The Grey-backed Shrike parents adopt brood survival strategy in both the egg and nestling phases

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Abstract

Background: Great diversity exists in the parenting pattern of altricial birds, which has long been considered as an adaptive response to specific environmental conditions but not to their life-history style.

Methods: We examined the egg-laying and nestling-raising pattern of the Grey-backed Shrike (Lanius tephronotus) that breeds only once a year on the Tibetan Plateau. We compared the dietary composition to that of its sympatric competitor, the Brown-cheeked Laughing Thrush (Trochalopteron henrici) that breeds twice a year.

Results: Female Grey-backed Shrikes produced a fixed clutch size of five, with increasing egg size by their laying sequence. The last offspring in the brood is disadvantageous in the size hierarchy because it hatches later. However, they had the largest fledgling body mass. These findings indicate that Grey-backed Shrikes adopt the brood survival strategy in both the egg and nestling phases. Moreover, males and females exhibit no sexual division in providing parental care as they made an equal contribution to the total amount of food delivered to their brood. This parenting pattern of Grey-backed Shrikes, as well as their dietary items, differ significantly from those of the Brown-cheeked Laughing Thrush.

Conclusions: We suggest that the differentiation in life-history style between sympatric competitors, rather than a behavioral response to specific environmental conditions, plays a decisive role in driving avian parenting strategy diversification.

Keywords: Brood survival strategy, Food types, Lanius tephronotus, Parenting pattern, Provisioning rate

Background

In altricial birds, parenting behaviors are critical for the survival of dependent offspring (Silver et al. 1985; Tal-lamy and Wood 1986) while they are costly for parents due to their time- and energy-consuming (Skutch 1949). It has long been suggested by the life-history theory that parents should optimize their parenting patterns under a specific environmental condition in a way that would maximize their reproductive fitness (Lack 1954). Empirical tests of the life-history theory have revealed an astonishing diversity in altricial birds’ parenting care patterns (Winkler and Wallin 1987; Martin 1995; Sæther and Bakke 2000; Eikenaar et al. 2003; Martin et al. 2011; Du et al. 2014). These patterns are generally considered as an adaptive response to various environmental conditions, such as food availability and nest predation risk (Martin 1995; Caro et al. 2016). However, few studies have attached importance to the differentiation in life-history style in driving avian parenting behaviors to diversify.

One fundamental characteristic of life-history style is the time that an individual can breed a year. Most
passerine species, only when the length of breeding season supports multiple broods could an individual select to breed one or more times (Verhulst and Nilsson 2008; Camfield et al. 2010; Du et al. 2014; Li et al. 2020a). Single- and multi-brooded breeders exhibit different tradeoffs in their parenting patterns. Multi-brooded individuals often adopt different strategies across broods. It will be beneficial for breeders to adopt a brood reduction strategy when the breeding conditions are poor, i.e. biasing investment towards stronger offspring, to ensure the lowest nest survival threshold (Mock and Forbes 1994; Forbes et al. 2001). When breeding conditions are better, it will be beneficial for breeders to adopt a brood survival strategy, i.e. biasing investment towards weaker offspring, so that they can raise many offspring as possible (Slagsvold et al. 1984, 1997; Forbes 2007). In contrast, single-brood species are more likely to adopt the brood survival strategy regardless of the breeding conditions, as they have only one chance to realize their fitness (Du et al. 2012; Li et al. 2020a). A comparison between sympatric single- and multi-brooded species’ parenting strategies would help understand the role of life-history style in the diversity of avian parenting behaviors.

Altricial birds have two important breeding phases, the egg and the nestling, to optimize their parenting strategies. In the egg phase, parents trade off the eggs’ number and size mainly based on the expected amount of available food (egg size strategy; Slagsvold et al. 1984; Christians 2002); whereas, they select their brood provisioning strategy based on the actual amount of food in the nestling phase (Decker et al. 2012). By modulating an egg’s size according to its laying sequence, female birds can affect the size hierarchy among newly-hatched offspring, which in turn affects the intensity of subsequent sibling rivalry (Shizuka and Lyon 2013; Mainwaring et al. 2014). When the egg size decreases with the laying sequence, as in the Giant Babax (Babax wadelli), the last hatching is smaller than its older siblings and hence at a disadvantage in the competition over parental investment within the brood (Du et al. 2012). In contrast, when the egg size increases with the laying sequence, as in the Azure-winged Magpie (Cyanopica cyanus), later hatchlings are larger at hatching than their older brood-mates. This can compensate, to some extent, for the disadvantage faced by later offspring in competing with their older siblings (Da et al. 2018). In the tradeoff between nestlings’ number and size, parents in many cases adopt different parenting strategies from that in egg-laying, particularly in an environment where the breeding conditions are unpredictable (Decker et al. 2012). For example, parents of the Giant Babax adopt the “brood reduction” strategy in laying eggs and the “brood survival” strategy in provisioning the nestlings (Du et al. 2012); while parents of the Horned Lark (Eremophila alpestris) adopt the “brood survival” strategy in laying eggs and the “brood reduction” strategy in provisioning the nestlings (Du et al. 2014; Da et al. 2018). The difference in individual tradeoffs between egg-laying and nestling-provisioning has become a common explanation for parenting pattern evolution, whereas the effect of life-history style has been largely neglected.

In this study, we addressed the role of life-history style in the evolution of parenting patterns in the Grey-backed Shrike (Lanius tephronotus), which is a small (approximately 40 g), carnivorous bird with no sexual dimorphism neither in size nor in plumage. It is the only Lanius species that can breed at the high elevation of the Tibetan Plateau (ranging between 2700 and 4500 m) (Lu et al. 2010). Generally, Grey-backed Shrikes produce only one brood a year. In a population distributed in their upper range limit (4000–4500 m), the clutch size (ranging from 3 to 5) decreases significantly with the elevation (Lu et al. 2010). In contrast, in the population distributed in the lower range limit on the Tibetan Plateau (2600–2900 m), females tend to produce a fixed clutch size of five (B. Du, unpublished data). It seems that Grey-backed Shrikes have made adaptive responses to the variation of breeding conditions; hence, it might be an ideal system to compare individual tradeoffs between egg-laying and nestling-provisioning. Several shrub-nesting bird species, such as the White-collared Blackbirds (Turdus albocinctus) (Fan et al. 2017) and the Brown-cheeked Laughing Thrush (Trochalopteron henrici) (Li et al. 2020b), are sympatric in the Grey-backed Shrike’s lower range limit. Both species mix with the Grey-backed Shrike in their nesting and foraging sites, but can breed twice a year. Therefore, it becomes possible to compare the parenting strategy of sympatric species, to identify the relative role of life-history style in the evolution of parenting behavior evolution.

To address whether Grey-backed Shrikes adopt different strategies between the egg and nestling phases, we first identified their egg-laying pattern and the growth pattern of nestling body mass with different hatching sequences. Then, we also performed a dietary investigation and food types between the Grey-backed Shrike and Brown-cheeked Laughing Thrush to examine the life-history style’s effect on the differentiation of parenting patterns.

Methods
Study area and population
This study was carried out in the Bayi town, Tibetan Autonomous Region, China (29°40′N, 94°20′E, mean altitude of 2900 m), during 2015–2019. This region has a typical cold (mean annual temperature 7 °C) and wet
(annual precipitation 500–700 mm) high-altitude climate. The temperature and precipitation change greatly through the year, with the highest temperatures and rainfall occurring between June and August (Fan et al. 2017). Local landscape is characterized by the evergreen Chuan-Dian Alpine Oak (Quercus aquifolioides) forest, mixed with some deciduous trees, such as the Aspen (Populus davidiana) and Tibetan Willow (Salix insignis), and shrubs composed mainly of the Lhasa Berberis (Berberis hemsleyana), roses (Rose spp.), azaleas (Rhododendron spp.), and powder-branched berry (Rubus biflorus). Our study area of 300 ha is located along the Niyang River, where Grey-backed Shrikes choose to build most of their nests in the shrubs.

Grey-backed Shrikes breed once a year in our study area. At the end of June, pairs start occupying territories and build their nests in the shrubs. Nest construction is carried out mainly by the female, while the male defends the territory. The outmost layer of a nest comprises small branches, withered grass, and plastic sheeting pieces, while the inner layer is lined with fine grass stems and animal hair. Females lay their eggs immediately after the nest is constructed. After the first egg is laid, the female would brood the nest, but only at night; after the penultimate egg was laid, all-day incubation is commenced. After that, both sexes contribute to the incubation. Hatching asynchrony occurs in the Grey-backed Shrike, with three or four chicks hatching in the first day and the remaining on the second day. Both sexes contribute to provisioning nestlings during the nestling period and approximately one month after they fledge.

**Data collection for reproductive parameters**

Data collection for reproductive parameters began with a systematic search for the Grey-backed Shrike nests at the end of June. The nest contents were checked daily to determine the clutch initiation date (the date when the first egg was laid) and the laying sequence of each egg. The laying sequence was marked on its eggshells with a non-toxic marker pen (Deli Company, Guangzhou, Guangdong Province, China). The fresh mass as an index for its size, was measured with an electronic balance to the nearest 0.1 g. The hatching sequence was marked on the chicks’ heads at hatching, and their body mass was measured (to the nearest 0.1 g). When two or more nestlings hatched on the first day, their hatching sequences could be determined according to their skin color. The darker the color, the earlier a nestling hatched. During the nestling period, nest content was checked every two days to measure the nestlings’ body mass. When the nestlings reached 30 g or were older than ten days, they were leg-banded with one numbered metal ring and two colored plastic rings. Nesting success was considered achieved when a social pair fledged at least one offspring.

Adults were captured after the nestlings have hatched, using a method that had been successfully adopted to capture other shrub-nesting species, such as the Azure-winged Magpie (Ren et al. 2016) and the White-collared Blackbird (Fan et al. 2017). Capturing the adults and measuring the nestlings were performed under the permission of the Tibetan Forestry Department (2016ZRY-NY-05). Each captured individual was sexed by the presence of a brooding patch, leg-banded with one numbered aluminum ring and two colored plastic rings, and weighed. Only one parent was captured in most nests to minimize the disturbance on parenting behaviors. This procedure fulfilled the parents’ sex identification requirement.

A dietary investigation was performed during the nestling period to examine the type and size of food that Grey-backed Shrikes delivered to their offspring. First, adults foraging behaviors were monitored to determine their foraging sites. Then, different types of food were sampled at the foraging sites by searching for insects on the ground and in the earth, and by gathering fruits berries as, in some cases, adults were found to feed on such plant food. The different types of food were weighed and classified based on their mean mass (Additional file 1: Table S1). These were considered candidate dietary items that parents might deliver to their offspring.

**Data collection for parental provisioning behaviors**

After adults were leg-banded, their provisioning behaviors were recorded automatically by digital camcorders (ZX1, Eastman Kodak Company, Rochester, NY, USA) for 3 h (9:00–12:00 a.m., China Standard Time) every two days. Each camcorder was mounted on a tripod that was fixed diagonally 0.8–1 m above a nest. The recording process caused the Grey-backed Shrike no adverse effects as there were no nest abandonment cases during the recording periods. A total of 256 h of adult provisioning behavior recordings were obtained (13.5 ± 0.7 h per nest, \( n = 19 \) nests).

Data on parental provisioning behaviors were extracted from videos by playing them back on a computer. This dataset included: (1) identity of the nest-visitor and whether it delivered food to the brood; (2) the type, number, size of food a provisioner delivered to the nestlings; (3) the predators’ species that were monitored preying on eggs or nestlings. Based on these data, an individual’s provisioning rate was calculated as the number of feeding bouts per hour; the food types and sizes identified in the video were assigned to the candidate food list (Additional file 1: Table S1), so that the amount of food a parent delivered to the brood per feeding bout could be calculated.
Statistical analysis
A generalized linear mixed model (GLMM) was fitted to test factors that might influence the fresh egg mass set as a dependent variable, with identical link function (Table 1). Fixed factors included the clutch initiation date and the egg laying sequence. The clutch size was not included in the model because it seemed stable among the nests. Random effects included the year and nest identity. A GLMM was also fitted to test factors that might influence the nestlings’ body mass set as a dependent variable, with identical link function (Table 2). Fixed factors included the clutch initiation date, nestling age, the hatching sequence. Random effects included the year, nest identity, and nestling identity. Factors that might influence the provisioning rate of males and females set as a dependent variable, with identical link function (Table 3), were tested by fitting two more GLMMs. Fixed effects included brood size and nestling age, and random effects included the year and nest identity. Similarly, factors that might influence the food amount delivered by males or females per feeding bout (set as dependent variable), with identical link function (Table 4) were tested by fitting GLMMs. Fixed effects included brood size, nestling age, the breeder’s provisioning rate, and random effects included the year and nest identity. During the process of fitting GLMMs, we did not introduce parental body conditions into the model because parents were captured in different nestling ages, and in most nests, only one parent was captured. As a substitute, we performed variance component analysis (VCA) to examine the relative contribution of between-nest difference (i.e. the random effect of nest identity) to the variance in each dependent variable. Multiple linear regression was used to test the multicollinearity of fixed effect variables before fitting the GLMMs. Variables were considered to have serious multicollinearity when their variance inflation factor (VIF) was larger than three.

A logistic model was fitted to monitor the nestlings’ growth pattern based on their body mass, using non-linear regression (Huin and Prince 2000): \[ W = K \]

### Table 1 Factors that might influence the egg's fresh mass in the Grey-backed Shrike

| Generalized linear mixed model parameters | Fixed effects | β ± SE | n  | t     | P    |
|------------------------------------------|--------------|-------|----|-------|------|
| Intercept                                | 4.274 ± 0.094| 151   | 45.618 | <0.001 |
| Clutch initiation date                   | –0.002 ± 0.003| 151 | –0.555 | 0.580 |
| Laying sequence                          | 0.058 ± 0.016| 151 | 3.697   | <0.001 |

| Random effects                           | β ± SD | n  | Results of VCA (%) |
|------------------------------------------|-------|----|---------------------|
| Nest identity                            | 0.097 ± 0.311 | 151 | 66.90               |
| Years                                    | 0.001 ± 0.001 | 151 | 0.69                |
| Residual                                  | 0.047 ± 0.217 | 151 | 32.41               |

SE of fixed effects is the standard error of the mean; SD of random effects is the square root of the variance. The explanations apply also to Table 2, 3 and 4.

### Table 2 Factors that might influence the nestling's body mass in the Grey-backed Shrike

| Generalized linear mixed model parameters | Fixed effects | β ± SE | n  | t     | P    |
|------------------------------------------|--------------|-------|----|-------|------|
| Intercept                                | 1.620 ± 1.086| 284   | 1.492 | 0.065 |
| Clutch initiation date                   | 0.042 ± 0.048| 284   | 0.870 | 0.284 |
| Hatching sequence                        | –0.186 ± 0.164| 284 | –1.134 | 0.101 |
| Nestling age                             | 2.264 ± 0.054| 284   | 42.234 | <0.001 |

| Random effects                           | β ± SD | n  | Results of VCA (%) |
|------------------------------------------|-------|----|---------------------|
| Nestling identity                        | 0.001 ± 0.001 | 284 | 0.006               |
| Nest identity                            | 3.694 ± 1.922 | 284 | 21.25               |
| Year                                     | 3.032 ± 1.741 | 284 | 17.44               |
| Residual                                  | 10.659 ± 3.265 | 284 | 61.31               |
In this equation, \( W \) is the nestling’s body mass, \( K \) is the asymptotic body mass that a fledgling could reach, \( a \) is the nestling’s exponential growth initiation date, \( b \) is the instantaneous growth rate, and \( A \) is the nestling age. The fledgling body mass of nestlings was compared by one-way ANOVA based on their hatching sequence.

The different food types frequencies delivered by parents to their offspring were tested to examine whether they were distributed evenly, using the one sample Kolmogorov–Smirnov test.

All analyses were conducted using SPSS (version 21.0; IBM Corp, Armonk, NY, USA) and R (version 3.3.4). Descriptive results are presented as mean ± standard error (SE). The null hypothesis was rejected when \( P < 0.05 \), and reported probabilities are two-tailed.

### Results

Over four years (2015–2018), 59 Grey-backed Shrikes’ nests were observed to complete their clutches in our study area. Females tended to produce a fixed clutch size of five (97% of the nests, 57/59). The mean fresh egg mass was 4.5 ± 0.4 g (\( n = 151 \), range 3.6–5.7 g). Thirty-one nests fledged at least one offspring, with a mean brood size of 4.0 ± 0.2 (\( n = 31 \) broods). The Domestic Cat (Felis catus) that preyed on the eggs and nestlings was found to be the main predator of the Grey-backed Shrike.

### Variation of egg size with their laying sequence

The fresh egg mass did not vary with the clutch initiation date, but it changed significantly with the laying sequence (Table 1). The later an egg was laid, the larger it was (\( F_{4,146} = 2.52, \ P = 0.04 \); Fig. 1). The variance of eggs’
fresh mass was greater between nests than between years (Table 1).

Growth pattern in nestlings’ body mass
A nestling’s body mass increased significantly with age but did not differ between clutch initiation date or its hatching sequence (Table 2). The variance of nestling’s body mass was greater between nests or years than between nestlings (Table 2).

The last nestlings in a brood were the biggest offspring, whereas the penultimate nestlings were the smallest ones (Fig. 2). As the last nestlings were usually one day younger than their brood-mates, the body mass growth patterns differed significantly among brood-mates (Fig. 2).

Provisioning patterns of the Grey-backed Shrike
The provisioning rate increased significantly with the nestling age in both males and females, but increased with brood size only in females (Table 3). Males’ provisioning rate variance was greater between nests than between years; whole for females, it was greater between years than between nests (Table 3).

Food amount delivered by males per feeding bout increased with the brood size but decreased with the provisioning rate in both males and females. It changed with the nestling age only in females (Table 4). The variance of males’ food amount per feeding bout is greater between nests than between years, while it was the same between nests and between years in females (Table 4).

Males’ provisioning rate (8.92 ± 0.58 bouts/h, n = 16 days) was significantly lower than that of females (11.17 ± 1.09 bouts/h, n = 16 days; t = -2.24, df = 15, P = 0.04; Fig. 3a). However, the amount of food males delivered per feeding bout (0.17 ± 0.003 bouts/h, n = 16 days) was significantly higher than that of females (0.15 ± 0.003 bouts/h, n = 16 days; t = 5.84, df = 15, P = 0.001; Fig. 3b). As a result, males’ contribution to provisioning of offspring (49.06 ± 2.86%, n = 16 days) was the same as that of females (50.94 ± 2.86%, n = 16 days; t = 0.33, df = 15, P = 0.75).

Dietary composition of the Grey-backed Shrike
The dietary investigation identified eleven types of food that parents had delivered to their offspring (Fig. 4). The
frequencies of the different food types are unevenly distributed ($Z = 1.48$, $n = 11$, $P = 0.03$). Lepidoptera larva and Hymenoptera adult were the two main components, contributing 29.07% and 24.27% to the diet, respectively; plant food, mainly powder-branched berry, was the least prevalent component, contributing only 0.15% to the diet (Fig. 4).

The dietary items of the Grey-backed Shrike exhibit two major differences compared with the Brown-cheeked Laughing Thrush. First, nearly one quarter of the Grey-backed Shrike’s diet is adult Hymenoptera that was not listed as food for Brown-cheeked Laughing Thrush. Yet, more than a quarter of the Brown-cheeked Laughing Thrush’s diet is adult Diptera, followed by powder-branched berry (Fig. 4). Second, Grey-backed Shrikes rely mainly on animal food, whereas Brown-cheeked Laughing Thrushes mix animal and plant food to raise their offspring (Fig. 4). Moreover, meat has contributed 5.12% to the Grey-backed Shrike diet (Fig. 4). We recorded one case in which parents tore down a chick that had died in the nest and fed it to the remaining nestlings.

Discussion

Grey-backed Shrike parents adopt a brood survival strategy in both the egg and nestling phases. Such a strategy is not only an adaptive response to the local environmental conditions, but also a consequence of the Grey-backed Shrike life-history style of one brood a year and a wide range of dietary items.

Brood survival strategy of the Grey-backed Shrike in egg-laying

In our study area, female Gray-backed Shrikes’ egg-laying strategy differs from the similar egg size strategy found in altricial birds, often manifested by a tradeoff between the number and size of eggs (Slagsvold et al. 1984; Martin 1987). Under preferable environmental conditions, such as lower nest predation risk and plentiful food supply, parents tend to lay larger clutches of smaller eggs; whereas under poor environmental conditions, such as higher nest predation risk and food scarcity, they are more likely to lay smaller clutches of larger eggs (Forbes 1993; Forbes et al. 2001; Du et al. 2012). Grey-backed Shrikes in our study area (i.e. the lower limit of their distribution on the Tibetan Plateau) produce clutches with a fixed size, unlike the population breeding in the upper limit of their Tibetan Plateau distribution. The population produces unstable clutches of decreasing size with altitude (Lu et al. 2010). Under such conditions, females only need to modulate the egg size based on the laying sequence. It would be a simpler strategy than trading off between the number and size of eggs in coping with the local environmental conditions. As the egg size increases...
with the laying sequence (Table 1), female Grey-backed Shrikes seem to adopt the brood survival strategy by modulating the egg size. This modulation facilitates parents to compensate for the disadvantage faced by the later offspring. After all, the greater the investment parents put into their later eggs, the more likely those offspring are to survive (Du et al. 2014; Da et al. 2018).

Brood survival strategy of the Grey-backed Shrike in brood provisioning

During the nestling period, the last offspring had higher growth rate than their brood-mates, implying that Grey-backed Shrike parents also adopt the brood survival strategy in brood provisioning. In many altricial birds, such as the Horned Larks (Du et al. 2014) and Black-collared Blackbirds (Fan et al. 2017), parents adopt the brood reduction strategy in provisioning their nestlings while they adopt the brood survival strategy when laying their eggs. This difference in parenting strategies between the egg and nestling phases may have resulted from a tradeoff between multiple yearly breeding cycles (Fan et al. 2017; Li et al. 2020a) or between the current and future reproduction (Trivers 1972; Palmer et al. 2004). In both tradeoffs, partial nestlings elimination can at least ensure the nest success; moreover, sacrificing these nestlings might increase the parents’ future reproductive prospects. By contrast, Grey-backed Shrikes in our study area raise only one brood a year. Hence, any starving nestling will reduce parental fitness, whereas sacrificing these nestlings seems unlikely to increase their future reproductive prospects because they produce fixed-sized clutches. Therefore, it is most beneficial for Grey-backed Shrikes to raise the entire brood in the current reproduction. In altricial birds, once hatching asynchrony occurs and size hierarchy is established within the brood, later offspring will be disadvantaged when competing for food with their older brood-mates (Du et al. 2012; Fan et al. 2017). However, we found the last Grey-backed Shrikes offspring to have the highest growth rate within the brood (Fig. 2). Although we obtained no direct behavioral evidence for it, we believe it to be a consequence of parental brood survival strategy. If the last nestlings had not obtained a larger food supply than the other nestlings, they could not keep up with the older nestlings’ growth.

The wide range of Grey-backed Shrike dietary items ensures that parents could support their entire brood. The dietary investigation revealed that parents delivered eleven types of food to their nestlings (Fig. 4), which cover the most common insects found in our study area (Li et al. 2020a) and some plant food types that are also used by other sympatric birds (Fan et al. 2017; Li et al. 2020a). The Grey-backed Shrike’s wide range of dietary items underlies their high provisioning rate and amount of food delivered per feeding bout. As a result, Grey-backed Shrikes can adopt the brood survival strategy in provisioning their offspring.

Life-history style underlying the parenting strategy of altricial birds

Differences in life-history style between sympatric birds, such as the Brown-cheeked Laughing Thrush and Grey-backed Shrike in our study area, could explain the differences in their parenting strategies. The Brown-cheeked Laughing Thrushes have a longer breeding season from early April to later September (Li et al. 2020b). They can, therefore, breed twice a year and adopt different parenting strategies in the two breeding attempts. For example, they deliver food evenly to the nestlings early in the breeding season (brood survival strategy), while bias food towards larger offspring later in the breeding season (brood reduction strategy; Li et al. 2020b). In contrast, Grey-backed Shrikes have a shorter breeding season from the end of June to early September. Hence, they can breed only once a year, so that a brood survival strategy could maximize their reproductive success. Under these conditions, consistent parenting strategies should be maintained between the egg and nestling phases.

Other behavioral responses to the local environmental conditions driven by the one brood a year life-history style also underlie the Grey-backed Shrike brood survival strategy. First, Grey-backed Shrike parents initiate their reproduction in late June, when most berries are ripe (Fan et al. 2017). A large proportion of the sympatric shrub-nesting birds’ diet, including the Brown-cheeked Laughing Thrush and White-collared Blackbird, is composed plant feed (Fan et al. 2017; Li et al. 2020b), indicating that the Grey-backed Shrike dietary composition differ from that of multi-brooded species in the area (Fig. 4). The Grey-backed Shrikes can broaden the range of items in their diet and thus reduce the competition with sympatric species. Moreover, the Brown-cheeked Laughing Thrushes and White-collared Blackbirds exhibit sexual division in provisioning the brood (Fan et al. 2017; Li et al. 2020b). The females often contribute less to provisioning the first brood, whereas the males often contribute less to provisioning the second brood, because they made a greater contribution to provisioning the first one. In contrast, Grey-backed Shrikes do not exhibit such sexual differences because both parents contribute equally to the total food supplied. Sexual division in brood provisioning would make it impossible for the Grey-backed Shrike parents to raise the entire brood by adopting a brood survival strategy.
Conclusions
By investigating the egg-laying and nestling growth pattern, we determined that the Grey-backed Shrike parents adopt the brood survival strategy during their parenting process. The one brood a year life-history style in this species, the delayed initiation of reproduction, and the absence of sexual division in brood provisioning might underlie the brood survival strategy adopted by the Grey-backed Shrike parents.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s40657-021-00244-x.

Additional file 1: Table S1. The mean mass of each food type delivered to the brood by the Grey-backed Shrikes in the Bayi town, Linzhi district. Table S2. Comparison of the percentage of each food type in the dietary items of the Grey-backed Shrike and Brown-cheeked Laughing Thrush that are sympatric in our study area.

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Authors’ contributions
BD and LF designed the study. LG, ZZ, XZ, HZ, WZ and JL collected the life-history data in fieldwork. BD and LG analyzed the data and wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The data used in the present study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
The ethical permits for the nest translocation experiment were issued by the Tibetan Forestry Department (2016ZR-NY-05). The procedures of animal measurement are under the Wildlife Conservation Law of P. R. China (20170101).

Consent for publication
Not applicable.

Competing interests
All authors declare no conflicts of interest to any other organization bodies.

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References
Camfield AF, Pearson SF, Martin K. Life history variation between high and low elevation subspecies of horned larks Eremophila spp. J Avian Biol. 2010;41:273–81.
Caro SM, Griffin AS, Hinde CA, West SA. Unpredictable environments lead to the evolution of parental neglect in birds. Nat Commun. 2016;7:10985.
Christians JK. Avian egg size: variation within species and inflexibility within individuals. Biol Rev. 2002;77:1–26.
Da XW, Xian LL, Luo JJ, Gao LF, Du B. Azure-winged magpies Cyanopica cyanus trade off reproductive success and parental care by establishing a size hierarchy among nestlings. Ibis. 2018;160:769–78.
Decker KL, Conway CJ, Fontaine JJ. Nest predation, food, and female age explain seasonal declines in clutch size. Ecol Ecol. 2012;26:683–99.
Du B, Zhao QS, Liu CJ, Guan MM, Liu NF. Giant babaxen mix brood reduction and brood survival strategies. J Ornithol. 2012;153:611–9.
Du B, Liu CJ, Yang M, Bao SJ, Guan MM, Liu NF. Horned larks on the Tibetan Plateau adjust the breeding strategy according to the seasonal changes in the risk of nest predation and food availability. J Avian Biol. 2014;45:646–74.
Eikenaar C, Berg ML, Komdeur J. Experimental evidence for the influence of food availability on incubation attendance and hatching asynchrony in the Australian red warbler Acrocephalus australis. J Avian Biol. 2003;34:419–27.
Fan LQ, Chen GL, Da XW, Luo JJ, Xian LL, Ren QM, et al. Seasonal variation in food availability influences the breeding strategy of White-collared Blackbirds Turdus albicollis on the Tibetan Plateau. Ibis. 2017;159:873–82.
Forbes LS. Avian brood reduction and parent-offspring “conflict.” Am Nat. 1995;142:84–117.
Forbes S. Sibling symbiosis in nesting birds Auk. 2007;124:1–10.
Forbes S, Glassy B, Thornton S, Earle L. The secondary adjustment of clutch size in red-winged blackbirds (Agelaius phoenicos). Behav Ecol Sociobiol. 2001;50:37–44.
Lack D. The natural regulation of animal numbers. London: Oxford University Press; 1954.
Li JC, Gao LF, Fan LQ, Wong SY, Wei C, Zhang HY, et al. Individual variation in parental tradeoffs between the number and size of offspring at the pre- and post-natal stages. Ibis. 2020a;162:1186–97.
Li JC, Gao LF, Fan LQ, Zhang HY, Zhang W, Du B. Field study of the relationship between personality and reproductive strategy in the White-collared Blackbird Turdus albicollis. Ibis. 2020b;162:245–9.
Lu X, Wang C, Yu T. Nesting ecology of the grey-backed shrike (Lanius tephronotus) in south Tibet. Wilson J Ornithol. 2010;122:399–8.
Mainwaring MC, Lucy D, Harley IR. Hatching asynchrony decreases the magnitude of parental care in domesticated zebra finches: empirical support for the peak load reduction hypothesis. Ethology. 2014;120:577–85.
Martin TE, Lloyd P, Bosque C, Barton DC, Biancucci AL, Cheng YR, et al. Growth rate variation among passerine species in tropical and temperate sites: an antagonistic interaction between parental food provisioning and nest predation risk. Evolution. 2011;65:1607–22.
Martin TE. Food as a limit on breeding birds: a life-history perspective. Ann Rev Ecol Syst. 1987;18:453–87.
Martin TE. Avian life-history evolution in relation to nest sites, nest predation and food. Ecol Monogr. 1995;65:101–27.
Mock DW, Forbes LS. Life-history consequences of avian brood reduction. Auk. 1994;11:115–23.
Palmier AG, Nordmeyer DL, Roby DD. Nestling provisioning rates of peregrine falcons in interior Alaska. J Raptor Res. 2004;38:9–18.
Ren QM, Luo S, Xu JX, Chen GL, Song S, Du B. Helper effects on the azure-winged magpie Cyanopica cyanus in relation to highly-clumped nesting pattern and high frequency of conspecific nest-raiding. J Avian Biol. 2016;47:449–56.
Saether BE, Bakke Ø. Avian life history variation and contribution of demographic traits to the population growth rate. Ecology. 2000;81:642–53.
Shizuka D, Lyon BE. Family dynamics through time: brood reduction followed by parental compensation with aggression and favouritism. Ecol Lett. 2013;16:315–22.
Silver R, Andrews H, Ball GF. Parental care in an ecological perspective: a quantitative analysis of avian subfamilies. Am Zool. 1985;25:823–40.

Skutch AF. Do tropical birds rear as many young as they can nourish? Ibis. 1949;91:430–55.

Slagsvold T, Sandvik J, Røstad G, Lorentsen Ø, Husby M. On the adaptive value of intraclutch egg-size variation in birds. Auk. 1984;101:685–97.

Slagsvold T. Brood division in birds in relation to offspring size: sibling rivalry and parental control. Anim Behav. 1997;54:1357–68.

Tallamy DW, Wood TK. Convergence patterns in subsocial insects. Ann Rev Entomol. 1986;31:369–90.

Trivers RL. Parental investment and sexual selection. In: Campbell B, editor. Sexual selection and the descent of man: 1871–1971. Chicago: Aldine; 1972. p. 136–79.

Verhulst S, Nilsson JA. The timing of birds’ breeding seasons: a review of experiments that manipulated timing of breeding. Philos Trans R Soc Lond B Biol Sci. 2008;363:399–410.

Winkler DW, Wallin K. Offspring size and number: a life history model linking effort per offspring and total effort. Am Nat. 1987;129:708–20.