

# COOPERATIVE BREEDING FAVORS MATERNAL INVESTMENT IN SIZE OVER NUMBER OF EGGS IN SPIDERS

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Received October 8, 2013 Accepted March 6, 2014

The transition to cooperative breeding may alter maternal investment strategies depending on density of breeders, extent of reproductive skew, and allo-maternal care. Change in optimal investment from solitary to cooperative breeding can be investigated by comparing social species with nonsocial congeners. We tested two hypotheses in a mainly semelparous system: that social, cooperative breeders, compared to subsocial, solitarily breeding congeners, (1) lay fewer and larger eggs because larger offspring compete better for limited resources and become reproducers; (2) induce egg size variation within clutches as a bet-hedging strategy to ensure that some offspring become reproducers. Within two spider genera, *Anelosimus* and *Stegodyphus*, we compared species from similar habitats and augmented the results with a mini-meta-analysis of egg numbers depicted in phylogenies. We found that social species indeed laid fewer, larger eggs than subsocials, while egg size variation was low overall, giving no support for bethedging. We propose that the transition to cooperative breeding selects for producing few, large offspring because reproductive skew and high density of breeders and young create competition for resources and reproduction. Convergent evolution has shaped maternal strategies similarly in phylogenetically distant species and directed cooperatively breeding spiders to invest in quality rather than quantity of offspring.

KEY WORDS: Adaptation, differential allocation, kin selection, parental manipulation, reproduction, social spiders.

EVOLUTION INTERNATIONAL JOURNAL OF ORGANIC EVOLUTION The transition to cooperative breeding is likely to alter the selective pressures on maternal strategies as parental care transforms into allo-parental care (Russell and Lummaa 2009). Several factors may influence how cooperatively breeding females invest optimally in their offspring. First, the density of breeders and amount of available resources determine whether juveniles compete for limited resources. Theoretical models by Brockelman (1975) and Parker and Begon (1986) show that optimal maternal investment is to produce few, large offspring when siblings and nonsiblings compete in a limited resource space, provided that larger offspring have a competitive advantage over smaller ones. Large clutches may further present a cost because more offspring may be more difficult for parents to tend and defend (Kam et al. 1998; Fox and Czesak 2000). These density-related effects could result in fewer offspring actually reaching the breeding age, which means that females producing fewer and larger offspring may gain higher fitness (Lack 1947; Noordwijk and Jong 1986; Godfray et al. 1991).

Second, cooperative breeding is usually characterized by some degree of reproductive skew (Keller and Reeve 1994). If larger offspring have a greater chance of becoming reproducers within a group, selection could also favor production of large offspring at the expense of offspring number (Brockelman 1975; Parker and Begon 1986). However, for many groupliving species, individuals' fitness is tightly linked to colony size, as larger colonies show lower risk of total group failure (e.g., Clutton-Brock et al. 1999; Kokko et al. 2001; Bilde et al. 2007). Hence, in species in which offspring stay in their natal colony as additional group members, selection would disfavor producing very small clutches. In this case, where selection favors both large offspring and large clutch size, reproductive resources might be allocated differentially into offspring to ensure that at least some will be large enough to become reproducers and that the group will be large enough to survive (diversified bet-hedging, e.g., Slatkin 1974; Philippi and Seger 1989; Einum and Fleming 2004). Maternal strategies in some animals do indeed include investing differentially in offspring within the same brood (Forbes 1999; Fox and Czesak 2000; Gibbs and Van Dyck 2009). This can be achieved, among other things, by manipulating egg size, nutritional packaging, or hatchling provisioning (e.g., Howe 1978; Crean and Marshall 2009).

Third, the probability that a female can produce a second clutch later, and the presence of nonreproductive helpers may alter optimal investment. A breeding female may strategically save resources for a subsequent breeding attempt by investing less in quality or quantity of her offspring as a plastic response if she has helpers at her nest that compensate for this decrease in fecundity (Russell et al. 2007; Taborsky et al. 2007; Russell and Lummaa 2009). This strategy, however, is not available for semelparous females that breed only once in a lifetime as they would not gain by saving reproductive resources for future reproduction.

We have learned from previous studies that alterations to social environments can induce a plastic response on maternal investment within some cooperatively breeding birds and fish, and that cooperative breeding across bird species correlates negatively with clutch size (Arnold and Owens 1998; Russell et al. 2007; Taborsky et al. 2007). Here, we take a different approach and investigate changes in maternal egg investment strategies that accompany the actual transition from solitary to cooperative breeding. Social spiders present an excellent system for testing maternal effect theories in the evolution of group living and cooperative breeding for several reasons: (1) Social spiders show allo-maternal care and reproductive skew: Less than half of all females in a colony reproduce, whereas the remaining females act as helpers (Vollrath 1986; Salomon and Lubin 2007; Salomon et al. 2008), and larger females are considered to be the ones that become reproducers (Vollrath and Rohde-Arndt 1983; Rypstra 1993; Salomon et al. 2008; Grinsted and Bilde 2013). Allomaternal care from mothers and helpers includes tending egg sacs and feeding hatchlings (Christenson 1984; Salomon and Lubin 2007). In the genus Stegodyphus, allo-maternal care is suicidal: the young finally consume their mother and all adult females of the colony (Seibt and Wickler 1987). (2) Permanent sociality has evolved from subsocial congeners multiple times within distantly

related spider families, and subsociality is still common allowing for comparative studies of social spiders and their ancestral state (Kullmann 1972; Kraus and Kraus 1988; Agnarsson 2006; Agnarsson et al. 2006; Johannesen et al. 2007). Subsocial spiders provide similar extended maternal care as social spiders, but juveniles show only a short period of cooperation in prey capture and feeding in the maternal nest before they disperse to live and breed solitarily (Avilés 1997; Lubin and Bilde 2007). Comparing traits from subsocial and social congeners can reveal valuable insights into the evolutionary consequences of the transition to permanent sociality and cooperative breeding from their subsocial ancestry. Comparing independent evolutionary origins of sociality from different spider genera may reveal evidence of convergent evolution and thus expose more general selective pressures on maternal investment. (3) Social and subsocial spiders usually reproduce only once in a lifetime (Lubin and Bilde 2007) so females need to optimize their investment in one clutch of eggs. Thus, the change in optimal maternal investment that accompanied the transition to cooperative breeding can be directly observed by comparing clutches of eggs from subsocial and social species.

We tested two, not mutually exclusive, hypotheses in the present study: first, if the transition to permanent group living and cooperative breeding was associated with competition among juveniles for resources and reproductive roles, we would expect that social females produce clutches of fewer, larger eggs than their subsocial congeners. Second, if social spider females use a bet-hedging egg-laying strategy to ensure that at least some of their own offspring become reproducers in the colony, we would expect that a social female produces a higher variation in egg sizes than a subsocial congener.

We investigated these hypotheses in two spider genera Anelosimus (Theridiidae) and Stegodyphus (Eresidae) to elucidate potential convergent evolution of maternal strategies. Both genera contain multiple independently derived social species that have evolved in distinctly contrasting habitats: Central and South American rainforest (Anelosimus) versus arid, open bush lands in Africa, the Middle East, and India (Stegodyphus) (Kraus and Kraus 1988; Agnarsson et al. 2006; Johannesen et al. 2007; Lubin and Bilde 2007). Within each genus, we compared maternal egg investment strategies of one social species with that of one or two closely related subsocial species that occurred in similar geographical regions and habitats. This allowed us to focus on evolutionary effects on maternal strategies while minimizing potentially confounding environmental effects. Based on previously published studies, we furthermore conducted a mini-meta-analysis on egg numbers in an additional nine species and have presented these data in reconstructed phylogenies of the two genera to augment the data in the present study.

## Methods study organisms

Social spiders live permanently in social groups in which females cooperate in prey capture and feeding, web building and web maintenance, and brood care (Bilde and Lubin 2011). Due to a lack of premating dispersal, social spiders breed with family members resulting in inbreeding and high levels of relatedness within colonies (Johannesen et al. 2002).

Anelosimus eximius occurs in the Americas from Panama to Argentina (Platnick 2012), where they form colonies of up to many thousands of individuals. Hence, A. eximius colonies are the largest of any social spider, although solitarily breeding females can occasionally be found (Vollrath 1982; Avilés 1997). Breeding occurs year round and thus colonies contain spiders of all instars through most of the year (Aviles 1986; Avilés 1997). Social A. eximius and subsocial A. baeza belong to the eximius group in the Anelosimus phylogeny (Agnarsson 2006) and thus are valid as a sister clade comparison. Additionally, spiders from both species used in this study were found in the same area in similar habitats. Anelosimus baeza occurs from Panama to Peru (Platnick 2012), where it mostly breeds solitarily, but can be found to live in multifemale colonies (L. Grinsted, pers. obs.). However, the level of cooperation, if any, within these colonies is unknown (Agnarsson 2006). Females of some subsocial Anelosimus species can produce two egg sacs in their lifetime (I. Agnarsson, pers. comm.), whereas others seem to only produce a second egg sac if the first one is abandoned (Marques et al. 1998). Females of social Anelosimus species are thought to usually produce only one egg sac in their lifetime although some might produce two (Aviles and Salazar 1999). Anelosimus spiders live for up to a year (Aviles and Tufino 1998).

Social S. sarasinorum occurs in India, Sri Lanka, and Nepal (Platnick 2012) in dry, shrubby habitats. Nests contain one female to several hundreds of individuals. The subsocial *S. tibialis* and *S.* pacificus also occur in India, and the individuals used in this study were found in the same area and in similar habitat as S. sarasinorum. Stegodyphus pacificus is the sister species of S. sarasinorum (Settepani et al., unpubl. data) and, hence, is appropriate for a sister clade comparison of the effects of social level on maternal strategies. Social and subsocial Stegodyphus females produce only one egg sac in their lifetime, unless the egg sac is lost after which they may produce another one, and they die when the young eventually consume them (Jacson and Joseph 1973). As these spiders only live for about a year and occur in seasonal habitats, breeding is restricted to one season in their lifetime and, hence, all spiders within colonies of social Stegodyphus spiders are approximately of the same age and life stage (Crouch and Lubin 2000; Lubin et al. 2009).

## **COLLECTIONS AND MEASUREMENTS** Stegodyphus

Nests refer to silken retreats consisting of either a single female with her single egg sac, or colonies with multiple females and multiple egg sacs. Nests of the subsocial *S. tibialis* ( $N_{nests} = 12$ ,  $N_{egg sacs} = 12$ ) and *S. pacificus* ( $N_{nests} = 10$ ,  $N_{egg sacs} = 10$ ) were collected from October to December 2010 near Kuppam in India ( $12^{\circ}48.854'N$ ,  $78^{\circ}15.964'E$ ). In the same area, colonies of the social *S. sarasinorum* were collected in January and February 2012. Whenever possible, up to five egg sacs per nest of *S. sarasinorum* were sampled ( $N_{nests} = 11$ ,  $N_{egg sacs} = 30$ , median  $N_{egg sacs/nest} = 2$ ). Upon collection, colonies were dissected and spiders were counted. Only females and males in the parental cohort were included in the colony size count; if colonies contained hatchlings, these were not included in the count. Colony sizes ranged from a single female to 106 spiders. An overview of species and sample sizes is available in the Supporting Information (Table S1).

The prosoma width of mothers was measured with a digital calliper (Toolmate) to the nearest 0.01 mm. Prosoma width is a widely accepted measure of body size in spiders as it is a sclerotized body part affected little by satiation state (Hagstrum 1971; Jakob et al. 1996). An estimate of mother size in multifemale colonies was obtained by taking the average prosoma width of up to 15 randomly chosen adult females in each colony (Bilde et al. 2007) or of all females in colonies smaller than 16.

#### Anelosimus

Egg sacs of the social *A. eximius* and subsocial *A. baeza* were collected near Sumaco in Ecuador (00°43.492'S, 77°38.665'W) in May and June 2011. The subsocial *A. baeza* sometimes formed multifemale nests, resembling social colonies, although adult females most likely did not cooperate in prey capture and brood care. Whenever possible, up to five egg sacs were sampled from nests of both *A. eximius* and *A. baeza* (*A. eximius*:  $N_{nests} = 33$ ,  $N_{egg sacs} = 136$ , median  $N_{egg sacs/nest} = 5$ ; *A. baeza*:  $N_{nests} = 21$ ,  $N_{egg sacs} = 30$ , median  $N_{egg sacs/nest} = 1$ ). An overview of sample sizes is available in the Supporting Information (Table S2).

Two measures of female body size were obtained: prosoma width and the combined length of tibia and patella of the first leg by measuring with a digital calliper to the nearest 0.01 mm either directly in the field or after collection of nests. The length of tibia + patella is recommended as a proxy for body size in *Anelosimus* spiders (e.g., Aviles 1986). The prosoma of *Anelosimus* spiders is relatively small (width: 0.9-1.7 mm) and therefore susceptible to higher measurement error when measured by hand, whereas the length of tibia + patella of the first leg is more easily measured (length: 2.0-3.8 mm). Estimates of mother size in social *A. eximius* were obtained by taking the average measure from up to 20 randomly chosen adult females in each colony or of all females

in colonies smaller than 21. When egg sacs were collected from multifemale nests of *A. baeza*, it was clear to which female the egg sac belonged, as only one female was in close proximity of the egg sac, and so the actual mother was measured.

The size of an *A. eximius* colony was obtained either by dissecting the colony and counting the spiders (22 nests) or estimated based on the physical dimension of the nest (11 nests: see Supporting Information). Colony sizes across the 33 nests ranged from a single female to 1934 nonjuvenile spiders (i.e., the number of subadult and adult males, and adult females and two different instars of subadult females). To determine the colony size of multifemale *A. baeza* nests, the transparent nests were visually inspected and adult and subadult females and males were counted. Colony sizes ranged from a single female to 54 nonjuvenile spiders.

### Egg number and egg sizes

In the laboratory, each egg sac was opened carefully and the eggs were spread out on a flat, black background. Broken and lost eggs were counted, and individual eggs with small, parasitic larvae attached were removed and counted. A photo was taken with a digital camera (Canon Cyber-shot DSC-W330 14.1 megapixel) at a set distance to the background. Immediately afterward a photo was taken of a piece of millimeter-scale paper for reference. The size of each egg in each of the clutches was measured, to nearest 0.01 mm<sup>2</sup>, with the use of a custom-fitted macro for ImageJ 1.45 (Abramoff et al. 2004; Gibbs et al. 2010), contrasting the light-colored egg against a black background. Egg numbers were obtained manually by counting eggs on each photograph and adding the number of lost or removed eggs for each clutch.

## EGG NUMBER AMONG SPECIES IN PHYLOGENIES

To supplement our results, we performed a literature search on reproductive strategies in *Anelosimus* and *Stegodyphus* species. We were also able to collect egg sacs from yet an extra *Stegodyphus* species (*S. mimosarum*, eight egg sacs collected from two nests in Madagascar, May 2012). In this way, we obtained egg numbers from an additional six *Anelosimus* and three *Stegodyphus* species. We then mapped the average egg number per egg sac from each species onto the phylogenies of the two genera so that closely related social and subsocial species could be compared. This meta-analysis was mainly meant for descriptive analysis and not for phylogenetic contrast analysis as sample sizes were low and the phylogenies were lacking branch lengths.

Specifically, we obtained egg numbers from the following social species: *A. domingo* (Aviles and Maddison 1991), *A. dubiosus* (Marques et al. 1998), *A. rupununi* (Aviles and Salazar 1999), *S. dumicola* (Aviles et al. 1999), and *S. mimosarum* (this

The as yet unpublished molecular phylogeny of *Stegodyphus* was based on nine independent nuclear loci (Settepani et al., unpubl. data). The nuclear loci were amplified with 13 primers designed from alignments of *S. lineatus*, *S. tentoriicola*, and *S. mimosarum* published in Mattila et al. (2012). The best substitution model for each locus was estimated with PartitionFinder (Lanfear et al. 2012). The phylogeny was constructed using the Bayesian method implemented in MrBayes 3.2 (Ronquist et al. 2012). MrBayes was run for 5 million generations with a sampling frequency of 500, a burn-in of 25%, and two chains. The partial *Anelosimus* phylogeny was based on the one published in Agnarsson (2006). The partial phylogenies presented here were drawn in TreeViewX.

#### STATISTICS

We used linear mixed effect models with the following three predictor variables: social level, mother size, and colony size. Response variables used were egg size (all individual egg sizes), variation in egg sizes within egg sacs (one value per egg sac), and egg number (one value per egg sac). The *lmer* function from the lme4 package (Bates et al. 2011) in R (version 2.14.2) was used (R Development Core Team 2011). For models with a Gaussian error structure, we checked whether the assumptions of normally distributed and homogenous residuals were fulfilled by inspecting qq-plots and the residuals plotted against fitted values. In each of these models, the response variable was transformed to optimize normality and homogeneity of the residuals (the different transformations are apparent in the Results, and presented in Supporting Information: Description of Statistical Models). We based model fitting on maximum likelihood rather than the default option of restricted maximum likelihood tests in these models. We used a Poisson error distribution with a loglink function in models in which egg number was the response variable and established that data were not overdispersed before proceeding.

For all constructed models, we confirmed that the model was robust and that there were no datapoints with a disproportionally large effect. We did this by excluding datapoints one by one and comparing the range of estimated coefficients derived with those obtained from the full model. We also determined the variance inflation factor for full models (reduced to linear models by excluding random effects) using the vif function from the *car* package (Fox and Weisberg 2011), ensuring the models did not suffer from multicollinearity. *P*-values were obtained by using likelihood ratio tests ( $\chi^2$ ) to compare full models with reduced models in which the main effect in question had been omitted. When continuous predictor variables were included in an interaction term, they were *z*-transformed to a mean of zero and an SD of 1 to facilitate interpretation of parameter estimates. The overall significance of the full model was established by comparing it to a null model that included all random effects and random slopes. Only if the full model was significant, we proceeded to test the significance of interaction terms and main effects. Only if interaction terms were found nonsignificant, the significance of the main effects involved in the interactions was tested.

For each of the three response variables, we started out by testing the effect of social level (i.e., social vs. subsocial) across genera by including all five species in the models. As female body size varies greatly among and within species, mother size was accounted for in the models by including it as a covariate. The interaction between social level and mother size was included to allow for the possibility that mother size had different effects on the response variable according to social level.

If any of the predictors showed a significant effect in tests including all five species, post hoc tests were performed with similar models testing the same predictors within genera. Colony size could be included into the models testing *Anelosimus* spiders, as both the social and subsocial *Anelosimus* species formed colonies. The effect of colony size was tested separately for the social *S. sarasinorum* (rather than in a test including all three *Stegodyphus* species) as the two subsocial *Stegodyphus* species always occurred solitarily. The effect of colony size was further examined in post hoc tests performed on each *Anelosimus* species by itself.

Whenever relevant, random effects and random slopes were included in the models. In some models, random slopes of mother size among genera, and mother size among species, were used. These allowed for random variation in the slope of the correlation between mother size and response variable among species and among genera.

We also examined the trade-offs between egg size and egg number at the level of individual mothers in each species separately. We did this by building generalized linear models (GLMs) with a negative binomial error distribution to account for overdispersion with egg number as the response variable and average egg size within egg sacs and mother size as predictor variables.

When including mother size in models containing all five species or only *Stegodyphus* species, prosoma width of females within colonies were used as a proxy for mother size. In models testing only *Anelosimus* species, length of tibia + patella was used as a proxy for mother size.

Detailed descriptions of all models tested are available in the Supporting Information (Description of Statistical Models). Based on the meta-analysis, we compared mean egg number from social species with that from their appropriate subsocial congener in a Wilcoxon test for matched pairs (see Table S3 for details on the species pairs). All raw data are available at Dryad Digital Depository (doi: 10.5061/dryad.t6k57).

## Results

## DOES SOCIAL LEVEL PREDICT EGG SIZE AND EGG NUMBER?

Social spider species laid larger eggs than subsocial congeners overall (Table 1 and Fig. 1A): level of sociality significantly predicted egg size in a model including all five species, but this was dependent on mother size (i.e., significant interaction between social level and mother size; Table 1). Specifically, post hoc tests within genera showed that although social level had a significant effect on egg size within both genera, mother size also significantly predicted egg size only within *Stegodyphus*, not within *Anelosimus*. Larger *Stegodyphus* females laid smaller eggs in both social and subsocial spiders (Table 1 and Fig. 1A). Colony size significantly positively correlated with egg size within both *Anelosimus* species (Fig. 2C, E), whereas this correlation was not significant in *S. sarasinorum* (Fig. 2A).

Social spider species laid significantly fewer eggs than their subsocial congeners in a model containing all five species (Table 2A and Fig. 1B). Mother size had a significant, positive effect on egg number overall (Table 2A). Within Stegodyphus species, this effect of mother size was highly significant (Figs. 1B, S1A) whereas social level showed a close-to-significant effect (Table 2A). In Anelosimus, both female size and social level significantly predicted egg number, dependent on the size of the colony (significant interaction between social level and colony size; Table 2A). This means that apart from larger mothers laying more eggs in both Anelosimus species (Fig. S1B), subsocial spiders laid more eggs than social spiders only in smaller colony sizes. When colony sizes reached their maximum for subsocial A. baeza, the egg numbers were similar to those of social A. eximius (Fig. 2D, F). Within each of the three group-forming species, A. eximius was the only one for which colony size had a significantly positive effect on egg number (Table 2A and Fig. 2B, D, F).

Social species laid significantly fewer eggs (grand mean 37.9 eggs per egg sac) compared to subsocial species (grand mean 127.8 eggs per egg sac) in sister clades (Table S3) in both *Stegodyphus* (Fig. 3) and *Anelosimus* (Fig. 4; Wilcoxon test for matched pairs, P = 0.0078).

We found a highly significant negative correlation between egg number and egg size, when mother size was taken into account, only in the subsocial *A. baeza* (Table 2B and Fig. S2). We also saw a negative correlation in subsocial *A. pacificus*, but this

Egg Size (individual egg sizes, log trans)					
GLMMs of the effect of social level and colony size		$\chi^2$	df	P-value	
Across genera (5 species, 85 nests, 13,282 eggs, 213 egg sacs)	Social level $\times$ <i>z</i> -trans prosoma width	5.07	11, 10	0.024	
Within <i>Stegodyphus</i> (3 species, 32 nests, 6407 eggs, 50 egg sacs)	Social level × z-trans prosoma width	0.90	9, 8	0.34	
	Social level	6.35	8, 7	0.012	
	z-Trans prosoma width	5.19	8,7	0.023	
Within social S. sarasinorum (29 egg sacs)	Full model (colony size + prosoma width)	5.34	5, 3	0.069	
Within <i>Anelosimus</i> (2 species, 52 nests, 6632 eggs, 157 egg sacs)	Social level × z-trans length of tibia+patella	1.69	9, 8	0.19	
	Social level $\times$ colony size	1.47	8,7	0.22	
	Social level	123.10	6, 5	<0.0001	
	z-Trans length of tibia + patella	0.81	7,6	0.37	
	Colony size	3.04	6, 5	0.081	
Within social A. eximius (127 egg sacs)	Colony size	9.08	5,4	0.0026	
	Length of tibia + patella	2.33	5, 4	0.13	
Within subsocial A. baeza (30 egg sacs)	Colony size	5.59	5, 4	0.018	
	Length of tibia + patella	0.66	5, 4	0.42	

Table 1. Results from generalized linear mixed models (GLMMs) testing the effect of various predictors on the response variable egg size.

The three columns to the right show test values and P-values. Significant P-values are highlighted in bold. The word "transformed" is shortened to "trans."

trend was not significant, and in the remaining three species, we found no correlation (Table 2B).

## DOES SOCIAL LEVEL PREDICT VARIATION IN EGG SIZES WITHIN CLUTCHES?

The variation of egg sizes within egg sacs of social species was not different from that of their subsocial congeners within both genera. Social level and *z*-transformed mother size had no effect on the coefficient of variation) among the five species ( $\chi^2 = 6.30$ , df = (8, 5), P = 0.10,  $N_{\text{egg sacs}} = 211$ ,  $N_{\text{nests}} = 85$ , Fig. 1C). Square root transformed colony size also did not affect the variation in egg sizes within egg sacs (log CV) in the three colony-forming species: *S. sarasinorum* ( $\chi^2 = 0.87$ , df = (4, 3), P = 0.35,  $N_{\text{egg sacs}} = 29$ ,  $N_{\text{nests}} = 11$ ), *A. eximius* ( $\chi^2 = 0.05$ , df = (4, 3), P = 0.82,  $N_{\text{egg sacs}} = 132$ ,  $N_{\text{nests}} = 32$ ), and *A. baeza* ( $\chi^2 = 0.02$ , df = (4, 3), P = 0.88,  $N_{\text{egg sacs}} = 30$ ,  $N_{\text{nests}} = 21$ ). Additionally, colony size in *A. eximius* did not affect the within-colony variation in average egg sizes per egg sac ( $\rho = -0.24$ , P = 0.25) or within-colony variation in egg number ( $\rho = -0.18$ , P = 0.38).

## Discussion

We found support for the hypothesis that with the transition from solitary to cooperative breeding, maternal investment strategies have been altered to invest in size of offspring on the expense of number. Social, cooperatively breeding spiders laid significantly

fewer, larger eggs than their subsocial congeners, when factoring out the effect of mother size. Our results suggest that the selection for laying fewer, larger eggs has acted similarly on females within two separate genera of spiders that evolved sociality in distinctly different geographical and environmental circumstances. Hence, we propose that convergent evolution has shaped maternal egg investment strategies in social spiders, likely due to competition among offspring for resources and reproduction in the transition to cooperative breeding. This may indicate that the evolution of cooperative breeding presents similar selective pressures more generally and that these may be applicable for a wide range of cooperatively breeding organisms. We found no support for the hypothesis that social spider females invest differentially in eggs within clutches. The variation of egg sizes within egg sacs was very low in both social and subsocial species, suggesting that females do not induce size variation among their offspring at the egg stage as a bet-hedging strategy.

# MATERNAL STRATEGIES IN THE EVOLUTION OF COOPERATIVE BREEDING

Theoretical models predict that there is an optimal balance between offspring number and size in animals (Smith and Fretwell 1974) and that the optimal egg size increases, whereas egg number decreases when competition among siblings and nonsiblings increases (Brockelman 1975; Sargent et al. 1987). Our results support these theoretical predictions as social spiders laid



**Figure 1.** Egg size (A), egg number (B), and variation in egg size (C) plotted against mother size (prosoma width, mm) of all five species. Egg size is the grand mean egg size (average egg sizes within egg sacs averaged within nests, mm<sup>2</sup>). Variation in egg size is the coefficient of variation in egg sizes within egg sacs, averaged within nests. Egg numbers are eggs per egg sac averaged within nests. *Stegodyphus* species are represented as squares and *Anelosimus* as circles. The social species *S. sarasinorum* and *A. eximius* are represented by black points and full regression lines. Subsocial species are represented by gray and white points and dotted lines: *S. tibialis* and *A. baeza* are gray; *S. pacificus* is white. Regression lines have only been drawn for statistically significant associations between variables, although here they represent correlations performed on averaged data and do not directly reflect the results from the statistical models.

fewer, larger eggs than their subsocial congeners. In social spider colonies, there may be high densities of breeding females, and both siblings and nonsiblings might have competed for limited resources in the transition to cooperative breeding. Larger offspring may have had a competitive advantage and, hence, grown enough to reproduce whereas smaller group members would fail to do so (Vollrath 1986; Lubin 1995; Ulbrich and Henschel 1999; Whitehouse and Lubin 1999). Hence, selection for producing large offspring may have been strong, even at the expense of offspring number. A comparable example may be found in the communally breeding banded mongoose, Mungos mungo, where young in large colonies compete for allo-maternal care. Larger, heavier young have higher competitive abilities and obtain more care and thus benefit from a significantly higher survival rate making them more likely to grow up to reproduce (Hodge et al. 2009). Hence, giving birth to larger pups is advantageous.

Cooperative breeding in animals is associated with costs and benefits that may influence how mothers optimally invest in their offspring. In social spider colonies, individual risk of mortality is diminished due to both a lack of dispersal and benefits of group living. These benefits include increased protection against predators when living permanently in a large, protective nest (Bilde et al. 2007; Lubin and Bilde 2007), and extensive brood care from mothers and helpers. Allo-maternal care provides clear fitness benefits to young in the form of higher survival and growth rates (Salomon and Lubin 2007). In mammals, fitness benefits of receiving help when reproducing also include increased growth and survival of offspring and allow for decreased interlitter intervals as females can breed more than once in their lifetime (Jennions and Macdonald 1994; Russell et al. 2003). Across bird species, cooperative breeding is associated with clutches of fewer eggs (Arnold and Owens 1998; but see Cockburn 2003), and in some fish, females lay clutches of smaller eggs when more helpers are present (Taborsky et al. 2007). This apparent lower fecundity in cooperatively breeding birds and fish seems to be compensated by significantly lower mortality rates due to saving of resources, allowing for the production of more clutches later in life (Arnold and Owens 1998; Taborsky et al. 2007). Small egg numbers in social spiders are often interpreted as a cost of group living on the reproductive output of individual spiders (Aviles and Tufino 1998; Bilde et al. 2007). Indeed, if social spiders as compared to subsocial spiders laid fewer eggs of a similar size, this could have been interpreted as a cost of cooperative breeding to reproductive output. However, our novel results suggest that investing in fewer offspring is compensated by an increase in egg size, and hence, may be an adaptation to, rather than a cost of, cooperative breeding and group living. That the altered maternal strategy is in fact an evolutionary response to social living is supported by our egg size data. Indeed, there was no overlap between the observed egg sizes of social females and that of their subsocial congeners, even though social



**Figure 2.** Egg size (A, C, and E) and egg number (B, D, and F) plotted against colony size (number of nonjuvenile spiders) for the three group-forming species: social *S. sarasinorum* (A and B), social *A. eximius* (C and D), and subsocial *A. baeza* (E and F). Egg size is the grand mean egg size (average egg sizes within egg sacs averaged within nests, mm<sup>2</sup>) and egg numbers are eggs per egg sac averaged within nests. Regression lines have only been drawn for statistically significant associations between variables, although here they represent correlations performed on averaged data and do not directly reflect the results from the statistical models.

Egg Number (eggs per egg sac)						
(A) GLMMs of the effect of social level and colony size (controlled for colony ID)		$\chi^2$	df	P-value		
Across genera (5 species, 212 egg sacs, 86 nests)	Social level $\times z$ -trans prosoma width	0.28	9, 8	0.59		
	Social level	4.21	8,7	0.04		
	z-Trans prosoma width	6.87	8,7	0.0088		
Within <i>Stegodyphus</i> (3 species, 51 egg sacs, 33 nests)	Social level $\times$ <i>z</i> -trans prosoma width	0.87	7, 6	0.35		
	Social level	3.61	6, 5	0.057		
	z-Trans prosoma width	12.30	6, 5	0.00045		
Within social <i>S. sarasinorum</i> (29 egg sacs, 11 nests)	Colony size	3.21	4, 3	0.073		
	Prosoma width	22.30	4, 3	<0.0001		
Within Anelosimus (2 species, 157 egg sacs, 52 nests)	Social level $\times$ <i>z</i> -trans length of tibia + patella	0.38	7, 6	0.54		
	Social level $\times$ colony size	5.97	6, 5	0.015		
	z-Trans length of tibia $+$ patella	7.64	6, 5	0.0057		
Within social A. eximius (127 egg sacs, 31 nests)	Colony size	7.25	4, 3	0.0071		
	Length of tibia + patella	8.02	4, 3	0.0046		
Within subsocial A. baeza (30 egg sacs, 21 nests)	Full model (colony size + length of tibia + patella)	4.28	4, 2	0.12		
(B) GLMs of the association with Egg Size (mother size included in all models)		z-Value	df	P-value		
Social S. sarasinorum (28 egg sacs)	Average egg size	-0.30	27, 25	0.76		
Subsocial S. tibialis (12 egg sacs)	Average egg size	0.85	11, 9	0.39		
Subsocial S. pacificus (9 egg sacs)	Average egg size	-1.81	8,6	0.070		
Social A. eximius (131 egg sacs)	Average egg size	1.16	130, 128	0.25		
Subsocial A. baeza (30 egg sacs)	Average egg size	-2.79	29, 27	0.0053		

Table 2. Results from models testing the effect of various predictors on the response variable egg number.

The three columns to the right show test values and *P*-values. Significant *P*-values are highlighted in bold. The word "transformed" is shortened to "trans". (A) Results from GLMMs. (B) Results from GLMs on each separate species. In these models, both mother size and average egg size were included as main effects although effects of mother size are not depicted here as they are similar to those depicted in the results from the GLMMs.

and subsocial female body sizes overlapped within both genera (Fig. 1A).

We found clear evidence of a trade-off between egg size and egg number in one species only, the subsocial *A. baeza*, of the five species examined. Trade-offs between egg size and number may be difficult to detect in species that use adult-acquired resources for reproduction or show parental care, because total reproductive effort is then difficult to quantify (Bernardo 1996; Fox and Czesak 2000). This is the case for subsocial spiders and to an even greater extent for social species with allo-maternal care, which may explain why we could not detect a trade-off in the remaining species.

## MATERNAL INFLUENCE ON SIZE VARIATIONS IN OFFSPRING

We found no difference in egg size variation within egg sacs between social spiders and their subsocial congeners, suggesting that social spider females do not induce size variation in their off-

spring at the egg stage. Recent evidence indicates that body size variation among group members in social spiders may be induced at an early life stage and remain more or less stable throughout the colony life cycle (Grinsted and Bilde 2013). These body size hierarchies lead to partitioning of reproduction, and if they arise at an early life stage, maternal effects might play a role in assigning reproductive roles in offspring. Although we found no evidence to suggest that females vary the sizes of their offspring at the egg stage, differential maternal investment in offspring can occur at later stages (Russell and Lummaa 2009). Maternal and allo-maternal feeding of young allows for potential maternally induced size differences among hatchlings by differential feeding. Alternatively, unsynchronized hatching of egg sacs could create these differences, as earlier hatched offspring would get a head start in weight gain (Laaksonen 2004). Further research on posthatching maternal effects and hatching asynchrony will add to the understanding of the mechanisms behind partitioning of reproductive roles in social spider colonies.



**Figure 3.** Partial phylogeny of *Stegodyphus* with average egg number  $\pm$  SD (rounded off to whole numbers; na indicates SD not available) given for the species from which data were available. Social species are presented in bold; the remaining species are subsocial. Stars indicate the species investigated in the present study. The tree represents topology only, that is, branch lengths do not indicate evolutionary distance. The topology is based on a preliminary molecular phylogeny from Settepani et al. (unpubl. data) constructed using 13 independent nuclear loci and analyzed with the Bayesian method implemented in MrBayes 3.2 (Ronquist et al. 2012).

## **PHYLOGENETIC DIFFERENCES**

Our results suggest similarities in maternal egg investment strategies in the convergent evolution of sociality in two phylogenetically distant spider genera. These two genera contain spiders with distinctly different natural histories and origins of sociality. Social and many subsocial Anelosimus species occur in nonseasonal rainforest habitats in the New World (Agnarsson et al. 2006), whereas social and subsocial Stegodyphus species occur in seasonal, arid, open shrub lands in the Old World (Kraus and Kraus 1988; Majer et al. 2013). Female Anelosimus spiders can usually reproduce throughout the year, and might produce a second egg sac after the first one (although it is unclear how often this occurs, Marques et al. 1998; Aviles et al. 2007). This means that resources obtained for egg production might not all be allocated to one brood, but may strategically be divided relative to future reproductive opportunities depending on environmental conditions, predicted survivability of the first brood, and probability of reproducing a second time. This situation is different for Stegodyphus species that only lay a second egg sac in case the first one is lost (Jacson and Joseph 1973). These differences in life-history traits between the two genera may explain why we found stronger as-



**Figure 4.** Partial phylogeny of *Anelosimus* with average egg number  $\pm$  SD (rounded off to whole numbers; na indicates SD not available) given for the species from which data were available. Social species are presented in bold; the remaining species are subsocial. Stars indicate the species investigated in the present study. When egg numbers were known from both solitary (sol.) and colony living (col.) females, two averages are given. The tree represents topology only, that is, branch lengths do not indicate evolutionary distance. The topology is based on the phylogeny published in Agnarsson (2006), which is a parsimony analysis of a morphological matrix (43 taxa, 147 characters).

sociations between mother size and egg number in *Stegodyphus* spiders compared to *Anelosimus* spiders, and also why mother size did not correlate with egg size in *Anelosimus*, whereas it did in *Stegodyphus*.

Another difference observed between the social species *A. eximius* and *S. sarasinorum* was the effect of colony size on reproductive output. In *A. eximius*, both egg size and number increased in larger colonies, whereas this effect was lacking in *S. sarasinorum*. Hence, living in larger groups seems to pose benefits to social *Anelosimus* but not to social *Stegodyphus*. It is possible that the extraordinarily high colony sizes reached in *A. eximius* (nest members in the thousands) mean higher assurance of offspring survival due to more allo-mothers and higher colony survival. Females may subsequently allocate more of their body resources into egg laying and less into maternal care. Although subsocial *A. baeza* also showed an increase in egg size in larger colonies,

this was accompanied with a (not significant) decrease in egg number. Subsocial spiders normally live solitarily, and *A. baeza* may not have adapted to group living in the same way as social *Anelosimus* species have. Thus, they may not enjoy the same benefits such as increased egg number when forming groups. However, as these spiders do occasionally occur in groups, perhaps they experience other benefits to group living such as higher protection within colonies leading to higher offspring and colony survival. Other aspects of group living appear to be similar between the social *Anelosimus* and *Stegodyphus*, including higher offspring survival and increased colony-level survival (Aviles and Tufino 1998; Bilde et al. 2007).

## Conclusion

Cooperative breeding is likely to alter optimal maternal strategies. In mammals, birds, and fish, the presence of allo-mothers may allow breeders to produce more clutches, which in turn can affect how females invest in size and number of offspring within clutches (Jennions and Macdonald 1994; Arnold and Owens 1998; Russell et al. 2003; Taborsky et al. 2007). Cooperatively breeding spiders commonly produce only one clutch in their lifetime, and hence need to optimize investment in this one clutch. In accordance with theoretical predictions, we show that cooperatively breeding social spiders within two different genera produce clutches of fewer, larger eggs than those of their subsocial, solitarily breeding congeners representing their ancestral state. We propose that this altered maternal investment strategy may reflect selection for producing larger offspring that are more likely to become reproducers due to a competitive advantage over smaller group members. Finally, we show that clutches of eggs in social and subsocial species showed similar low variation in egg sizes, suggesting that mothers do not induce size variations in their offspring at the egg stage as a bet-hedging strategy. Our study shows how convergent social evolution can shape maternal strategies similarly in phylogenetically distant species, and highlights how permanent group living and cooperative breeding can direct maternal investment from quantity to quality of offspring.

## ACKNOWLEDGEMENTS

We would like to thank R. Balakrishnan and her students as well as Dr. Shibu and the Agastya International Foundation for great help with organization in India, L. Avilés for valuable help and for sharing her expertise in Ecuador, and V. Settepani for precious help in the field and for providing the partial phylogeny of *Stegodyphus*. We thank everyone in the Spider Lab for a stimulation work environment. This study was supported by the Danish Research Council FNU 495997, Oticon Foundation, and School of AGSoS, Aarhus University.

## DATA ARCHIVING

The doi for our data is 10.5061/dryad.t6k57.

#### LITERATURE CITED

- Abramoff, M. D., P. J. Magalhães, and S. J. Ram. 2004. Image processing with ImageJ. Biophotonics Intl. 11:36–42.
- Agnarsson, I. 2006. A revision of the New World eximius lineage of Anelosimus (Araneae, Theridiidae) and a phylogenetic analysis using worldwide exemplars. Zool. J. Linn. Soc. 146:453–593.
- Agnarsson, I., L. Aviles, J. A. Coddington, and W. P. Maddison. 2006. Sociality in Theridiid spiders: repeated origins of an evolutionary dead end. Evolution 60:2342–2351.
- Arnold, K. E., and I. P. F. Owens. 1998. Cooperative breeding in birds: a comparative test of the life history hypothesis. Proc. R. Soc. B Biol. Sci. 265:739–745.
- Aviles, L. 1986. Sex-ratio bias and possible group selection in the social spider Anelosimus-Eximius. Am. Nat. 128:1–12.
- 1997. Causes and consequences of cooperation and permanentsociality in spiders. Pp. 476–498 in J. C. Choe and B. J. Crespi, eds. The evolution of social behavior in insects and arachnids. Cambridge Univ. Press, Cambridge, U.K.
- Aviles, L., and W. Maddison. 1991. When is the sex-ratio biased in social spiders—chromosome-studies of embryos and male meiosis in *Anelosimus* species (Araneae, Theridiidae). J. Arachnol. 19:126– 135.
- Aviles, L., and P. Salazar. 1999. Notes on the social structure, life cycle, and behavior of *Anelosimus rupununi*. J. Arachnol. 27:497–502.
- Aviles, L., and P. Tufino. 1998. Colony size and individual fitness in the social spider *Anelosimus eximius*. Am. Nat. 152:403–418.
- Aviles, L., C. Varas, and E. Dyreson. 1999. Does the African social spider Stegodyphus dumicola control the sex of individual offspring? Behav. Ecol. Sociobiol. 46:237–243.
- Aviles, L., I. Agnarsson, P. A. Salazar, J. Purcell, G. Iturralde, E. C. Yip, K. S. Powers, and T. C. Bukowski. 2007. Natural history miscellany altitudinal patterns of spider sociality and the biology of a new midelevation social *Anelosimus* species in Ecuador. Am. Nat. 170:783–792.
- Bates, D., M. Maechler, and B. Bolker. 2011. lme4: linear mixed-effects models using S4 classes. Available at http://CRAN.R-project.org/package=lme4.
- Bernardo, J. 1996. The particular maternal effect of propagule size, especially egg size: patterns, models, quality of evidence and interpretations. Am. Zool. 36:216–236
- Bilde, T., and Y. Lubin. 2011. Group living in spiders: cooperative breeding and coloniality. Pp. 275–306 in M. E. Herberstein, ed. Spider behaviour, flexibility and versatility. Cambridge Univ. Press, New York.
- Bilde, T., K. S. Coates, K. Birkhofer, T. Bird, A. A. Maklakov, Y. Lubin, and L. Aviles. 2007. Survival benefits select for group living in a social spider despite reproductive costs. J. Evol. Biol. 20:2412–2426.
- Brockelman, W. Y. 1975. Competition, fitness of offspring, and optimal clutch size. Am. Nat. 109:677–699.
- Christenson, T. E. 1984. Behavior of colonial and solitary spiders of the Theridiid species Anelosimus-Eximius. Anim. Behav. 32:725–734.
- Clutton-Brock, T. H., D. Gaynor, G. M. McIlrath, A. D. C. Maccoll, R. Kansky, P. Chadwick, M. Manser, J. D. Skinner, and P. N. M. Brotherton. 1999. Predation, group size and mortality in a cooperative mongoose, *Suricata suricatta*. J. Anim. Ecol. 68:672–683.
- Cockburn, A. 2003. Cooperative breeding in oscine passerines: does sociality inhibit speciation? Proc. R. Soc. B Biol. Sci. 270:2207–2214.
- Crean, A. J., and D. J. Marshall. 2009. Coping with environmental uncertainty: dynamic bet hedging as a maternal effect. Philos. Trans. R. Soc. B 364:1087–1096.
- Crouch, T. E., and Y. Lubin. 2000. Effects of climate and prey availability on foraging in a social spider, *Stegodyphus mimosarum* (Araneae, Eresidae). J. Arachnol. 28:158–168.

- Einum, S., and I. A. Fleming. 2004. Environmental unpredictability and offspring size: conservative versus diversified bet-hedging. Evol. Ecol. Res. 6:443–455.
- Forbes, L. S. 1999. Within-clutch variation in propagule size: the double-fault model. Oikos 85:146–150.
- Fox, C. W., and M. E. Czesak. 2000. Evolutionary ecology of progeny size in arthropods. Ann. Rev. Entomol. 45:341–369.
- Fox, J., and S. Weisberg. 2011. An R companion to applied regression. Sage, Thousand Oaks, CA.
- Gibbs, M., and H. Van Dyck. 2009. Reproductive plasticity, oviposition site selection, and maternal effects in fragmented landscapes. Behav. Ecol. Sociobiol. 64:1–11.
- Gibbs, M., C. J. Breuker, H. Hesketh, R. S. Hails, and H. Van Dyck. 2010. Maternal effects, flight versus fecundity trade-offs, and offspring immune defence in the Speckled Wood butterfly, *Pararge aegeria*. BMC Evol. Biol. 10:345.
- Godfray, H. C. J., L. Partridge, and P. H. Harvey. 1991. Clutch size. Annu. Rev. Ecol. Syst. 22:409–429.
- Gonzaga, M. O., and J. Vasconcellos-Neto. 2001. Female body size, fecundity parameters and foundation of new colonies in *Anelosimus jabaquara* (Araneae, Theridiidae). Insectes Soc. 48:94–100.
- Grinsted, L., and T. Bilde. 2013. Effects of within-colony competition on body size asymmetries and reproductive skew in a social spider. J. Evol. Biol. 26:553–561.
- Hagstrum, D. W. 1971. Carapace width as a tool for evaluating rate of development of spiders in laboratory and field. Ann. Entomol. Soc. Am. 64:757–760.
- Hodge, S. J., M. B. V. Bell, F. Mwanguhya, S. Kyabulima, R. C. Waldick, and A. F. Russell. 2009. Maternal weight, offspring competitive ability, and the evolution of communal breeding. Behav. Ecol. 20:729–735.
- Howe, H. F. 1978. Initial investment, clutch size, and brood reduction in the common grackle (*Quiscalus-Quiscula* L.). Ecology 59:1109–1122.
- Jacson, C. C., and K. J. Joseph. 1973. Life-history, bionomics and behavior of social spider *Stegodyphus sarasinorum* Karsch. Insectes Soc. 20:189– 203.
- Jakob, E. M., S. D. Marshall, and G. W. Uetz. 1996. Estimating fitness: a comparison of body condition indices. Oikos 77:61–67.
- Jennions, M. D., and D. W. Macdonald. 1994. Cooperative breeding in mammals. Trends Ecol. Evol. 9:89–93.
- Johannesen, J., A. Hennig, B. Dommermuth, and J. M. Schneider. 2002. Mitochondrial DNA distributions indicate colony propagation by single matri-lineages in the social spider *Stegodyphus dumicola* (Eresidae). Biol. J. Linn. Soc. 76:591–600.
- Johannesen, J., Y. Lubin, D. R. Smith, T. Bilde, and J. M. Schneider. 2007. The age and evolution of sociality in *Stegodyphus* spiders: a molecular phylogenetic perspective. Proc. R. Soc. B Biol. Sci. 274:231–237.
- Kam, Y. C., C. F. Lin, Y. S. Lin, and Y. F. Tsal. 1998. Density effects of oophagous tadpoles of *Chirixalus eiffingeri* (Anura: Rhacophoridae): importance of maternal brood care. Herpetologica 54:425–433.
- Keller, L., and H. K. Reeve. 1994. Partitioning of reproduction in animal societies. Trends Ecol. Evol. 9:98–102.
- Kokko, H., R. A. Johnstone, and T. H. Clutton-Brock. 2001. The evolution of cooperative breeding through group augmentation. Proc. R. Soc. B Biol. Sci. 268:187–196.
- Kraus, O., and M. Kraus. 1988. The genus *Stegodyphus* (Arachnida, Araneae). Sibling species, species groups, and parallel origin of social living. Verh. Naturwiss. Ver. Hamb. 30:151–254.
- Kullmann, E. J. 1972. Evolution of social behavior in spiders (Araneae— Eresidae and Theridiidae). Am. Zool. 12:419–426.
- Laaksonen, T. 2004. Hatching asynchrony as a bet-hedging strategy—an offspring diversity hypothesis. Oikos 104:616–620.

Lack, D. 1947. The significance of clutch-size. Ibis 89:668-668.

- Lanfear, R., B. Calcott, S. Y. W. Ho, and S. Guindon. 2012. PartitionFinder: combined selection of partitioning schemes and substitution models for phylogenetic analyses. Mol. Biol. Evol. 29:1695– 1701.
- Lubin, Y. 1995. Is there division of labour in the social spider *Achaearanea wau* (Theridiidae). Anim. Behav. 49:1315–1323.
- Lubin, Y., and T. Bilde. 2007. The evolution of sociality in spiders. Adv. Study Behav. 37:83–145.
- Lubin, Y., K. Birkhofer, R. Berger-Tal, and T. Bilde. 2009. Limited male dispersal in a social spider with extreme inbreeding. Biol. J. Linn. Soc. 97:227–234.
- Majer, M., J. C. Svenning, and T. Bilde. 2013. Habitat productivity constraints the distribution of social spiders across continents—case study of the genus *Stegodyphus*. Front Zool. 10:9.
- Marques, E. S. A., J. Vasconcelos-Netto, and M. B. de Mello. 1998. Life history and social behavior of *Anelosimus jabaquara* and *Anelosimus dubiosus* (Araneae, Theridiidae). J. Arachnol. 26:227–237.
- Mattila, T. M., J. S. Bechsgaard, T. T. Hansen, M. H. Schierup, and T. Bilde. 2012. Orthologous genes identified by transcriptome sequencing in the spider genus *Stegodyphus*. BMC Genomics. 13:70.
- Noordwijk, A. J. V., and G. D. Jong. 1986. Acquisition and allocation of resources: their influence on variation in life history tactics. Am. Nat. 128:137–142.
- Parker, G. A., and M. Begon. 1986. Optimal egg size and clutch size—effects of environment and maternal phenotype. Am. Nat. 128:573–592.
- Philippi, T., and J. Seger. 1989. Hedging Ones evolutionary bets, revisited. Trends Ecol. Evol. 4:41–44.
- Platnick, N. I. 2012. The world spider catalog, version 12.5. American Museum of Natural History, New York.
- Pruitt, J. N., and M. C. O. Ferrari. 2011. Intraspecific trait variants determine the nature of interspecific interactions in a habitat-forming species. Ecology 92:1902–1908.
- R Development Core Team. 2011. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
- Ronquist, F., M. Teslenko, P. van der Mark, D. L. Ayres, A. Darling, S. Hohna, B. Larget, L. Liu, M. A. Suchard, and J. P. Huelsenbeck. 2012. MrBayes 3.2 efficient bayesian phylogenetic inference and model choice across a large model space. Syst. Biol. 61:539–542.
- Russell, A. F., and V. Lummaa. 2009. Maternal effects in cooperative breeders: from hymenopterans to humans. Philos. Trans. R. Soc. B 364:1143– 1167.
- Russell, A. F., P. N. M. Brotherton, G. M. McIlrath, L. L. Sharpe, and T. H. Clutton-Brock. 2003. Breeding success in cooperative meerkats: effects of helper number and maternal state. Behav. Ecol. 14:486–492.
- Russell, A. F., N. E. Langmore, A. Cockburn, L. B. Astheimer, and R. M. Kilner. 2007. Reduced egg investment can conceal helper effects in cooperatively breeding birds. Science 317:941–944.
- Rypstra, A. L. 1993. Prey size, social competition, and the development of reproductive division-of-labor in social spider groups. Am. Nat. 142:868– 880.
- Salomon, M., and Y. Lubin. 2007. Cooperative breeding increases reproductive success in the social spider *Stegodyphus dumicola* (Araneae, Eresidae). Behav. Ecol. Sociobiol. 61:1743–1750.
- Salomon, M., J. Schneider, and Y. Lubin. 2005. Maternal investment in a spider with suicidal maternal care, *Stegodyphus lineatus* (Araneae, Eresidae). Oikos 109:614–622.
- Salomon, M., D. Mayntz, and Y. Lubin. 2008. Colony nutrition skews reproduction in a social spider. Behav. Ecol. 19:605–611.
- Sargent, R. C., P. D. Taylor, and M. R. Gross. 1987. Parental care and the evolution of egg size in fishes. Am. Nat. 129:32–46.

Seibt, U., and W. Wickler. 1987. Gerontophagy versus cannibalism in the social spiders *Stegodyphus mimosarum* Pavesi and *Stegodyphus dumicola* Pocock. Anim. Behav. 35:1903–1905.

Slatkin, M. 1974. Hedging ones evolutionary bets. Nature 250:704-705.

- Smith, C. C., and S. D. Fretwell. 1974. Optimal balance between size and number of offspring. Am. Nat. 108:499–506.
- Taborsky, B., E. Skubic, and R. Bruintjes. 2007. Mothers adjust egg size to helper number in a cooperatively breeding cichlid. Behav. Ecol. 18:652– 657.
- Ulbrich, K., and J. R. Henschel. 1999. Intraspecific competition in a social spider. Ecol. Modell. 115:243–251.
- Vollrath, F. 1982. Colony foundation in a social spider. Z. Tierpsychol. 60:313– 324.
- ——. 1986. Eusociality and extraordinary sex-ratios in the spider Anelosimus eximius (Araneae, Theridiidae). Behav. Ecol. Sociobiol. 18:283–287.
- Vollrath, F., and D. Rohde-Arndt. 1983. Prey capture and feeding in the social spider Anelosimus eximius. Z. Tierpsychol. 61:334–340.
- Whitehouse, M. E. A., and Y. Lubin. 1999. Competitive foraging in the social spider *Stegodyphus dumicola*. Anim. Behav. 58:677–688.

### Associate Editor: L. Sundström

## Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's website:

Table S1. Overview of species and sample sizes.

 Table S2. Overview of references and sample sizes used for egg counts in the phylogenies.

Table S3. Matched pairs of social and subsocial species and their mean egg counts used for Wilcoxon test for matched pairs.

Figure S1. Egg number (eggs per egg sac), averaged within colonies, plotted against mother size.

Figure S2. The trade-off between egg size and number for each of the five species.