Provisioning mass by females of the maritime earwig, Anisolabis maritima, is not adjusted based on the number of young

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Abstract

The amount of parental provisioning is thought to reflect the need of offspring. This hypothesis was tested in the case of provisioning food mass to young with controlled clutch size using the maritime earwig, Anisolabis maritima Bonelli (Dermaptera: Anisolabididae). The female provisioned a constant mass of food to the young irrespective of the number of nymphs and the distance of food carrying. In addition, the survival rate of young did not change with adjusted clutch size. This study showed that A. maritima females appear to provide food mass to their nymphs independent of their number.

Keywords: begging signal, maternal care, parent–offspring conflict

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Introduction

Provisioning has been reported for several insect species (Costa 2006). Progressive provisioning, when parents repeatedly transport food to their young, can be regarded as a special form of parental care that is found in some insect species (Filippi-Tsukamoto et al. 1995; Filippi et al. 2001). Parental care can only spread if the benefits of care outweigh the offspring number advantage of non–caring individuals. Food provisioning to the young by parents is a widespread phenomenon that enhances the survival of the young, but preparing the food can be very costly to the parents (Clutton-Brock 1991). Thus, the mechanism of food allocation to the young is an important issue of evolutionary biology. It is possible that some parents have a flexible investment dependent upon their young’s need, in order to minimize the costs of current reproduction (Sæther et al. 1993).

The amount and duration of parental investment are also influenced by a conflict between parents and offspring for the need of investment (Trivers 1974). The evolution of offspring begging signals is predicted based on an evolutionary resolution of this conflict (Parker et al. 2002). Begging signals are thought to reflect the needs of offspring, which parents use to adjust their food allocation (Godfray 1991). Recently, the presence of food begging signals for parents has been recognized in some insect species (Kölliker et al. 2005; Smiseth and Moore 2004). All the species of earwig (Dermaptera) studied to date exhibit parental care (Lamb 1976), though the extent of care varies greatly from species to species (Vancassel 1984). Furthermore, the females of some species have been reported to provision their nymphs (Shepard et al. 1973; Lamb 1976; Rankin et al. 1996; Staerkle and Kölliker 2008). Mas et al. (2009) reported the presence of chemical begging signals to their mother, this group will be suitable material for examining the regulation of female–offspring interactions.

Anisolabis maritima Bonelli (Dermaptera: Anisolabididae) is a cosmopolitan earwig species that exhibits subsocial behavior, where the females tend clutches of eggs in soil burrows (Bennett 1904). The females provision nymphs progressively, and providing food to the nymphs increases the nymph survival (Suzuki 2010). Suzuki (2010) reported that A. maritima females provision increasing amounts of food as the number of days from hatching increases. Provisioning in A. maritima may play a crucial part in the maintenance of family groups, because the survival rate of nymphs decreased greatly in the absence of provisioning by the female (Suzuki 2010). Thus, it is possible that females control food mass or provisioning times for nymph needs. Since A. maritima females lack kin recognition ability (Suzuki, unpublished data), their clutch size can be manipulated easily, and the manipulation can be expected to cause a change in the intensity of begging signals by the young. The aim of this study was to determine, by controlling clutch size, whether A. maritima females regulate their food mass provisioning in response to their young’s needs. The effect of the size of food mass on the survival of nymphs was also studied.

Materials and Methods

All A. maritima earwigs were caught in a field at the coast of Izumozaki, Niigata prefecture, Japan (37° 32’ 11’’ E, 138° 42’ 10’’ N) from late April to early May in 2009. All females were paired with a male for more than one day.
prior to the start of the experiment. After measuring body length, females were placed in a polyethylene container (8 × 5 × 4 cm) on a 1-5 mm layer of moist sand. Containers were placed under dim light at room temperature with sufficient humidity. All individuals were fed with turtle food pellets ad libitum. All containers including females were checked daily, and when egg masses were found, the containers containing the masses were subjected to the experiments.

Experiment 1 (adjusted clutch size): The mean size of the typical A. maritima first clutch is 58.0 eggs, with a range of 19-125 (Suzuki, unpublished data). Two or more clutches were produced within two days, and eggs were moved carefully with forceps. Clutch size was adjusted to 30, 60, or 90 eggs (adjusted clutch size) after determining the original clutch size. Since A. maritima females cannot distinguish their own eggs from a stranger’s eggs (Suzuki, pers. obs.), they treated the adjusted clutches normally. 16 clutches were adjusted as 30 eggs, 21 clutches as 60 eggs, and 16 clutches as 90 eggs. When hatched nymphs were found, food for nymphs was placed beside them. Bottle caps (25 mm diameter, 10 mm depth) placed at a distance of 2-3 cm from the burrow were used as food containers. Immature (before dispersal) nymphs could not enter the bottle cap and eat the food (Suzuki 2010). The food provided was 15 turtle food pellets (average 0.1 g), a mass that is 1.5 times or more than the need of nymphs in a normal–sized brood (Suzuki 2010). Each container was checked daily, at which time the uneaten pellets were counted and replenished by adding up to 15 pellets as needed. When more than half of the nymphs had left the nests, or some nymphs were present in the bottle cap, the brood was recorded as dispersed. Since the day of hatching cannot be synchronized within an adjusted clutch, the time span ranging from the first day of food carrying to dispersal day was regarded as the duration of care. After nymph dispersal, the number of surviving nymphs was counted.

Experiment 2 manipulated the distance between food source and nest to potential verify the cost of provisioning. After confirming egg mass, the females and eggs were placed in larger boxes (20 × 7 × 6 cm) and clutch sizes were adjusted to 60 (N = 11). Bottle caps (25mm diameter, 10mm depth) placed at a distance of 15 cm from the burrow were used as food containers. Food was provided for hatched nymphs once they were found. Using same procedure of Experiment 1, 15 turtle food pellets were provided. Each container was checked daily, at which time uneaten pellets were counted and replenished by adding up to 15 pellets as needed.

**Results**

Experiment 1 (adjusted clutch size): The original clutch size (before adjustment) did not differ among adjusted clutch size (30: 54.7 ± 9.3; 60: 60.9 ± 14.2; 90: 58.3 ± 15.9, one–way ANOVA, $F = 0.91$, df = 2, $p = 0.4$) and did not correlate with the body size of the adults ($r = 0.19$, $p = 0.37$). Thus, the effect of body size was excluded from the analysis. Despite adjusted clutch size, females carried most of the food from the bottle cap and provisioning increased over time regardless of clutch size (Figure 1, GLM using REML analyses with individuals as a random factor, date: $F = 15.9; df = 4$, $p < 0.01$, clutch size difference: $F = 1.71$, df = 2, $p = 0.12$, date*clutch interaction: $F = 0.14$, $p = 0.70$). The survival rate of nymphs was not significantly different among broods of adjusted clutch size (one–way ANOVA after arcsine transformation, $F = 0.6$, df = 2, $p = 0.9$). Duration of care was also not
Figure 1. Daily changes and difference among adjusted clutch size in the average ± SE amount of food provisioning by Anisolabis maritima. High quality figures are available online.

Figure 2. Daily changes in the average ± SE amount of food provisioning with increased distance from the food source by Anisolabis maritima. High quality figures are available online.

significantly different among broods of adjusted clutch size (30: 4.9 ± 0.4; 60: 4.8 ± 0.4; 90: 4.9 ± 0.3, one-way ANOVA $F = 0.16$, df = 2, $p = 0.8$). However, since the day of hatching cannot be synchronized within an adjusted clutch, this duration of care seemed not to be exact.

In Experiment 2, where the cost of care was manipulated by varying the distance to the food source, females still carried most of the food from the food cap regardless of the increased distance to the food source. Furthermore, the number of carried food pellets was not different from that found in Experiment 1 in which the distance to the food source was shorter (Repeated–measures ANOVA, $F = 3.3$, df = 1, $p = 0.08$, Figure 2).

Discussion

Providing food to their offspring may cost parents not only because of higher risks of predation while foraging, but also increased energy expenditure. Parents reportedly gauge the young’s need by measuring the level of offspring begging (Kilner and Johnstone 1997). Provisioning the young improves their survival rate but may decrease female energy intake required for future reproduction (Mas et al. 2008). However, the results of this experiment indicated that A. maritima females provision a constant mass of food to the young regardless of brood size, even though the survival rate of nymphs was independent of brood size, and females carried much more food than the young needed. Though provisioning improves the survival of young (Suzuki 2010), females of A. maritima will not control the mass of provision precisely.

In species where the parents perform food provisioning, reciprocal parent–offspring interaction drives the evolution of offspring begging signals. Begging signals are thought to reflect the hunger level of offspring, which parents use to adjust their food allocation (Godfray 1991). Empirical studies conducted on birds showed that feeding the offspring increased their begging (Kilner and Johnstone 1997). Food begging behavior by the young has been reported in some insect species (den Boer and Duchateau 2006; Kölliker et al. 2005, 2006; Smiseth et al. 2003). However, these studies have postulated both the presence of a begging signal and the ability of parents to recognize the begging signal. Unlike the burying beetle (Smiseth et al. 2003) or the Common European earwig (Staerkle and Kölliker 2008), A. maritima
does not allocate food to individual young but brings food back for the whole brood (Suzuki 2010), similar to the burrower bug (Knight 1997). In such types of provisioning, the benefit of maternal food provisioning is simultaneously shared among all offspring, and competition among the young can disturb food allocation by the parents. Since begging signal has been reported in the nymphs of burrower bug (Kölliker et al. 2005), absence of individual food allocation will not be necessary to the evolution of the begging signal.

Progressive provisioning has been reported in some Cydnidae species (Filippi et al. 2001; Kölliker et al. 2006) that carry plant seeds as food for their young. In field conditions, A. maritima will eat small invertebrates (Bennett 1904; Suzuki, personal observation). Live and/or dead invertebrates are protein rich but seem to be difficult to collect constantly, although their availability in the field is unknown. As a resource this type of food can easily go rotten and is difficult to preserve. Foods for this experiment in the laboratory also easily go rotten.

The benefit of provisioning in A. maritima is substantial (Suzuki 2010), although the details of the cost have not been clarified. Since females seem to provide too much food for the young, females may pay an extra price for provisioning. The cost of care in some subsocial insects has been reported; for example, the interval from the first to the second clutch was shorter without care in the Dermapteran species F. auricularia (Vancassel and Foraste 1980; Vancassel 1984; Kölliker 2007) and E. annulipes (Rankin et al. 1996). Such delay may be common in Dermapteran species and may reduce the opportunity for reproduction. Agrawal et al. (2005) showed a very small cost of provisioning in the burrower bug (Sehirus cinctus). If the cost of provisioning food by A. maritima is negligibly small, it is unnecessary to minimize provision dependent upon the young’s need. If this hypothesis is correct, females can provision in excess of their young’s need, since provisioning more than is needed does not have a negative effect on reproduction. These experiments were planned to provide excess food for young’s consumption (see Suzuki 2010). All experiments were conducted in artificial condition; the cost of provisioning may be lower in field conditions. There needs to be further investigation to verify this hypothesis.

In summary, this study asserts the possibility of the absence of controlling food provisioning in A. maritima. Although the current evolutionary theory on parent–offspring conflict resolution has generally assumed that offspring begging signals advertise need, this result showed the possibility that parents provide food in excess of need. Detailed behavioral experiments will be required to fully understand how to determine food provisioning to the young.

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References

Agrawal AF, Combs N, Brodie ED. 2005. Insights into the costs of complex maternal care behavior in the burrower bug (Sehirus cinctus). Behavioral Ecology and Sociobiology 57: 566-574.
Bennett C. 1904. Earwigs (Anisolabis maritima Bon.). *Psyche* 11: 47-53.

Clutton-Brock TH. 1991 *The Evolution of Parental Care*. Princeton University Press.

Costa JT. 2006. *The Other Insect Societies*. Harvard University Press.

den Boer S, Duchateau M. 2006. A larval hunger signal in the bumblebee Bomibus terrestris. *Insectes Sociaux* 53: 369-373.

Filippi L, Hironaka M, Nomakuchi S. 2001. A review of the ecological parameters and implications of subsociality in Parastrachia japonensis (Hemiptera: Cydnidae), a semelparous species that specializes on a poor resource. *Population Ecology* 43: 41-50.

Filippi-Tsukamoto L, Nomakuchi S, Kuki K, Tojo S. 1995. Adaptiveness of parental care in Parastrachia japonensis (Hemiptera: Cydnidae) *Annals of Entomological Society America* 88: 374-383.

Godfray HCJ. 1991. Signaling of need by offspring to their parents. *Nature* 352: 328-330.

Kight SL. 1997. Factors influencing maternal behaviour in a burrower bug, Sehirus cinctus (Heteroptera: Cydnidae). *Animal Behaviour* 53: 105-112.

Kilner R, Johnstone RA. 1997. Begging the question: are offspring solicitation behaviours signals of need? *Trends in Ecology and Evolution* 12: 11-15.

Kölliker M, Chuckalovcak JP, Brodie EDI. 2006. Maternal food provisioning in relation to condition-dependent offspring odours in burrower bugs (Sehirus cinctus). *Proceedings of the Royal Society of London B—Biological Sciences* 273: 1523-1528.

Lamb RJ. 1976. Parental behavior in the Dermaptera with special reference to Forficula auricularia (Dermaptera: Forficulidae). *Canadian Entomologist* 108: 819-824.

Mas F, Kölliker M. 2008. Maternal care and offspring begging in social insects: chemical signaling, hormonal regulation and evolution. *Animal Behaviour* 76: 11-21.

Parker GA, Royle NJ, Hartley IR. 2002. Begging scrambles with unequal chicks: interactions between need and competitive ability. *Ecology Letters* 5: 206-215.

Rankin SM, Storm SK, Pieto DL, Risser AL. 1996. Maternal behavior and clutch manipulation in the ring–legged earwig (Dermaptera: Carcinophoridae). *Journal of Insect Behavior* 9: 85-103.

Roff DA. 1992. *The Evolution of Life Histories*. Chapman and Hall.

Sæther BE, Andersen R, Pedersen HC. 1993. Regulation of parental effort in a long–lived seabird: an experimental manipulation of the cost of reproduction in the Antarctic petrel, Thalassoica antarctica. *Behavioral Ecology and Sociobiology* 33: 147-150.

Shepherd M, Waddil V, Kloft W. 1973. Biology of the predaceous earwig Labidura riparia (Dermaptera: Labiduridae). *Annals of
the Entomological Society of America 66: 837-841.

Smiseth PT, Darwell CT, Moore AJ. 2003. Partial begging: an empirical model for the early evolution of offspring signaling. *Proceedings of the Royal Society of London B—Biological Sciences* 270: 1773-1777.

Staerkle M, Kölliker M. 2008. Maternal food regurgitation to nymphs in earwigs (*Forficula auricularia*). *Ethology* 114: 844-850.

Suzuki S. 2010. Progressive provisioning by the females of the earwig, *Anisolabis maritima*, increases the survival rate of the young. *Journal of Insect Science* 10:184. Available online, [http://insectscience.org/10.184](http://insectscience.org/10.184)

Trivers RL. 1974. Parent–offspring conflict. *American Zoologist* 14: 249-264.

Vancassel M. 1984. Plasticity and adaptive radiation of Dermapteran parental behavior: results and perspective. *Advances in the Study of Behavior* 14: 51-80.