Seasonal Change in Nest Site and Nest Success of Bull-headed Shrikes

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Seasonal changes in nesting vegetation, nest height, and nest success of the Bull-headed Shrike Lanius bucephalus were studied in Hokkaido, northern Japan from 1992 to 1995. Bull-headed Shrikes nested in dwarf bamboo and vine bushes in early breeding season; however, as the season progressed they moved their nest sites to a variety of deciduous shrubs as the foliage of these nesting plants increased. Nest height gradually increased throughout the breeding season, an observation that was attributed to seasonal changes in the use of nesting vegetation types. Bull-headed Shrikes showed a seasonal change of nest sites with the progress of plant phenology, and the change decreased the probability of predation.

Key words: Bull-headed Shrike, Lanius bucephalus, Nest height, Nest success, Seasonal change of Nest sites

Nest predation is considered to be the major cause of nest failure for many species of birds (NILSSON 1984, MARTIN 1992). It has been suggested that various characteristics of nest sites (e.g., concealment, tree species, height, distance from the forest edge, and so on) influence nest success (e.g., GATES & GYSEL 1978, MARTIN 1987, GRAHAM 1988, KNOPF & SEDGWICK 1992). In temperate zones, the environment for breeding birds changes drastically as nesting trees and herbs foliate. Many authors (e.g., YAMAGISHI 1970, YARNER & SCOTT 1988, GAWLIK & BILDSTEIN 1990, FILLIATER et al. 1994) have stressed that the relationship between plant phenology and predation risk is an important factor for nest site selection by open-cup nesters. However, there is only one study where the effects of leafing were measured quantitatively in relation to the predation risk (SANTOS & TELLERIA 1991). In order to study habitat selection by a bird species in temperate zones, we should pay attention to the temporally changing factors.

In this paper, we describe breeding habitat of Bull-headed Shrikes Lanius bucephalus in Hokkaido, northern Japan. Bull-headed Shrikes are monogamous, single-brooded (although a few are double-brooded), open-cup nesters that replace depredated clutches during breeding season. Females incubate alone, but both sexes feed the young. The diameter of their nests measures about 15 cm. Clutch sizes, which range from 3 to 6 eggs. Eggs are laid between mid-April - early July (TAKAGI unpublished data). Because of their protracted breeding season, Bull-headed Shrikes are suitable subjects for studying seasonal change in nest sites. The objectives of this paper are (1) to clarify alliances between temporal changes in nest height, types of nesting vegetation and plant growth (as foliage increases), (2) to investigate relationships among nest height, types of nesting vegetation, and nest
success.

STUDY SITE AND METHODS

The study was conducted during April-July, 1992-1995 in Oyafuru about 15 km north of Sapporo (43°13'N, 141°20'E). The area consisted mainly of pastures and crop lands with wind shelter-belts composed of ash Fraxinus mandshurica, painted maple Acer mono, and Japanese alder Alnus japonica. Manchurian crab Malus baccata, panicle hydrangea Hydrangea paniculata, and willows Salix spp. were distributed among fallow fields where dwarf bamboo Sasa senanensis and goldenrod Solidago altissima were dominant plants.

To detect temporal changes in nest sites, the clutch initiation date was determined, the types of nesting vegetation was identified, and the height of each nest above the ground was measured. Nest height is defined as the distance from the outside bottom of the nest cup to the ground.

In order to determine the clutch initiation date and nest success, nest contents were checked at one- to six-day intervals until they either fledged or failed. If the exact clutch-initiation date could not be confirmed, it was estimated by backdating from the hatching date, assuming an incubation period of 15 days (as determined in a study by TAKAGI 1994). For those nests where it was not possible to determine hatching date, we calculated clutch initiation dates by aging the nestlings, using the growth mass of known-age nestlings (TAKAGI 1994) and back calculating. We categorized nests either as successful (nests that fledged at least one young, including those where the nestling fledged even though the nest was parasitized by the Common Cuckoo Cuculus canorus) or depredated. If all eggs or nestlings disappeared between nest visits, we considered the nest to be depredated.

We found 150 nests during the four breeding seasons. We excluded 22 nests from analyses of the relationships between each character of nest sites and nest fate, because 8 fell down before nest heights had been measured, and 14 were deserted during incubation. Therefore, 128 nests were available for the analysis. On the other hand, 136 nests were used in the study of the seasonal change in nesting vegetation, as only 14 nests were excluded. Because of the small sample size in each year (i.e., 1992: 18 nests, 1993: 24 nests, 1994: 52 nests, and 1995: 42 nests), data from four years of the study were pooled for the analysis. To detect the seasonal change in types of nesting vegetation, we divided the breeding season into six periods. However, to secure enough data for the statistical analysis of the relationships among nesting fate, types of nesting vegetation, and nest height, the six periods (two each of earlier, middle, and later periods) were collapsed into three (early, middle, and late) periods.

To determine the seasonal change in the foliation of shrubs and vines, we measured the length of shoots for three nesting deciduous shrub species (Manchurian crab, panicle hydrangea, and red-berried elder Sambucus sieboldiana) and vines to the nearest 0.1 mm, once a week from late April to early July 1994. The growth of shoots corresponds to the extent of leaf-out in these plant species. The shrubs and vines were the shrikes’ main nesting substrates (TAKAGI 1994). We define the shoot length as the distance from the base of the shoot to the tip. We randomly selected 15 shrub patches from each shrub species and 10 from vines. Eight shoots were selected and measured from four directions (north, south, east, and west) of
the upper and lower parts of each shrub that divided into two substrates. The timing of foliation did not significantly differ among the four years of our study.

In 1994 and 1995, we estimated the type of nest predators during egg laying, incubation, and nestling phases by the traces of predation.

Statistical analyses were performed using StatView® (Abacus Concepts Inc., Berkeley, CA). The statistical tests were two-tailed with a significance level at .05. When we compared height of nests in each type of vegetation, the standard Bonferoni technique was applied at the adjusted significance levels (as suggested in RICE 1990).

RESULTS

1) Temporal changes in nesting vegetation, nest height, and plant growth

Shrikes selected their nest sites in dwarf bamboos covered by old leaves and dense vine bushes during earlier breeding periods (Fig. 1). They also nested in coniferous shrubs, dwarf plum-yew Cephalotaxus harringtonia and Japanese yew Taxus cuspidata with evergreen leaves. As the season progressed, shrikes nested increasingly in deciduous shrubs (Fig. 1). The main nesting shrubs were Manchurian crab, panicle hydranger, red-berried elder, and willows. In contrast, shrikes’ nesting attempts in dwarf bamboo declined over the breeding season. The predominant nest substrate differed among breeding periods (G-test, G=32.6, df =4, P<0.0001). As for the phenology of nesting vegetation, shoots of the four plant species grew continually from mid-May to mid-June (Fig. 2). The portion of deciduous shrubs used by shrikes for nesting was proportional to the increase in leafing of the particular plant species, while portion of dwarf bamboo used varied

Fig. 1. Seasonal change of types of nesting vegetation. Deciduous shrubs consisted of Manchurian crab Malus baccata, panicle hydranger Hydrangea paniculata, red-berried elder Sambucus sieboldiana, and Willows Salix spp. “Others” are composed by dwarf plum-yew Cephalotaxus harringtonia, Japanese yew Taxus cuspidata, and Japanese larch Larix leptoleps. Data were collected from 1992 to 1995.
Fig. 2. Growth of shoots in three shrub species (red-berried elder, panicle hydranger, and Manchurian crab) and vines. Y-axis, shoot expansion represents that shoot lengths on each measuring day x 100/the length on the last measuring day (%). The survey was conducted in May-July 1994. Number of shoot of Manchurian crab, Panicle hydrangea, Red-berried elder, and vine were 89, 100, 79 and 35, respectively.

Fig. 3. Seasonal change in nest height. Data were collected from 1992 to 1995.

 inversely with the increase of foliage in the deciduous shrubs.

To examine seasonal change in nest height, the relationship between clutch initiation date and nest height was analyzed. Shrikes nested increasingly higher positions as the breeding season progressed (Kendall’s rank correlation coefficient, $\tau = 0.26$, $z = 4.33$, $P < 0.0001$). The seasonal change in nest height corresponded with the expansion of leaves on the various deciduous shrub species (Figs. 2, 3).

The relationship between the clutch initiation date and the nest height in each
vegetation was analyzed separately. The height of nests constructed in vines and dwarf bamboo did not change throughout the season (Vines, $\tau = -0.01, z = -0.08, \text{NS}$; Dwarf bamboo, $\tau = 0.12, z = 1.14, \text{NS}$). However, shrikes nested in higher positions in deciduous shrubs and others, as the season progressed (Deciduous shrub, $\tau = 0.32, z = 2.62, P < 0.008$; Others, $\tau = 0.53, P < 0.05$). Mean nest height $\pm$ SD (n) in dwarf bamboo, vines, deciduous shrub, and others was 77.7 cm $\pm$ 20.4 (59), 107 cm $\pm$ 40.2 (27), 122.8 cm $\pm$ 52.7 (33) and 137.4 cm $\pm$ 79.6 (9), respectively. Nest height in dwarf bamboo were significantly lower than those in three other vegetation types (i.e., dwarf bamboo vs. vines: $U = 388.5, z = -3.8, P < 0.05$; deciduous shrubs: $U = 331.0, z = -5.2, P < 0.05$; others: $U = 133.0, z = -2.4, P < 0.05$). There were no differences in nest height between other types of vegetation. It was suggested that the seasonal increase in nest height was caused by the seasonal change in the composition of the vegetation types used in nesting.

2) The relationships among nesting vegetation, nest height, and nest success

In the analyses of nest success in relation to the types of vegetation, early and late periods were combined, as nest success rates in the two periods were similar. Nest success rates were significantly higher in the late period than in the early/middle periods for both deciduous shrub and others (Fisher's exact probability test, deciduous shrub, $P = 0.04$; others, $P = 0.01$, see Table 1). However, there were no significant differences in nest success between early/middle and late periods in dwarf bamboo and vines (Fisher's exact probability test, dwarf bamboo, $P > 0.99$; Vines, $P = 0.44$, Table 1).

The difference in mean height between successful and depredated nests was not significant in the earlier breeding period. Heights of successful nests in the late period were remarkably higher than those of depredated nests (Table 2), though height of successful nests in the middle period were lower than those of depredated nests.

3) Predators

Potential nest predators in the study area were red foxes *Vulpes vulpes*, weasels *Mustela* spp., rats *Rattus* spp., domestic cats *Felis catus*, snakes *Elaphe quadrivirgata*, *E. climacophora*, Crows *Corvus corone*, *C. macrorhynchos* and

<table>
<thead>
<tr>
<th>Breeding period</th>
<th>Nesting vegetation</th>
<th>Total</th>
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<tbody>
<tr>
<td>Dwarf bamboo</td>
<td>Vines</td>
<td>Deciduous Shrub</td>
</tr>
<tr>
<td>Early</td>
<td>48.4 (15/16)</td>
<td>50 (6/6)</td>
</tr>
<tr>
<td>Middle</td>
<td>52.2 (12/11)</td>
<td>36.4 (4/7)</td>
</tr>
<tr>
<td>Late</td>
<td>44.4 (4/5)</td>
<td>28.6 (2/5)</td>
</tr>
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Table 1. Nest success, number of successful nest x 100/total number of nest (%), in relation to vegetation type in each period. Data collected during April-July, 1992-1995. Numerals in parentheses show: number of successful nests/number of depredated nest.
Table 2. Mean nest height ± SD (n) in relation to nest fate in each period. Data collected during April-Juuy, 1992-1995.

<table>
<thead>
<tr>
<th>Breeding period</th>
<th>Successful nests</th>
<th>Depredated nests</th>
<th>Mann-Whitney test</th>
</tr>
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<tbody>
<tr>
<td>Early</td>
<td>91.9±26.6 (20)</td>
<td>79.1±27.5 (25)</td>
<td>U=186, z=-1.5, NS</td>
</tr>
<tr>
<td>Middle</td>
<td>70.5±31.4 (20)</td>
<td>98.9±19.0 (28)</td>
<td>U=107, z=-3.6, P&lt;0.001</td>
</tr>
<tr>
<td>Late</td>
<td>159.1±29.9 (20)</td>
<td>105.9±68.3 (15)</td>
<td>U=75.5, z=-2.5, P&lt;0.01</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th>Breeding period</th>
<th>Predation by ground mammals</th>
<th>Predation by unknown predators</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fell down/ collapsed</td>
<td>Teeth marks on eggs*</td>
</tr>
<tr>
<td>Early</td>
<td>10 (47.6)</td>
<td>4 (19)</td>
</tr>
<tr>
<td>Middle</td>
<td>4 (21.1)</td>
<td>2 (10.5)</td>
</tr>
<tr>
<td>Late</td>
<td>5 (33.3)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

* The eggs were eaten by rats or weasels.

Northern Goshawks Accipiter gentilis. Apparently, most of the predation in the early period were caused by mammals (Table 3). On the other hand, many of the predations were without any trace of damage in the middle and late periods (Table 3). In the aggregate of the three periods, 25 nests of 55 depredated nests were those certainly attacked by mammalian predators based on the fact that 19 nests had been knocked down and/or collapsed and the other 6 contained eggs with teeth marks of rats or weasels. With the other 30 nests, nest materials were extracted in 3 nests and all eggs were stolen by unknown predators without any trace of nest disturbance in the remaining 27.

DISCUSSION

Loggerhead Shrikes Lanius ludovicianus build their nests at low sites early in the breeding season despite the probably increased risk of possible predation. Their site selection presumably results in protection of their nests from early spring weather (GAWLIK & BILDSTEIN 1990). Although there was no nest that was damaged by strong wind or heavy rain in our study area, we could not find evidence as to whether or weather influenced nest success.

It has been suggested that the concealment of eggs, young, and adults has a major anti-predator function against avian predators in arboreal breeding species.
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(Westmoreland & Best 1985, Martin & Roper 1988, Knopf & Sedgwick 1992, but see Holway 1991, Gtmach et al. 1995). Our Bull-headed Shrikes would also build their nests in sites concealed by leaves or twigs, most likely in order to avoid predation. In earlier breeding season, dwarf bamboo with dense foliage was mainly available for nest sites because deciduous shrub species had not yet leafed-out (Fig. 2). However, since dwarf bamboo was not a higher substrate, it did not provide as safe a site for nesting as did the deciduous shrubs later in the season. Thus, nesting in the early period would result in the higher probability of mammalian predation (Table 3).

It is suggested that the predation probability of artificial avian nests above ground is lower than that on the ground (Yahner et al. 1989). However, there is evidence that in some cases the opposite situation has been found (Santos & Telleria 1991). Furthermore, in a study by Ratti & Reese (1988) no difference was found in predation probability between ground and arboreal nests. In Loggerhead Shrikes, nest height was affected by the vegetation, and the nest success was influenced by only the type of nesting vegetation (Kridelbaugh 1983). These suggest that the difference in predation probability in relation to nest position varying with the type of predators in the area. In the present study 46% of total nests were depredated by ground mammals. Some of the other depredated nests without any trace of damage could be attributable to snakes. The snake species E. quadrievirgata does not disturb the open-cup nests when it depredates eggs or nestlings (M. Ueta, personal communication, July, 25, 1996). Thus, terrestrial predators as well as avian predators may influence the nest failures of shrikes. Shrikes must build their nests at higher positions to avoid terrestrial predators. Since the nests were constructed at higher positions in the middle and late periods, they would be safe from ground mammals (Table 3).

In the middle period, the height of depredated nests was higher than successful nests, a finding that appeared to be inconsistent with our hypothesis. As leaves of higher layer in deciduous tree species were not fully grown during this period, nests built in deciduous shrubs might be easily detected by avian predators. Hence, shrikes might have been confronted by the dilemma that while they needed to build their nests in higher positions to avoid nest predation by ground predators higher substrate was not available for nesting and, during the earlier season at least, this left them vulnerable to attack from avian predators. With shoots expansion in the shrubs, the increasing availability of nest sites would induce shrikes to change types of nesting vegetation.

As a consequence, Bull-headed Shrikes show a seasonal change in the use of nesting vegetation in relation to the plant phenology, and the shifts of nesting sites would allow shrikes to increase their nest success.

We are indebted to Drs. H. Abe, Y. Watanuki, N. Sodhi, and Miss E. C. Hayashi for their valuable comments and improvement of this manuscript, to Mr. M. Ueta for providing his observational data, and especially Dr. C. K. Curtis for improving the English in the manuscript. We specially thank the Hasegawa family for many kindnesses during our field study.
LITERATURE CITED


(Accepted 1 October 1996)
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の南緯1822, 湿度50%ではほぼ一定にして, 光周期のみをLD10:14からLD15:9まで, 1週間に30分づつ明暗を短く, 闇明を短く変化させた。そして飼育期間中に両種とも3～6個体のふんを定期的に採取し, 一定量を蒸留水でホモジェナイズし, このホモジェネートを遠心した後, 上澄みをジェチルエーテルで抽出し, この抽出物中のアンドロジェンをラジオイムノアッセイにより定量した。

その結果, オオジリンにおけるふん中のアンドロジェン含量はLD10:14とLD11:13では低いレベルであったが, LD12:12で急激に増加した。その後, やや減少したがLD14:10までは比較的高いレベルを維持した。一方, オオジロでは光周期の変化に対してふん中のアンドロジェン含量は有意な変化を示さなかった。オオジリンで観察されたように, LD12:12からLD14:10までの高いアンドロジェンレベルを示す光周期は, 渡りの衝動がおきる直前の光波が活発になる時期と一致した。さらに, ふん中のアンドロジェン含量と血液中のアンドロジェン濃度が増加すると考えられる。

ミヤマシトド Zonotrichia leucophrys gambelli を用いた一連の実験から, アンドロジェンは春の渡りの調節要因の一つであることが示唆されている（Rankin, 1991）。したがって, 本研究の結果からオオジリンでも春の渡りにおいてアンドロジェンは重要な役割を果たしているものと推察される。今後, 実際にアンドロジェンが春の渡りの調節要因であるかどうか, さらに検討する必要がある。

モズの営巣場所と巣立ち成功の季節変化
(Seasonal Change in Nest Site and Nest Success of Bull-headed Shrikes. 45: 167-174)

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鳥類の繁殖失敗の最大の原因は卵や雛の捕食であり, 多くの鳥類は捕食による繁殖失敗を回避できるような場所を営巣場所として選択する。営巣場所の様々な特性, たとえば植物による果の被覆の度合い, 樹種, 果の地上からの高さ, 林縁からの距離などの捕食の受け易さ, 受け難さに関係している。しかし, 温帯域で繁殖する鳥類にとっての営巣環境は季節の進行と共に樹木や草本が開花, 伸長するので大きく変化する。したがって, 鳥類の営巣場所選択の研究においては, 営巣環境の季節変化と鳥が選択する巣場所の季節変化をともに解明するが必要がある。多くの研究者は植物の季節性が島型の果実で繁殖する鳥類の捕食に関与して重要であることを強調しているが, 捕食と植物の季節性の関係を定量的に明らかにした論文はほとんどない。本研究では, 1) モズ Lanius bucephalus が営巣した植生の種類および果の地上からの高さの季節変化と営巣植物の開花の季節変化の対応関係を明ら
かにし，2）果実的成功を評価関数にして営農植物，および果の高さの季節変化の意義を検討した。
調査は1982年から1995年の4月から7月まで，北海道石狩郡の農耕牧草地で行った。モズが営農
する植生の利用頻度の季節変化を分析した結果，繁殖期前後には前年の果で覆われたクマイザサ
Sasa senanensiやツツジ植物のツルで覆われた場所に営農し，穂性の常緑針葉樹 ハイヌガヤ
Cephalotaxus harringtoniaとも営農し，落葉広葉樹の低木の利用
頻度は低かった（Fig. 1）。繁殖期が進行するにつれてササおよびツツジ植物の利用頻度はともに低下
したのに対し，落葉広葉樹の低木であるエゾノコリンポ Malus baccata，ノリウツギ Hydrangea
paniculata，エゾツツジ Sambucus sieboldiana，ヤナギ類 Salix spp. の利用頻度は上昇し
た（Fig. 1）。繁殖期後期ではササの利用頻度がさらに低下し，落葉低木の利用頻度は上昇した
（Fig. 1）。これらの落葉低木は繁殖期中には一気に伸長し，後期にかけて開花しきる状態になって
おり，モズが営農する植生の季節変化は，植物の開花の進行に対応していた（Figs. 1, 2）。
果の地上からの高さ，季節の進行とともに高くなった（Fig. 3）。それぞれの植物ごとに果の高さ
の季節変化を示したところ，ササツツジ植物は季節を通じて果の高さに変化は認められなかったが，
落葉広葉樹低木とその他の場所における果の地上高は季節の進行とともに高くなった。ササに造られ
た果の地上高は他の果に造られたものよりも有意に低かった。果の地上高の季節変化も植物の開花の
進行に対応していた（Figs. 2, 3）。繁殖期初期の果が低い位置に造られたのは，より高い位置に果
を被覆する葉が開花していなかったためと考えられる。
営農植物のタイプごとに果実成功の季節変動を比較した。その結果，ササおよびツツジ植物では季
節的な差は認められなかったが，落葉広葉樹低木とその他の果では後期が前/中期に比較して果実成功
が有意に高かった（Table 1）。捕食を受けなかった果と捕食を受けた果の地上からの高さを比較した
結果，繁殖期前後では差が認められなかったが，中期では捕食を受けた果で高くなり，後期では捕
食を受けなかった果の地上高が高かった（Table 2）。地上性の果実によって捕食率が前期に高かっ
たのは，果の地上高が低かったためと考えられ，中，後期に低くなっしたのは果の地上高が高くなった
めであろう（Table 3）。
他に鳥の研究では，地上の果よりも樹上の果の方が捕食圧が低いことが実験的に明らかにされてい
る。一方，逆の結果も示されており，捕食圧は調査域の捕食者の種類によって異なると考えられる.
そこで，1995，96年に捕食を受けた55果の状況から捕食者を推定した。その結果，46%（n=25）が
地上性は乳類によって捕食されていた，また，果の構造に全く掠乱の痕跡がない捕食者が49%（27），
果材が引き出された捕食者が5%（3）認められた。これらの捕食者は特定できなかったが，掠乱の痕
跡がない捕食にはヘビ類よ捕食者が含まれると考えられたので，本調査域では地上捕食者の捕食圧が
高いため，モズが地上捕食者による捕食を逃れるためには営農場所を高くする必要があい一方で，
植物の開花の季節性に果場所の利用可能性が制約されていると考えられた。
これらのことから，モズは植物の季節変化にしたがって，捕食を回避できるように営農場所を変化
させていることが示唆された。