

Breeding Ecology of the Black Redstart Phoenicurus ochruros at a Tibetan Site, with Special Reference to Cooperative Breeding

Authors: Lu, Xin, Ke, Dianhua, Guo, Yuanyuan, Tang, Shiyi, Zhang, Lixia, et al.

Source: Ardea, 99(2) : 235-240

Published By: Netherlands Ornithologists' Union

URL: https://doi.org/10.5253/078.099.0215

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <u>www.bioone.org/terms-of-use</u>.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Breeding ecology of the Black Redstart *Phoenicurus ochruros* at a Tibetan site, with special reference to cooperative breeding

Xin Lu^{1,*}, Dianhua Ke¹, Yuanyuan Guo¹, Shiyi Tang¹, Lixia Zhang¹ & Chen Wang¹



Lu X., Ke D., Guo Y., Tang S., Zhang L. & Wang C. 2011. Breeding ecology of the Black Redstart *Phoenicurus ochruros* at a Tibetan site, with special reference to cooperative breeding. Ardea 99: 235–240.

This paper provides data on the breeding biology and social system of the Black Redstart Phoenicurus ochruros at a Tibetan site (4300 m elevation). Egglaying occurred between early May and early July during which pairs bred once (10 out of 12 marked pairs) or twice (2). Clutch size varied between 4 and 5 eggs (4.56 \pm 0.51 SE), but a seven-egg clutch was recorded. Incubation by the female lasted 12-14 d (13.0 ± 1.0 SE). Nestlings were provisioned by both parents for 13-18 d (16.7 ± 1.6 SE). Nestlings fledged when they were slightly heavier than adult females. Compared to their European counterparts, the high-elevation redstarts had similar clutch size, incubation or nestling periods and tended to produce fewer clutches of larger eggs. This suggests a life history strategy adapted to maximize offspring survival in the harsh highland environments. Most nests were attended by one male and one female, while at several nests the pair was accompanied by a third individual in female-like plumage. One of these helpers was molecularly sexed as a male and its plumage pattern indicated it to be a yearling. Such cooperative groups occurred in two of the 12 closely monitored nests and two additional nests based on opportunistic observations. One of the four trios was confirmed to raise two broods during a single breeding season and another produced the seven-egg clutch. This is the first confirmation of cooperatively breeding among the 11 Phoenicurus species.

Key words: cooperative breeding, elevational gradient, life history, nesting parameter, *Phoenicurus ochruros*

¹Department of Zoology, College of Life Sciences, Wuhan University, Wuhan 430072, China; *corresponding author (luxinwh@gmail.com)

Elevations can generate environmental gradients along which abiotic (ambient temperature, seasonality and oxygen availability) and biotic conditions (predator and food supply) become harsher and less predictable (Potapov 2004). For avian species that occur across different elevations, life-history patterns are expected to vary in response to local selective pressures. Relative to their lowland counterparts, birds nesting at high elevations have shorter breeding seasons, produce fewer broods, smaller clutches of larger eggs, invest more in parental care and enjoy higher annual survival rates (Badyaev 1997, Badyaev & Ghalambor 2001, Lu 2005, Bears et al. 2009). However, our knowledge about the evolution of avian life history along elevational gradients is limited compared with the knowledge about latitudinal variation (Lack 1968).

The Black Redstart *Phoenicurus ochruros* is an insectivorous passerine breeding in Europe, Asia and Africa. Originally the species was restricted to rocky mountainous areas where they nested in natural hollows. However, they have successfully colonized human settlements, both rural and urban, where they use building crevices as nest sites (Glutz & Bauer 1988, Cramp 1988). This species now has a wide elevational distribution, ranging from near sea level up to 5000 m in the Tibetan plateau (Zheng *et al.* 1983), being one of the largest elevational distributions of any passerine.

Breeding ecology and social behaviour of the Black Redstart is well-described in Europe (Cramp 1988, Weggler 2000, 2001, 2006, Sedlacek *et al.* 2004, 2007, Draganoiu *et al.* 2005, 2006, Dolenec 2006, Sedlacek & Fuchs 2008, Schwarzová *et al.* 2010), but has rarely been studied in Asia (Zhang 1982, Mu *et al.* 2008). Here we present such data collected in a Tibetan settlement at 4300 m elevation, where the birds are summer visitors. We compare the data from this high-elevation site with literature data from lower elevations, aiming to identify possible adaptations of life history and social system of this species.

Methods

Our field work was conducted in Dangxiong (30°28'N, 91°05'E; 4300 m elevation), northern Tibet, during the breeding seasons of 2004–2007 and 2010. The study area is a human settlement of about 100 ha surrounded by alpine meadows. Based on the records of a local weather station over the period 1980–2009, the area has an annual average temperature of 1.7°C, mean summer (June–August) temperature of 10.6°C and mean winter (December–February) temperature of ~7.4°C. Annual precipitation averages 441 mm, 72% of which falls during summer. Further details of the study area are available elsewhere (Ke & Lu 2009).

The Black Redstart is the only Phoenicurus species breeding in the study area. There are approximately 20 redstart territories per year in our study plot of 100 ha. We searched for their nests mainly using behavioural cues. Nests were visited at least once every 5 days to record clutch size, egg size, hatching success, nestling growth and fledging success. For those nests that were estimated to approach hatching or fledging dates, we increased the frequency of nest inspection to once every 1 or 2 days to determine when these events occurred. At 12 nests, at least one of the two parents was captured near their nests using a mist net. Captured individuals were weighed and measured for body dimensions, and marked with 1-3 colour plastic rings and one aluminium ring to allow individual identification. From each of them, a blood sample of $10-15 \ \mu l$ was taken via brachial venipuncture and saved in pure ethanol for molecular sexing. Nestlings were marked when they were older than 10 days. Mate guarding, incubation and nestling provisioning were recorded through focal observation at selected nests. Focal observations lasted 25-170 min and were made 20-30 m away from the nests to avoid disturbance to nesting activities.

Yearling male Black Redstarts are indistinguishable in plumage from females in the field. For one nest with three attendants, we obtained blood samples of the two nest attendants with female plumage. We established their sex molecularly using the primer-pairs P2/P8 and 2550F/2718R (Du & Lu 2010). This method was valid because the assigned sex of all other 21 adults was in agreement with that assessed from plumage and behaviour in the field.

In total, 56 redstart nests were located during this study. However, sampling efforts differed among years and nests. Only the nests found before hatching were used to calculate clutch size. First-egg dates were either observed directly, or backdated based on egg weight or nestling weight. The relationships between these parameters and incubation time and nestling age, respectively, were established at other nests. Only nest-lings that were regularly weighed (once per day) through the complete nestling period were used for estimating growth parameters (logistic model; Ricklefs 1968). Due to small sample size in each year, we pooled the data from the five breeding seasons. Values are presented as means \pm SE.

Results

Redstarts arrived in the study area in early April, and left by early November. More males than females were seen during the initial 10 days since the first arrival record (36 males, 10 females). Among the 26 adults that were metal-ringed in 2006 (11 males and 15 females), one male was recaptured in 2007 (as one of the five individuals captured that year) and another in 2010 (one of the four individuals captured). None of the individuals marked as nestlings (2006, 45; 2007, 8) was rediscovered in the subsequent years.

Most nests (53 of 56) were placed in house crevices, and a few in abandoned burrows of the Tibetan Ground Tit *Parus humilis* or the Sand Martin *Riparia riparia*. Egg-laying occurred between early May and early July, with one peak around late May and another around mid-June (Fig. 1). Clutch start date did not differ between years (ANOVA, $F_{4,50} = 2.09$, P = 0.10). Thus, the double peak did not result from very late breeding during one year. This interpretation was supported by the fact that out of the 12 closely monitored parents, two (each female was marked) produced two broods during a single breeding season.

BREEDING PARAMETERS

Clutch size usually varied between 4 and 5 eggs (4.56 \pm 0.51, n = 27), but one nest contained 7 eggs. Excluding the 7 egg clutch, clutches initiated before 1 June (4.71 \pm 0.47, n = 14) were slightly larger than those initiated after 1 June (4.38 \pm 0.51, n = 13; $t_{25} =$ 1.76, P = 0.09). Eggs were pearl blue. The dimensions of 83 eggs were 20.50 \pm 0.78 mm (range 16.69–22.33) in length and 14.49 \pm 0.42 mm (range 13.54–15.80) in breadth. Brood size at hatching averaged 4.30 \pm 0.80 (range 3–5, n = 20) and at fledging 3.75 \pm 1.08

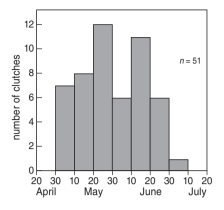


Figure 1. Temporal distribution of the date of laying of the first egg by Black Redstart, southern Tibetan plateau. The time of clutch initiation is arranged in 10-day periods. The data from the five breeding seasons are combined.

(range 2–5, n = 32). Out of the 36 known-fate nests, 32 (89%) fledged at least one nestling.

Only the female built the nest. Incubation was undertaken solely by the female for $12-14 \text{ d} (3.0 \pm 1.0, 100)$ n = 3). Observations at one nest showed that the female spent 48.8% of her daily time (total observation time 678 min) attending the nest, with on-nest bouts lasting 15.8 \pm 7.3 min (range 7–32, n = 21) and offnest bouts 9.0 \pm 4.6 (range 2–22, n = 26). Nestlings were provisioned by both parents for 13-18 d (16.7 \pm 1.6, n = 12 nestlings from four nests, day 0 = hatching date). Provisioning rates (observations of 279 min at four pairs) did not differ statistically between the male (25.4 \pm 18.9 trips/h) and the female parent (18.5 \pm 6.9 trips/h; Wilcoxon test, z = 0.73, P = 0.47). Growth rate constants of the logistic model for the 8 nestlings from 3 broods averaged 0.44 \pm 0.04 g/day (range 0.38-0.50). Shortly before fledging, nestlings weighed 18.2 ± 1.1 g (range 15.3-21.2, n = 62), corresponding to 104% of the adult females (17.5 \pm 0.4 g, range 14.6–21.6, n = 19).

BREEDING SYSTEM

Of the 12 closely monitored breeding units where at least one adult was marked, ten bred as pairs, and two had three attendants. The latter two consisted of one breeding male, one breeding female and one individual with female-like plumage. We captured the breeding female and the female-like attendant from one of the cooperative groups. We found that only the former had a brood patch and was responsible for incubation (twice we observed the female entering the nest cavity and remaining there during the incubation period). The other bird was determined to be a male using molecular sexing. All three individuals in each cooperative group delivered food to the nestlings. In two other nests where we failed to mark any attendants, we observed trios consisting of the pair members and one additional nest attendant with female-like plumage, which all carried food to the nest. We confirmed that one of the two cooperative groups in which one adult male and one adult female were marked produced two broods during a single breeding season. The first nest contained 5 eggs and fledged 5 nestlings on 22 June; the second nest had 5 eggs hatched on 7 July but the nestlings died at day 5. Nests of the three other cooperative groups were located on 8 June (with 5 nestlings), 6 July (7 eggs) and 17 July (3 newly hatched nestlings and 1 egg), respectively. Brood size at fledging was 4.33 ± 0.37 (range 3–5) for three cooperative nests.

Discussion

Birds breeding at high elevations have evolved life histories that favour relatively few offspring of high quality in response to the harsh, unpredictable environments (Bears et al. 2009). The prediction is supported by comparisons of breeding parameters of birds nesting on the Tibetan plateau with their lowland conspecifics or congeners (Lu 2005, 2006, 2008, 2011, Lu et al. 2008, 2009, 2010a, b). Here we compare breeding parameters of two Black Redstart populations on the Tibetan plateau with those reported for their European counterparts at lower elevation (Table 1). We find that the Tibetan redstarts are similar to the European ones in clutch size and length of incubation and nestling periods. However, the higher-elevation redstarts raise fewer broods per season, and lay larger eggs than lower-elevation ones, in particular when taking body size (measured with wing length) into account. The large eggs observed on the northeastern Tibetan plateau may be attributed to potentially higher food availability due to more annual precipitation (782 mm) compared with the drier southern Tibetan plateau (441 mm). The number of annual nesting attempts of birds is mainly limited by the length of the season available for reproduction (Lack 1968). Therefore, given a lower annual reproductive output, higher-elevation redstarts should put more energy into individual offspring to increase their chances of survival. Such a life history strategy is adopted by many organisms breeding in stressful environments (Roff 2002).

An interesting finding of this study was that some redstart nests (16.7%) were cooperatively cared for by both parents and one helper. The helpers were all in female-like plumage, but one individual that was molecularly sexed proved to be a male. Unfortunately, the

	S Tibetan plateau	NE Tibetan plateau	W Europe
Latitude (°N)	30.9	34.1	50
Altitude (m)	4300	3470	<500
Female wing length (mm)	83.4 (78–90)		85.4 (83–90)
Breeding season	Early May – early July	Early May – mid-July	Early April – late July
Duration of breeding season (d)	67	75	110
No. of clutches	1–2	1–2	2–3
Clutch size	4.6 (4–5)	4.8 (3–6)	4–6
No. of eggs/female/season ^a	5–9	5–10	10–15
Fresh egg mass (g)	2.34	2.51	2.16
Egg volume (cm ³) ^b	2195	2659	2052
Incubation period (d)	13 (12–14)	13–14	13–17
Nestling period (d)	16.7 (13–18)	16.9 (15-18)	12–19
Reproductive success ^c	89%	83%	
Reference	This study	Mu et al. 2008	Cramp 1988

 Table 1. Breeding parameters of Black Redstart in different regions.

^aAverage clutch size multiplied by the minimum and maximum numbers of clutches/female/season.

^bCalculated as Egg volume = $0.51 \times \text{length} \times \text{breadth}^2$ (Hoyt 1979).

^cPercentage of nests from which at least one young fledged.

relatedness of the helpers with the helped breeders remained unknown. Three of the four cooperative nests we observed had a normal brood size. The seven eggs present in a single nest cared for by a cooperative group (an uncommon clutch size for the study population) had probably been laid by more than one female through intraspecific brood parasitism.

Cooperative breeding has been found in a few hundred bird species world-wide (Ligon & Burt 2004). However, the behaviour is unevenly distributed across taxa, with species of cooperative lineages usually being poor migrants and inhabiting tropical and subtropical regions (Cockburn 2003). However, a few cooperative species in the Tibetan plateau have recently been reported (Lu 2004, Lu et al. 2007, Du & Lu 2009, Johannessen et al. 2011). None of the 11 members of the genus Phoenicurus has been listed as cooperative breeders (Ligon & Burt 2004), although some of them have been extensively studied. Trios associated with the same nest have occasionally been recorded in two European Black Redstart populations out of at least 15 populations studied (Cramp 1988). Our results indicated that cooperative breeding occurred at a relatively high frequency on the Tibetan population. The presence of helpers may improve reproductive success of the nest by preventing brood reduction (Hatchwell 1999). For the Tibetan Black Redstart population, it appeared that more nestlings on average fledged from group-fed nests (4.3 fledglings per nest) than from pairfed ones (3.6 fledglings per nest). Moreover, parents with helpers may benefit from workload lightening so that they are able to produce more broods during a single breeding season (Brown *et al.* 1978; Russell & Rowley 1988). This seems to be applicable to our study species where the double-brooded case occurred in a cooperative group. We acknowledge that conclusions about social system of the birds based on the current results should be viewed with caution because of limited field samples.

Our study was approved by Law of the People's Republic of China on the Protection of Wildlife (Reference 19881108) and Provisions of the Tibet Autonomous Region for the Protection of Wild Animals and Wild Plants (Reference 20091001). Financial support was provided by the National Sciences Foundation of China (Grants 30425036 and 30670337). We thank XY Ma, LQ Fang, ZW, Peng, JJ Liang, YY Guo, TL Yu, R Hou, C Wang, and B Tao for enthusiastic field work. We also thank Dr. M. Weggler for his valuable comment on the manuscript.

References

- Badyaev A.V. 1997. Avian life history variation along altitudinal gradients: an example with cardueline finches. Oecologia 111: 365–74.
- Badyaev A.V. & Ghalambor C.K. 2001. Evolution of life histories along elevational gradients: trade-off between parental care and fecundity. Ecology 82: 2948–2960.

- Bears H., Martin K. & White G.C. 2009. Breeding in high-elevation habitat results in shift to slower life-history strategy within a single species. J. Anim. Ecol. 78: 365–375.
- Brown J.L., Dow D.D., Brown E.R. & Brown S.D. 1978. Effects of helpers on feeding and nestlings in the Grey-crowned Babbler (*Pomatostomatus temporalis*). Behav. Ecol. Sociobiol. 4: 43–59.
- Cockburn A. 2003. Cooperative breeding in oscine passerines: does sociality inhibit speciation? Proc. R. Soc. Lond. B. 270: 2207–2214.
- Cramp S. 1988. Handbook of the birds of Europe, the Middle East and North Africa, Vol. V. Oxford University Press, Oxford.
- Dolenec Z. 2006. Egg dimensions variation in relation to the laying order in Black Redstart (*Phoenicurus ochruros* Gmelin, 1774) in NW Croatia. Belg. J. Zool. 136: 257–258.
- Draganoiu T., Nagle, L., Musseau R. & Kreutzer M. 2005. Parental care and brood division in a songbird, the Black Redstart. Behaviour 142: 1495–1514.
- Draganoiu T., Nagle, L., Musseau R. & Kreutzer M. 2006. In a songbird, the black redstart, parents use acoustic cues to discriminate between their different fledglings. Anim. Behav. 71: 1039–1046.
- Du B. & Lu X. 2009. Bi-parental vs. cooperative breeding in a passerine: fitness-maximizing strategies of males in response to risk of extra-pair paternity? Mol. Ecol. 18: 3929–3939.
- Du B. & Lu X. 2010. Sex allocation and paternity in a cooperatively breeding passerine: evidence for the male attractiveness hypothesis? Behav. Ecol. Sociobiol. 64: 1631–1639.
- Glutz U.N. & Bauer K.M. 1988. Handbuch der Vögel Mitteleuropas, Vol. 11/1. Aula, Wiesbaden.
- Hatchwell B.J. 1999. Investment strategies of breeders in avian cooperative breeding systems. Am. Nat. 154: 205–219.
- Hoyt O.F. 1979. Practical methods of estimating volume and fresh weight of bird egg. Auk 96: 73–77.
- Johannessen L.E., Ke D.H., Lu X. & Lifjeld J.T. 2011. Geographical variation in patterns of parentage and relatedness in the co-operatively breeding Ground Tit *Parus humilis*. Ibis 153: 373–383.
- Ke D.H. & Lu X. 2009. Burrow use by Tibetan Ground Tits *Pseudopodoces humilis*: coping with life at high altitudes. Ibis 151: 321–331.
- Lack D. 1968. Ecological adaptations for breeding in birds. Methuen, London.
- Ligon J.D. & Burt D.B., 2004. Evolutionary origins. In: Koenig W.D. & Dickinson J.L. (eds) Ecology and evolution of cooperative breeding in birds. Cambridge University Press, Cambridge, pp. 5–34.
- Lu X. 2004. Conservation status and reproductive ecology of Giant Babax *Babax waddelli* (Aves, Timaliinae), endemic to the Tibet plateau. Oryx 38: 418–425.
- Lu X. 2005. Reproductive ecology of Blackbirds (*Turdus merula maximus*) in a high-altitude location, Tibet. J. Ornithol. 146: 72–78.
- Lu X. 2006. Abundance and breeding ecology of Brown Accentors *Prunella fulvescens* in Lhasa, Tibet. Acta Ornithol. 41: 121–128.
- Lu X. 2008. Breeding ecology of an Old World high-altitude warbler, *Phylloscopus affinis*. J. Ornithol. 149: 41–47.

- Lu X., Gong G.H. & Zeng X.H. 2008. Reproductive ecology of Brown-cheeked Laughing Thrushes (*Garrulax henrici*) in Tibet. J. Field Ornithol. 79: 152–158.
- Lu X., Ke, D.H., Zeng X.H. & Yu T.L. 2009. Reproductive ecology of two sympatric Tibetan snowfinch species at the edge of their altitudinal range: response to more stressful environments. J. Arid Environ. 73: 1103–1108.
- Lu X., Ma X.H. & Fan L.Q. 2007. Nesting and cooperative breeding behaviours of a high-altitude babbler, Tibetan Babax *Babax koslowi*. Acta Ornithol. 42: 181–185.
- Lu X., Yu T.L., Liang W. & Yang C.C. 2010a. Comparative breeding ecology of two White-bellied Redstart populations at different altitudes. J. Field Ornithol. 81: 167–175.
- Lu X., Wang C. & Yu T.L. 2010b. Nesting ecology of the Greybacked Shrike (*Lanius tephronotus*) in south Tibet. Wilson J. Ornithol. 122: 395–398.
- Lu X. 2011. Reproductive ecology of three Tibetan waterbird species, with special reference to life-history alterations along elevational gradients. Zool. Stud. 50: 192–202.
- Mu H.Y., Liu N.F. & Yang M. 2008. Breeding of the Black Redstart *Phoenicurus ochruros rufiventris* in the Southeastern Qingzang (Qinghai-Tibetan) plateau. Acta Zool. Sin. 54: 201–208.
- Potapov R.L. 2004. Adaptation of birds to life in high mountains in Eurasia. Acta Zool. Sin. 50: 970–977.
- Ricklefs R.E. 1968. Patterns of growth in birds. Ibis 110: 419-451.
- Roff D.A. 2002. Life history evolution. Sinauer Associates, Inc., Sunderland, Massachusetts.
- Russell E. & Rowley I. 1988. Helper contributions to reproductive success in the Splendid Fairy-wren (*Malurus splendens*). Behav. Ecol. Sociobiol. 22: 131–140.
- Schwarzová L., Štros P., Frynta D. & Fuchs R. 2010. Arrival timing in subadult and adult Black Redstart males: competition-dependent behaviour? Ethol. Ecol. Evol. 22: 111–118.
- Sedlacek O, Fuchs R., Exnerova A. 2004. Redstart *Phoenicurus phoenicurus* and black redstart *P. ochruros* in a mosaic urban environment: neighbours or rivals? J. Avian Biol. 35: 336–343.
- Sedlacek O., Fuchs R., Exnerova A. 2007. Differences in the nestling diets of sympatric Redstarts *Phoenicurus phoenicurus* and Black Redstarts *P. ochruros*: species-specific preferences or responses to food supply? Acta Ornithol. 42: 99–106.
- Sedlacek O. & Fuchs R. 2008. Breeding site fidelity in urban Common Redstarts *Phoenicurus phoenicurus*. Ardea 96: 261–269.
- Weggler M. 2000. Reproductive consequences of autumnal singing in Black Redstarts (*Phoenicurus ochruros*). Auk 117: 65–73.
- Weggler M. 2001. Age-related reproductive success in dichromatic male Black Redstarts *Phoenicurus ochruros*: why are yearlings handicapped? Ibis 143: 264–272.
- Weggler M. 2006. Constraints on, and determinants of, the annual number of breeding attempts in the multi-brooded Black Redstart *Phoenicurus ochruros*. Ibis 148: 273–284.
- Zhang X.A. 1982. Studies on breeding biology of 10 species of passerine birds in alpine meadow, northern Qinghai. Acta Zool. Sin. 28: 190–199.
- Zheng Z.X., Li D.H., Wang Z.X., Wang Z.Y., Jiang Z.H. & Lu T.C. 1983. The avifauna of Tibet. Science Press, Beijing.

Samenvatting

Broeden in het hooggebergte vergt andere aanpassingen dan broeden in het laagland. De Zwarte Roodstaart *Phoenicurus ochruros* broedt van zeeniveau in Nederland tot op 5000 meter hoogte in Tibet. Het is daarmee een ideale soort om aanpassingen aan het leven langs een hoogtegradiënt te bestuderen. In dit onderzoek maten de onderzoekers verschillende broedbiologische parameters in een populatie op de Tibetaanse hoogvlakte. De Tibetaanse Zwarte Roodstaarten legden minder broedsels per seizoen en grotere eieren dan hun Europese soortgenoten. Ook werden er enkele gevallen van coöperatief broeden en een mogelijk geval van broedparasitisme vastgesteld. Dit is het eerste bevestigde geval van coöperatief broeden onder de 11 soorten van het slacht *Phoenicurus*. Broedparasitisme treedt uiterst zeldzaam op in Europese Zwarte Roodstaarten. Het lijkt erop dat Tibetaanse Zwarte Roodstaarten inzetten op minder jongen, maar meer investeren per jong dan hun soortgenoten dichter bij zeeniveau. (KK)

Corresponding editor: Ken Kraaijeveld Received 11 October 2010; accepted 19 April 2011