Sex-biased parental investment is correlated with mate ornamentation in eastern bluebirds

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Males typically have greater variance in reproductive success than females, so mothers should benefit by producing sons under favourable conditions. Being paired with a better-than-average mate is one such favourable circumstance. High-quality fathers can improve conditions for their offspring by providing good genes, good resources, or both, so females paired to such males should invest preferentially in sons. Ornamentation has been linked to male quality in many birds, and, in support of differential allocation theory, females of several avian species invest more in entire broods when paired to attractive mates. Additionally, the females of some bird species apparently manipulate the primary sex ratio of their broods in relation to the attractiveness of their mates. However, empirical support for a link between mate ornamentation and preferential feeding of sons (another form of biased investment) is lacking. We tested for correlations between sex-biased parental investment and mate plumage colour in the eastern bluebird, Sialia sialis, a species in which juveniles have sexually dichromatic UV blue plumage. We found that the proportion of maternal feeding attempts to fledgling sons (versus fledgling daughters) was positively correlated with structurally coloured plumage ornamentation of fathers. Additionally, paternal feeding attempts to sons were correlated with plumage ornamentation of mothers and increased in fathers that had breast plumage characteristics typical of older males. These results provide further support for the idea that parental strategies are influenced by mate attractiveness and provide the first evidence that mate ornamentation can influence parental behaviour even after offspring have left the nest.

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When the benefits of producing male and female offspring vary depending on context, parents are expected to maximize reproductive success by investing differentially in sons and daughters depending on their circumstances (Trivers & Willard 1973; Charnov 1982). Differential investment in sons and daughters can occur by varying the ratio of sons and daughters produced, or through differential investment of energy in sons and daughters after birth or hatching. Animals have been shown to adjust investment in sons versus daughters in relation to season (Dijkstra et al. 1990; Sakisaka et al. 2000; Schultz 2008), diet (Bradbury & Blakey 1998; Opit & Throne 2008), maternal age (Blank & Nolan 1983; Isaac et al. 2005), mate quality (Svensson & Nilsson 1996; Roed et al. 2007) and mate attractiveness (Sheldon et al. 1999). In birds, differential investment in sons and daughters can take the form of primary sex-ratio manipulation via changes in the proportion of male eggs in a given brood. Alternatively, sex-biased investment strategies can manifest as differential resource allocation, which may occur if parents invest time and effort differently in sons and daughters after hatching.

The disparity in the value of males versus females stems from different reproductive opportunities for males and females of different qualities. In many species of animals, males have greater variance in reproductive output than females because the investment in offspring by females is larger than the investment by males (Bateman 1948; Clutton-Brock 1988). Greater variance in male reproductive success arises because poor-quality males are likely to produce fewer offspring than are poor-quality females, whereas high-quality males can produce more offspring than can high-quality females. Therefore, parental investment in sons should be higher if (1) mothers are in good condition (Trivers & Willard 1973), or (2) if sons are fathered by high-quality males (Charnov 1982). Because high-quality males can provide a suite of direct (e.g. increased levels of food provisioning and nest defence; Hoelzer 1989) and indirect (e.g. heritable genetic quality) benefits to their
offspring (Andersson 1994), and because these benefits increase the likelihood that superior offspring will be produced, relative levels of parental investment in sons should also be influenced by the quality of their father.

Total parental investment in birds, typically measured in terms of investment to an entire brood, has been shown to vary with mate ornamentation in several species including blue tits, Cyanistes caeruleus (Limborg et al. 2004), barn swallows, Hirundo rustica (de Lope & Møller 1993), and zebra finches, Taeniopygia guttata (Burley 1988). Female blue tits, for example, feed broods at higher rates when mated to males with bright UV coloration (Limborg et al. 2004) and defend their nests more vigorously than do females paired to males with dull UV plumage (Johnsen et al. 2005). In contrast to these examples, female eastern bluebirds, Sialia sialis, do not feed nestlings at higher rates when mated to highly ornamented males (Siefferman & Hill 2003). Although no evidence has yet been published that supports the relationship between mate attractiveness and sex-biased provisioning for any avian species, bluebird parents may assess the ornamentation of their mates when making feeding decisions within a brood, altering relative investment in sons versus daughters.

We examined the potential for sex-biased parental investment via differential provisioning to sons and daughters in the eastern bluebird, a strongly sexually dichromatic species with sexually dichromatic offspring (Gowaty & Plissner 1998; Siefferman & Hill 2008). Because conspicuous sexual dichromatism is rare in juvenile birds (Kllner 2006) and presents a clear mechanism by which parents can distinguish sons from daughters, bluebirds present an ideal system in which to study sex-biased provisioning for any avian species. Additionally, extrapair paternity is high in some populations of eastern bluebirds (Gowaty & Plissner 1998), potentially providing the reproductive variance between males and females required by the Trivers & Willard (1973) investment theory.

The bright UV blue structural plumage coloration of adult male eastern bluebirds is positively correlated with male provisioning rates to incubating females (Siefferman & Hill 2005a), male provisioning rates to nestlings (Siefferman & Hill 2003), body condition and age (Siefferman et al. 2005). Melanin-based, orange-coloured, breast plumage of males also varies with age, with older males possessing brighter breast feathers with lesser chroma (Siefferman et al. 2005). Because males with more colourful structural plumage feed incubating females and nestlings at higher rates and are generally older and more experienced, we predicted that females would adjust their resource allocation to sons and daughters according to the ornamentation (perceived quality) of their mates. Specifically, females mated to colourful males should increase their investment in sons because having a bright male as a mate should increase the quality of those sons. Because the structural plumage of adult female eastern bluebirds is a condition-dependent trait that varies with food intake and predicts reproductive success in the wild (Siefferman & Hill 2005b), males also stand to benefit by modifying allocation to sons and daughters relative to the coloration of their mates. Thus, we also predicted that adult male bluebirds should increase their investment in sons when mated to more colourful females.

To test these predictions we examined the provisioning rate of bluebird parents to fledgling sons and daughters. When offspring were of fledging age, we placed one daughter and one son in a divided cage and allowed the parents to forage freely. Parents quickly adapted to the trial set-up and provisioned offspring through the wire cage. We analysed feeding rates of parents as a proportion of feeding attempts to sons versus daughters and examined the relationship between this proportion and the plumage ornamentation of each individual’s mate.

**MATERIALS AND METHODS**

**Study Population**

We studied a banded population of eastern bluebirds (hereafter, bluebirds) in Lee County, Alabama, U.S.A. (32°35′N, 82°28′W) between March and August 2008. Bluebirds are a sexually dimorphic passerine species (family Turdidae) that breeds throughout eastern North America (Gowaty & Plissner 1998). Adult male bluebirds have rich blue coloration on their heads, backs, rumps, tails and wings. The upper breasts of adult males are orange, and the bellies are white. Adult females have blue–grey upper parts with dull blue wings and tails and pale orange breasts.

We monitored nestboxes throughout the breeding season to determine when nests were in use by bluebirds, as well as to monitor the age and development of bluebird offspring. Bluebirds typically begin laying during the month of March in central Alabama and can produce up to three broods, averaging approximately four eggs per clutch (7 ± SE = 3.75 ± 1.1; Siefferman & Hill 2007), over the course of the breeding season. Nestlings typically hatch after 14 days of incubation and fledge 15–20 days after that (Gowaty & Plissner 1998).

**Trial Procedures**

Because fledglings spend much of their time hidden, it is typically difficult to observe patterns of parent–offspring interactions after young have fledged. In our study, we constrained the movements of fledging bluebirds to an observation arena while simultaneously allowing their parents to forage normally. Between 16 and 18 days of age, we selected one male and one female nestling from nests that contained one male nestling and at least one female nestling. We determined the sex of fledglings using plumage characteristics in the field, but later performed molecular techniques that allowed us to double-check our field assignments. Eastern bluebird nestlings typically fledge near this stage of development (Gowaty & Plissner 1998), and we chose specific trial dates on a brood-by-brood basis depending on the development and size of nestlings in each nestbox. When broods contained multiple female nestlings, we selected the individual with the mass closest to that of the male nestling.

On the day of the trial, we gathered all nestlings from each box to measure their mass. To minimize the effects that different hunger levels might have on fledging begging rate and intensity, as well as the effects that such differences might have on parental provisioning patterns, we fed each nestling one mealworm (Tenebrio molitor) before all members of the brood were returned to the natal nestbox. At this point, we sealed the entrance to the nestbox to prevent any feeding by parents, as well as any premature fledging attempts. We then left the immediate area for 30 min to allow the nestlings to digest the recently consumed mealworm. After the 30 min pretrial period, we returned to the nestbox, selected the predetermined male and female, and placed them separately in a divided wire cage near the natal nestbox (Fig. 1). A solid partition prevented physical and visual contact between siblings in the wire cage. To create a location from which bluebird parents could assess their offspring, we placed a 50 cm tall perch 1 m away from the front of the cage. We kept all remaining nestlings in a cloth box and fed them mealworms throughout the duration of the trial.

Parent bluebirds quickly adjusted to the trial set-up and began to feed their offspring through the wire mesh of the cage. We used a tripod mounted video camera (Sony Hi-8) to record parent and chick interactions for 1 h, at which point we reversed the position of the fledglings in the cage (to control for possible effects of cage location) and resumed recording for one additional hour. After each trial, we removed the fledglings from the cage and returned them to their nestbox along with their siblings.
We quantified parental investment from videotapes without knowledge of the sex of the fledglings or the identity of the parents in each trial. Although we were unable to reliably assess the size of food items brought to fledglings during the trial, previous research indicates that prey size does not vary with feeding rate in this population (Siefferman & Hill 2007). Food handling and transfer difficulties between parents and offspring, exacerbated by the wire mesh separating them, often caused parents to temporarily abandon feeding one fledgling and begin attempting to feed the other fledgling. Because of the inconsistency of food transfer, we used long-distance parental approaches to juveniles as a proximate measure of investment. Every time an adult directly approached one of the juveniles from outside the frame of the video screen, or from the perch 1 m in front of the cage, we scored the event as a feeding attempt. This scoring method best captured the choices that parents made when delivering food and it minimized the effects that delivery complications and fledgling behaviour had on parental feeding decisions. In another study using similar methodology (R.A. Ligon & G.E. Hill, unpublished data), begging by juveniles prior to parental approach was rare (twice in 366 feeding attempts), limiting the effect of this behaviour on feeding decisions by parents.

In total, we recorded 49 feeding trials throughout the course of the breeding season. However, five of these trials were conducted with bluebird parents that had already been tested earlier in the year. We excluded these duplicate trials and used each set of bluebird parents only once in our analyses. Additionally, we were forced to exclude several trials from analysis because we failed to capture either the mother or the father bluebird. These failures precluded analysis of plumage colour for these individuals. One additional trial was excluded from analysis because the sex assigned to fledglings in the field was later determined to be incorrect using molecular sexing techniques.

**Colorimetric Data**

We plucked feathers from adult bluebirds in March and April 2008, and measured plumage colour from these feathers using an S-2000 spectroradiometer with a deuterium-halogen lamp (Ocean Optics, Dunedin, FL, U.S.A.) following the procedures described in Siefferman & Hill (2003). When adults were captured, we collected nine breast feathers, nine rump feathers and two outer tail feathers from each bird. Feather samples were plucked from the same location on all birds and were later placed on black paper for spectrophotometric analysis. Breast and tail feathers (contour feathers) attain colour by superposition of several feathers (sensu Quesada & Senar 2006), so we arranged them in a manner that mimicked the way these feathers lay naturally on bluebird bodies. One researcher (R.A.L) then recorded spectral data using a micron fibre optic probe at a 90° angle to the feather surface.

We used the spectral processing program Colour (v1.7, Queens, Ontario, Canada) to calculate three standard descriptors of reflectance spectra: UV chroma, hue and brightness. Brightness (total amount to light reflected by the feather) is the summed reflectance from 300 to 700 nm. Chroma was calculated differently for UV blue and chestnut coloration because of the inherent reflective properties of the two colours. For the rump and tail feathers, UV chroma, a measure of spectral purity, was calculated as the proportion of reflectance within the UV range to the total reflectance (R\textsubscript{300–400}/R\textsubscript{300–700}). For the chestnut breast feathers, red chroma was calculated as the ratio of the total reflectance in the red range to the total reflectance of the entire spectrum (R\textsubscript{675–700}/R\textsubscript{500–700}). Hue is the principal colour reflected by the feather. For structural coloration (rump and tail), hue was defined as the wavelength of maximum reflectance (λ\textsubscript{R}\textsubscript{max}). Because hue (calculated as maximum slope) of the chestnut breast feathers expressed very little variation among males, we do not report hue for breast coloration.

Although structural plumage colour varies seasonally in blue tits (Örnborg et al. 2002), no such relationship has been found in eastern bluebirds (Siefferman & Hill 2005c). This fact, coupled with the relatively short period (<2 months) over which adults were captured, suggests that effects of seasonality on our analyses in the present study were minimal.

Using Principal Components Analysis to Describe Plumage Colour

We performed separate principal components analyses (PCA) on measures of breast, rump and tail coloration. We used PCA analysis because the plumage characteristics (brightness, chroma, hue) of each body region were correlated and because this analysis enabled us to reduce the number of colour variables to a more manageable number (from eight plumage variables to three). The results of the principal components analyses are summarized in Table 1. The first principal component captured, suggests that effects of seasonality on our analyses in the present study were minimal.

**Male plumage**

The melanic breast coloration of males (brightness and chroma) reduced to one principal component that explained 64% of the variation among these variables (brightness loading, 0.80; red chroma loading, −0.80). A male with a high positive PC1 score for breast plumage had brighter feathers and less-red chroma and was, therefore, less ornamented. The first principal component for structural rump coloration explained 64% of the variation among these variables and received strong loadings from brightness, hue and UV chroma (0.45, −0.93 and 0.92, respectively). An individual with a high positive PC1 score for rump colour was more ornamented with brighter feathers, left-shifted hues and greater UV chroma. The first principal component for tail coloration, which explained 60% of the variation among these variables, also received strong loadings from brightness, hue and UV chroma (0.73, 0.73, −0.86). An individual with a high positive PC1 score for tail colour displayed bright feathers with right-shifted hues and decreased UV chroma (less ornamented).
Distribution and logit link functions in our models, as is typical in biological investigations of binary data using generalized linear models (Donazar et al. 1993; Bustamante 1997; Martinez et al. 2003). Differences in sample sizes (e.g. number of feeding attempts per trial) are also accounted for in generalized linear models, an additional benefit of using this approach (Wilson & Hardy 2002). Using these models we were able to examine how the plumage coloration of the parents (their mates and themselves), the number of feeding attempts to sons by each parent’s mate, the number of offspring in each brood, the difference in mass between sons and daughters, and the age of the nestlings influenced the proportion of feeding attempts to sons. Our final model selection was based on Akaike’s Information Criterion (AIC) values wherein models with the lowest AIC values were considered the most parsimonious. For the sake of simplicity, we plotted only the relationships between parental feeding behaviour and the most significant variable in each model.

**Ethical Note**

We began each trial early in the morning to minimize the time that fledglings were exposed to high temperatures, and we returned all juveniles to their nestboxes within 2.5 h of the start of each trial. Field observations of fledglings following preliminary trials indicated that our caging protocol did not influence survival to the fledging stage. This study was approved by the Auburn University Internal Animal Care and Use Committee (IACUC project registration no. 2008-1341) and conducted under Alabama State and U.S. Fish and Wildlife permits.

**RESULTS**

During our study trials, mothers fed their offspring significantly more than fathers did (paired t test: \( t_{33} = 2.82, P < 0.01 \)). Mothers fed their offspring a mean \( \pm \) SD of 25.47 \( \pm \) 22.48 times/trial \( (N = 43) \), while fathers averaged only 14.60 \( \pm \) 12.69 feeds/trial.

The generalized linear model with the lowest AIC value, built using data from 34 independent trials, included five predictive variables that explained a significant amount of variation in the proportion of feeding attempts that mother bluebirds directed towards their sons (Table 2). Specifically, we found significant relationships between the proportion of mothers’ feeding attempts directed towards sons and the first principal component (PC1) of father’s rump coloration, PC1 of father’s tail coloration, PC1 of father’s breast color, the mother’s own PC2 for tail coloration and the number of offspring in the brood (Table 2). Females mated to...
males with more highly ornamented rumps (brighter, left-shifted hues, increased UV chroma) increased their investment in sons (Fig. 2), as did females mated to males with greater tail ornamentation (left-shifted hues, increased UV chroma; not shown). Females also increased investment in sons when they were mated to males with brighter, less-red breast plumage, when they themselves (mothers) had darker tails, and when brood sizes were smaller (Table 2).

The generalized linear model with the lowest AIC value for paternal feeding behaviour, constructed using data from 34 feeding trials, contained only two predictive variables that explained a significant amount of variation in the proportion of feeding attempts that father bluebirds directed towards their sons (Table 3). Fathers fed sons relative to maternal breast ornamentation and in relation to their own breast plumage. Males mated with females with redder, darker breast plumage fed their sons more than their daughters (Fig. 3), as did males with brighter, less-red breast plumage (not shown).

**Assortative Mating**

None of the correlation coefficients between colour patches of mated fathers and mothers was statistically significant after we corrected the probability values for simultaneous tests by using the Bonferroni procedure (Table 4). The lack of assortative mating with respect to plumage ornamentation agrees with the results previously obtained examining this possibility in the same study population at an earlier time (Siefferman & Hill 2005b).

**DISCUSSION**

Although the potential benefits associated with sex-ratio manipulation in birds are numerous (Hasselquist & Kempenaers 2002), support for this possibility has not been universal (e.g. Saino et al. 1999; Dreiss et al. 2006). Nevertheless, we found a significant relationship between the plumage ornamentation of adult bluebirds and the parental feeding decisions of their mates. Specifically, females mated to males with more ornamented rump and tail feathers increased their feeding attempts to fledgling sons relative to daughters. Maternal investment in sons also increased when females were mated to males with brighter, less-red breast plumage, when mothers had darker tail feathers and when broods were smaller. Additionally, paternal feeding decisions appear to be influenced by the breast coloration of their mates such that males mated to females with redder, darker breast plumage showed increased provisioning effort to sons. Taken together, these results support our predictions regarding the influence of mate ornamentation on parental feeding decisions.

Female eastern bluebirds should benefit from choosing brighter, generally older, males as mates because of the increased provisioning investment provided by such males (Siefferman & Hill 2003) and the inherent quality of males that have survived to an advanced age (Kokko & Lindström 1996). In this study, female bluebirds behaved as if they assessed the plumage colour of their mates when making reproductive decisions. Mothers fed sons at higher rates when mated to males with highly ornamented structural plumage (increased UV chroma, left-shifted hues) and increased feeding attempts to sons when mated to males with melanin-based breast plumage qualities typical of older males (brighter plumage, less-red chroma; Siefferman et al. 2005). However, experimental mate choice studies have shown that female bluebirds do not consistently choose males with brighter

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**Table 2**

Optimized generalized linear model (lowest AIC value) for the proportion of feeding attempts that eastern bluebird mothers directed towards sons (out of their total feeding attempts)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate</th>
<th>SE</th>
<th>Wald 95% CI</th>
<th>x²</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>0.55</td>
<td>0.25</td>
<td>0.05, 1.04</td>
<td>4.74</td>
<td>0.03</td>
</tr>
<tr>
<td>Father rump PC1*</td>
<td>0.19</td>
<td>0.07</td>
<td>0.06, 0.32</td>
<td>8.66</td>
<td>&lt;0.01</td>
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<tr>
<td>Father tail PC1*</td>
<td>-0.16</td>
<td>0.06</td>
<td>-0.28, -0.04</td>
<td>7.25</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Father breast PC1*</td>
<td>0.21</td>
<td>0.08</td>
<td>0.05, 0.38</td>
<td>6.51</td>
<td>0.01</td>
</tr>
<tr>
<td>Mother tail PC2</td>
<td>-0.16</td>
<td>0.07</td>
<td>-0.29, -0.03</td>
<td>5.85</td>
<td>0.02</td>
</tr>
<tr>
<td>Brood size</td>
<td>-0.16</td>
<td>0.07</td>
<td>-0.29, -0.03</td>
<td>5.74</td>
<td>0.02</td>
</tr>
</tbody>
</table>

* Principal component plumage colour scores for the respective body regions.

**Table 3**

Optimized generalized linear model (lowest AIC value) for the proportion of feeding attempts that eastern bluebird fathers directed towards sons (out of their total feeding attempts)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate</th>
<th>SE</th>
<th>Wald 95% CI</th>
<th>x²</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>-0.12</td>
<td>0.10</td>
<td>-0.31, 0.07</td>
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<td>0.22</td>
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<tr>
<td>Mother breast PC1*</td>
<td>-0.24</td>
<td>0.09</td>
<td>-0.42, -0.06</td>
<td>7.04</td>
<td>0.01</td>
</tr>
<tr>
<td>Father breast PC1*</td>
<td>0.22</td>
<td>0.10</td>
<td>0.03, 0.40</td>
<td>5.20</td>
<td>0.02</td>
</tr>
</tbody>
</table>

* Principal component plumage colour scores for the respective body regions.

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**Figure 2**

Relationship between paternal rump colour (PC1) and the proportion of maternal feeding attempts to sons. Higher PC1 scores indicate increased ornamentation. To facilitate a more accurate interpretation of the relative weights of each trial to the final model, symbol sizes are proportional to the number of maternal feeding attempts in each trial.

**Figure 3**

Relationship between maternal breast colour (PC1) and the proportion of paternal feeding attempts to sons. Lower PC1 scores indicate increased ornamentation. Symbol sizes are proportional to the number of paternal feeding attempts in each trial.
plumage (Liu et al. 2007, 2009). The lack of preference for traits that apparently signal multiple aspects of male quality (Siefferman & Hill 2003, 2005a, 2005c) presents something of a conundrum. One possible explanation is that females assess territory quality rather than male ornamentation when choosing mates. If highly ornamented males hold higher-quality territories (Siefferman & Hill 2005c), then the observed relationship between paternal ornamentation and maternal investment in sons could arise if females make parenting decisions based solely on habitat quality. Only by experimentally manipulating adult plumage can we be entirely sure of the direct impact of male plumage on parental feeding decisions, and such manipulations are an obvious next step in examining the factors influencing sex-biased provisioning.

In this study, bluebird mothers varied investment in sons and daughters relative to the ornamentation of their mates, but females' provisioning decisions were also tied to their own tail coloration. Mothers with darker (less bright) tail feathers increased provisioning attempts to sons. Given that the brightness of structural plumage is condition dependent in female eastern bluebirds (Siefferman & Hill 2005b), this relationship runs counter to predictions of the Trivers & Willard (1973) hypothesis. Females in poor condition (less ornamented plumage) should invest more in daughters (Trivers & Willard 1973), given the lower variance in reproductive success for daughters relative to sons. At this point, we have no simple explanation for why drab females feed sons more.

We predicted that bluebird fathers, like bluebird mothers, would adjust provisioning to sons and daughters relative to the ornamentation of their mates. Indeed, we found that fathers increased investment in sons when mated to females with darker, redder breast plumage, a finding supporting our general prediction. Melanin-based plumage has frequently been found to serve a signalling role in antagonistic intraspecific interactions (Senar 2006), although no such function has been shown in eastern bluebirds. If, in fact, melanin breast plumage of female bluebirds conveys information about how individuals fare in intrasexual competition (for mates or resources), then males that assess such signals when making offspring investment decisions might increase their own fitness. Contrary to our findings regarding melanin-based plumage, we found no relationship between the structural plumage colour of mothers and the feeding decisions of fathers, despite the fact that such plumage appears to be condition dependent in female bluebirds (Siefferman & Hill 2005b). However, fathers performed significantly fewer feeding attempts per trial than mothers, suggesting that males follow a different overall provisioning strategy compared to females. The contribution of eastern bluebird fathers to fledglings may come primarily in the form of nest and territory defence (e.g. Rytkönen et al. 1993; Hogstad 2005; Gibson & Moehrenschlager 2008).

An advantage of selective feeding is that offspring can be fed differentially throughout the nesting period, parents should benefit by becoming increasingly selective about which of their offspring receive food. If costliness of provisioning offspring increases choosiness by parents with respect to which offspring they feed, then the increased selectivity shown by provisioning bluebird mothers (i.e. the correlation between paternal ornamentation and maternal investment in sons) fits with previously discovered sex-specific costs of reproduction in bluebirds, where costs of reproduction are greater for mothers than for fathers (Siefferman & Hill 2008).

While the potential mechanisms of primary sex-ratio manipulation in birds are poorly understood (reviewed in Pike & Petrie 2003), the mechanisms by which parental behaviours can influence sex allocation and sex ratio are relatively straightforward. To change secondary offspring sex ratio, parents can (1) selectively destroy offspring of one sex at an early stage (Charnov 1982), (2)
selectively incubate eggs of different sexes (Ligon & Ligon 1990; Pike & Petrie 2003) or (3) give more food to offspring of a particular sex (Charnov 1982). There are obvious costs to the first and second strategies, namely the loss of energy and time expended in producing a fertile egg and/or incubating it. However, the third strategy is plastic and allows parents to match allocation decisions with current environmental conditions. Such a benefit might be particularly important for species with extended parental care, when the optimal investment strategy (sons versus daughters) may change through time.

Ornamental traits have been a focus of interest among evolutionary biologists since the discussions of Darwin and Wallace (Cronin 1991), but most studies of such traits in birds have focused on ornaments in the context of female mate choice (Hill 2006) or male–male competition (Senar 2006). However, if ornamental traits are condition-dependent signals of quality, then they should be assessed in contexts other than mate choice and competition. Here we show evidence that male and female bluebirds assess mate quality via expression of plumage coloration as a means to optimize resource allocation among offspring. It seems probable that assessment of condition-dependent ornaments occurs across a wide range of contexts, but additional experimental tests are required to confirm such possibilities.

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