within the spermatheca for years before it will be able to fertilize eggs. It would thus seem obvious that glandular secretions from the queen’s spermathecal glands are also important for sperm viability, but no explicit tests have been done to quantify such effects.

In a recent study on Atta leafcutter ants we showed that male accessory gland (AG) secretions, that were inferred to contribute most of the seminal fluid, have a positive effect on sperm viability even when sperm is only exposed to minute quantities of these secretions (Den Boer et al., 2008). In the present study we use the honeybee Apis mellifera to further examine this effect, by focusing on proteins within the AG secretion of males. We also test the effect of queen spermathecal secretion on sperm viability and examine whether the proteins produced in male and queen secretions are sex-specific. Honeybees have been a long standing model organism in biology, so that many basal aspects of its mating biology have been developed as a semen extender used for artificial insemination in honeybees and is a relatively simple saline solution that represents an environment that is similar to the inorganic fraction of the ejaculate (Schley, 1987). Because it has no added proteins, carbohydrates, fatty acids or amino acids, Hayes is expected to impose some physiological stress on the sperm cells. This makes Hayes saline ideal as a control solution to examine the positive effects of seminal fluid and spermathecal fluid proteins on sperm survival in an osmotically suitable but slightly suboptimal environment (see also Den Boer et al., 2008).

Sperm viability was measured using a Live/Dead™ sperm viability kit (L-7011, Molecular Probes; Collins and Donoghue, 1999; Den Boer et al., 2008). The kit consists of two fluorescent dyes that allow the experimenter to distinguish live (green emission, using SYBR-14 dye) from dead sperm cells (red emission, using propidium iodide). We used a Leica fluorescence microscope (blue excitation filter at λ = 490 nm) at 400× magnification and counted the number of live (green), dead (red) and dual-stained sperm cells for at least 400 sperm cells for each sample. Dual stained sperm cells represented on average 0.13 ± 0.03% (mean ± S.E.M.) of the total sperm population and were therefore excluded from the data.

2. Material and methods

2.1. Sampling of bees

Bees used for the experiments originated from several colonies of Apis mellifera carnica that were kept at the University of Western Australia. Mature males and virgin queens became available during the Australian summer between October 2007 and January 2008, a time span that includes the natural reproductive season of local honeybees. Mature males were collected from male-producing colonies. Virgin females were obtained by grafting, whereby 4-day-old female larvae where reared into virgin queens using queenless colonies. Virgin queens were used for experiments at an age of 5–8 days after hatching, which is the typical period for nuptial flights (Ruttner, 1975).

2.2. Dissections and sperm viability measurements

Dissections were carried out with Inox 5 (Biology) watchmaker forceps in Hayes solution (9 g NaCl, 0.2 g CaCl2, 0.2 g KCl and 0.1 g NaHCO3 in 1000 ml H2O, pH 8.7). Hayes solution was originally developed as a semen extender used for artificial insemination in honeybees and is a relatively simple saline solution that represents an environment that is similar to the inorganic fraction of the ejaculate (Schley, 1987). Because it has no added proteins, carbohydrates, fatty acids or amino acids, Hayes is expected to impose some physiological stress on the sperm cells. This makes Hayes saline ideal as a control solution to examine the positive effects of seminal fluid and spermathecal fluid proteins on sperm survival in an osmotically suitable but slightly suboptimal environment (see also Den Boer et al., 2008).

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2.3. The effect of AG secretion on sperm viability

To investigate the effect of male AG secretion on sperm viability we collected 20 mature males. Each male was killed and his two accessory glands were dissected (Fig. 1A) and placed in 1 ml Hayes saline. The glands were carefully ruptured to help the gland content dissolve into the Hayes saline. The sample was then vortexed and centrifuged for 3 min at 3000 × g to separate the soluble gland secretion from the remaining gland tissue. We also collected two sperm samples from the same male by puncturing each of his two accessory testes (also referred to as seminal vesicles, Snodgrass, 1956) in a drop of 4 µl Hayes and obtaining 2 µl aliquots of the out flowing sperm in Hayes, using a micropipette. One of these was then dissolved in the 1 ml Hayes solution containing AG secretion as described above, whereas the second sperm sample was dissolved in 1 ml of Hayes only (control). Sperm viability was then estimated using 5 µl of each sample from which >400 sperm cells were counted (see above).

![Fig. 1. The sexual organs of an Apis mellifera male and queen: (A) The testes (T) degenerate shortly after hatching and sperm is stored in the accessory testes (AT, also known as seminal vesicles). A major part of the seminal fluid is produced in the accessory glands (AG) and mixed with the sperm when an ejaculate is transferred via the ejaculatory duct (ED). The accessory glands also produce a less soluble component, which is typically referred to as mucus and forms a mating plug in honeybees. (B) The spermatheca (S) is covered by a network of trachea and the paired spermathecal glands (SG).](image-url)
2.4. The effect of proteins in accessory gland secretion on sperm viability

To test whether the protein fraction of the seminal fluid had a particularly distinct effect on sperm viability, we collected 25 mature males from several colonies. Their accessory glands were dissected and pooled in 500 μl Hayes saline. After rupturing the glands the solution was vortexed and centrifuged for 20 min at 20,000 × g and 4 °C to separate the soluble secretion from the non-soluble gland tissue and mucus. We then centrifuged the supernatant for 30 min at 10,000 × g and 20 min at 12,000 × g and 4 °C with a Millipore Ultrafree centrifugal filter device with a 5 kDa cut off membrane. Because the smallest seminal fluid protein identified so far is 11 kDa (Baer et al., in press), the vast majority of proteins, including all the ones hypothesized to be involved in sperm viability, did not pass through the membrane. Consequently we ended up with two subsamples that we refer to as the protein fraction (proteins above 5 kDa) and the non-protein fraction (peptides and other metabolites below 5 kDa) of the AG secretion. From a total of 180 μl of protein fraction obtained, we used 30 μl (equivalent to 4 males) for SDS-PAGE gel runs (see below) to visualize the protein profile, whereas the remaining 150 μl (equivalent to 21 males) was diluted in 2 ml of Hayes saline and used for sperm viability assays (see below). Of the 300 μl of non-protein fraction obtained, we used 48 μl (equivalent to 4 males) for gel runs and the remaining 252 μl (equivalent to 21 males) for sperm viability assays after dilution in 2 ml Hayes.

To test for the effects of AG proteins on sperm viability we used another 19 males. Three sperm samples of 1 μl each were collected from each male, by puncturing the accessory testes in a 3 μl drop of Hayes saline and collecting the out flowing sperm with a micropipette. The sperm samples were diluted in 100 μl of (1) the protein fraction, (2) the non-protein fraction and (3) Hayes saline (control). Sperm viability was then estimated for a 5 μl subsample of each treatment.

To visually compare the protein profile of AG secretion with that of seminal fluid of male ejaculates, seminal fluid was collected from an additional set of males according to a standardized protocol as outlined below and used for SDS-PAGE gel runs. Males were killed in chloroform, which stimulates ejaculation and thus allowed us to collect entire ejaculates. A total of 20 μl of pooled ejaculated sperm from 20 to 30 males was collected and diluted in 50 μl Hayes solution, mixed and centrifuged for 25 min at 850 × g and 4 °C and afterwards for 10 min at 18,620 × g and 4 °C to separate sperm cells from the seminal fluid.

2.5. The effect of spermathecal secretions on sperm viability

To examine the effect of spermathecal fluid and spermathecal gland secretion on sperm viability, we used thirty virgin sister queens. The spermathecae and spermathecal glands of these queens were collected in 200 μl Hayes saline each (Fig. 1B). Both spermathecal glands and spermathecae were ruptured, vortexed and centrifuged for 3 min at 3000 × g to separate tissue from secretion. In addition, AG secretion was obtained from ten mature males as described in the previous sections. From each of these males we also collected four samples of 0.5 μl sperm, by puncturing their accessory testes in 2 μl of Hayes and collecting the out flowing sperm. The samples were dissolved in 100 μl of (1) the spermathecal fluid secretion, (2) the spermathecal gland secretion, (3) male AG secretion and in (4) Hayes saline (control). Sperm viability was then estimated using 5 μl aliquots.

SDS-PAGE was used to visualize the protein profiles of male AG secretions, seminal fluid, spermathecal glands and spermathecal fluid, using Biorad Criterion precast gels (10–20% [w/v] Acrylamide, HCl, 1 mm, 18 comb). Gels were run at 30 mA, fixed in a solution of 40% methanol and 10% acetic acid for an hour, and stained overnight with colloidal Coomassie blue (G 250). A total of 27 μg of protein was loaded onto the gel for the seminal fluid, the AG secretion and the protein fraction of the AG secretion. For the non-protein fraction of the AG secretion, we loaded 15 μl of sample. For the spermathecal samples we loaded 20 μl of spermathecal gland fluid and 8 μl of spermathecal fluid on the gel.

2.6. Statistics

Statistical analyses were carried out using SAS 9.1 for Windows. We examined the overall effect of treatment on sperm viability in all experiments using a generalized linear model with a binomial error distribution and a logit-link function, with treatment (the solution sperm was dissolved in) as repeated measure on the same male. The data were over-dispersed so we estimated the dispersion parameter from the scaled Pearson Chi-square. We used pair-wise contrasts to examine differences between treatment levels. To test for differences in overall sperm viability between males, we used the same generalized linear model, but with male as an independent class variable. χ² values are presented for the treatment effects, linear contrasts and difference between males.

3. Results

Sperm viability was significantly higher in the presence of male AG secretions compared to the control treatment (χ² = 15.76, df = 1, p < 0.001, Fig. 2A), but we found no significant difference between individual males in overall sperm viability (χ² = 26.09, df = 19, p = 0.128).

When we tested for effects of the protein and the non-protein fraction of male AG secretion, we found significant overall differences (χ² = 15.19, df = 2, p < 0.001, Fig. 2B). Sperm viability was significantly higher when exposed to the protein fraction of AG secretion compared to the non-protein fraction (χ² = 15.25, df = 1, p < 0.001). Sperm viability was also significantly higher in the non-protein fraction compared to the control treatment (χ² = 5.92, df = 1, p = 0.015), indicating that metabolites and peptides that were able to pass through the filter (<5 kDa) enhance sperm viability as well, but that this effect is smaller. Similar to our first experiment we did not detect significant differences in sperm viabilities between individual males (χ² = 11.28, df = 18, p = 0.882).

Visual inspection of 1D SDS-PAGE gels confirmed the absence of larger proteins in the non-protein fraction of the AG secretion (Fig. 3A, fourth lane) and showed that the profile of the protein fraction is similar to that of ejaculated seminal fluid, confirming that AG secretion is indeed the major contributor to seminal fluid in the honeybee.

When we tested the respective sperm viability enhancing effects of spermathecal fluid and spermathecal gland secretion, we found that they were similar, both to each other and to the effect of male AG secretion (χ² = 0.46, df = 2, p = 0.796), whereas all three were significantly higher than the control (χ² = 7.77, df = 1, p = 0.005, Fig. 2C). Once more, there was no difference in overall sperm viability between the males tested (χ² = 7.05, df = 9, p = 0.632).

Inspection of the 1D SDS-PAGE gels (Fig. 3B) suggested that the protein profiles of spermathecal fluid and spermathecal gland secretion are visually comparable to those reported elsewhere (Klink et al., 2004; Baer et al., submitted for publication). However, while the profiles for the spermathecal gland secretion and the spermathecal fluid were similar, both appear to be different from the profile of the male seminal fluid. This corresponds to the findings of Baer et al. (in press, submitted for publication), who
shows that the proteomic profiles of seminal fluid and spermathecal fluid hardly overlap.

4. Discussion

The transfer and storage of highly viable ejaculates is particularly important in social insects where queens never remate to replenish their sperm supply and sperm therefore needs to be kept viable for years (Boomsma et al., 2005). Adaptations in queens and males that prevent sperm death are therefore expected. Our study indeed shows that honeybee males and females both contribute glandular secretions that enhance sperm viability. In the sections below, we evaluate the implications of our findings, both by comparing our findings with older literature on sperm transfer and sperm storage in the honeybee, and by identifying novel questions of potential evolutionary trade-offs that may have shaped ejaculate and sperm storage traits. Finally, we discuss the molecular mechanisms that need to be clarified for a full understanding of the potency of these reproductive gland secretions.

4.1. Male accessory gland effects on sperm survival

Our work confirms that the AG secretions of males are of significant importance for sperm viability in the honeybee. As in our previous study on Atta leafcutter ants (Den Boer et al., 2008), our present data indicate that AG secretions are highly potent and express their positive effects on sperm survival even when diluted in a saline solution. In addition, we provide further evidence that the AG secretions are a main component of seminal fluid and are indeed transferred to the female as part of the ejaculate.

Secondly, we provide the first empirical evidence in insects for the hypothesis that proteins within the AG secretion are particularly important in maintaining sperm viable and that the non-protein fraction has a lesser, but also significantly positive, effect. Amino acids or sugars are likely candidates for producing this positive effect of the non-protein fraction, as they have previously been shown to increase sperm motility and sperm survival in honeybee semen (Verma, 1981; Poole and Edwards, 1970). We hypothesize that the effect of these metabolites on sperm survival may be relatively short-term (e.g. the duration of our experiment), while the proteins might be more important for maintaining sperm viability for the 40 h that sperm spend within the bursa copulatrix of queens before they are transferred to the spermatheca. An experiment to quantify the duration of the effect of the protein and non-protein seminal fluid components on sperm viability would therefore be interesting for future work. Likewise, it would be interesting to quantify whether the positive effect of the protein fraction is relatively more important in Apis mellifera honeybees than in A. florea and A. andreniformis dwarf honeybees (Koeniger and Koeniger, 1991; Koeniger et al., 1989) which, similar to Atta leafcutter ants (Baer and Boomsma, 2006), have sperm stored in
the spermatheca without prestorage in the female reproductive tract.

4.2. Queen spermathecal gland effects on sperm survival

Similar to male seminal fluid, the queen spermathecal fluid and spermathecal gland secretion have positive effects on sperm viability. We found no difference between the treatments with spermathecal fluid and spermathecal gland secretion (Fig. 2C), which matches the observation that their protein profiles are very similar (Fig. 3B). The epithelium of the spermatheca itself lacks glandular structures, (Dallai, 1975) which underlines that the spermathecal glands are the source of the proteins found in the spermathecal fluid (also see Klenk et al., 2004). In addition, smaller molecules within the spermathecal environment seem important to keep sperm alive whilst in storage. For example, the honeybee spermatheca is known to have significantly higher concentrations of Na+ and K+ ions compared to the surrounding haemolymph, (Bass and Gessner, 1976), which has been hypothesized to act as a reversible inhibitor of sperm motility (Verma, 1973). The spermathecal lumen has also been found to have a surprisingly alkaline pH of 8.6 (Gessner and Gessner, 1976), which may induce sperm dormancy (Lensky and Schindler, 1967). Furthermore, elevated levels of anti-oxidant enzymes found in the spermatheca are likely to protect sperm cells from oxidative stress (Weirich et al., 2002).

By visually comparing the gels (Fig. 3B), we can conclude that the protein profiles of male seminal fluid and queen spermathecal fluid are likely to differ considerably, which would indicate that the sexes accomplish their support functions for sperm viability in fundamentally different ways. Proteomic work by Baer et al. (in press, submitted for publication), where individual seminal fluid and spermathecal fluid proteins have been identified, confirms that there is little overlap between proteins produced in the male and queen glandular secretions. We hypothesize that this difference is related to the different time windows of male and female reproduction, so that seminal fluid primarily affects short-term sperm viability and spermathecal fluid is more important for long-term viability after storage.

4.3. Understanding the functional significance of sperm support compounds

Male AG secretions and seminal fluid are complex biochemical mixtures and are therefore likely to generate costs when males have to produce considerable amounts of these substances during their short adult lives. The peculiarities of eusocial hymenopteran mating systems imply that males do not renew their sperm and AG secretions (Boomsma et al., 2005), so that these costs may be a relatively constant proportion of total reproductive investment. However, this may be different in promiscuous mating systems where males are as long lived as females. A recent study on crickets, (Baer et al. in press; Collins et al., 2006) showed that colony-founding queens of Atta leafcutter ants may pay a substantial cost for storing higher than average amounts of sperm as this apparently trades off with their immune response during this vulnerable solitary stage in their life. Such trade offs are less likely to apply in honeybees, who found new colonies by swarming so that young queens are always well provisioned. However, both Apis honeybees and Atta leafcutter ants have queens that are considerably more long-lived than queens of sister groups such as Bombus bumblebees and Trachymyrmex fungus-growing ants. It would thus be highly interesting to obtain comparative data on the identity and production costs of spermathecal proteins that secure the viability of stored sperm for one or a few years, relative to those active in queens that live for decades.

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