SEXUAL DIMORPHISM IN THE DEVELOPMENT OF SCENT STRUCTURES FOR THE OBLIGATE MONOGAMOUS EUASIAN BEAVER (CASTOR FIBER)

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Scent marking is a common form of territorial demarcation in mammals and differential marking by the sexes may select for differences in scent gland developmental rates or size. Rates of marking reflect level of defense and so behavioral dimorphism in marking by the sexes may select for dimorphism in scent structures. Eurasian beavers (Castor fiber) are monogamous, territorial mammals that show some behavioral dimorphism in territoriality. Beavers have 2 primary scent structures, anal glands and castor sacs. Anal gland secretion appears to serve as a family or individual identifier. Castor fluid is the main source of chemical signals used for marking territories. We examined beavers collected from 3 different habitats, namely streams, creeks and tarns (small lakes), which varied in the density of beaver sites. The objectives of this study were to examine if morphological attributes of Eurasian beaver scent structures reflected differences in adult beaver territorial function or in the ontogeny of the sexes. We hypothesized that male beavers would have larger castor sacs and anal glands, and that beavers from sites of greater density, and thus greater rates of intrusion, would have larger castor sacs and anal glands. Our results showed that investment did not differ by habitat type. Although the combined weight of the scent structures did not differ between the sexes, males developed significantly larger anal glands but somewhat smaller castor sacs than females. Total investment to scent structures may be limited but allocation is flexible. Males patrol the territory more extensively and may benefit by releasing anal gland secretion to signal their identity to potential intruders. Higher rates of marking may decrease castor sac size by reducing the amount of stored scent material. Hence, even in monogamous mammals, sexual dimorphism can arise from behavioral dimorphism.

Key words: Castoridae, chemical ecology, ontogeny, scent marking, territory

Monogamous mammals show reduced or no apparent sexual dimorphism relative to polygamous species (Kleiman 1977; Ralls 1977). Both sexes of monogamous pairs provide parental investment (Trivers 1972), and contribute to territorial defense (Clutton-Brock 1989). However, males and females may demonstrate different activity patterns and play different roles. Minimally, female mammals nurture the young during lactation, whereas males purportedly perform other tasks more frequently, such as territory maintenance and defense. Scent marking is a common form of territory demarcation and announcement of status in mammals (Ralls 1971). Sexual selection may shape such olfactory communication (Blaustein 1981; Darwin 1871), resulting in differential investment in scent structures and their products between the sexes (Heymann 1998; Jannett 1986).

Genetic variation in the sexes could account for differences in developmental rates and size of scent structures. Environmental components also could be important. In general, habitat quality influences developmental rates, adult size, and susceptibility to disease of animals (Dixon and Kundu 1998; Judson and McFarlane 1998) in the classic sense of allocation and life history trade-offs (Cody 1966; Stearns 1976). Hence, it is plausible to hypothesize that variation in nutrition among the sexes could influence investment in secondary characteristics such as scent structures. In addition, behavioral factors could influence marking rates and resultant investment in scent structures and chemical signal production (Collins et al. 2001). For example, if one sex ranged further than the other, the former may encounter more intruders or unmarked regions and respond by marking at higher rates. Selection could then favor individuals of this sex with relatively larger or more active scent structures.

Eurasian (Castor fiber) and North American (C. canadensis) beavers are obligate monogamous mammals and are relatively long lived; they pair bond for multiple years to life and exhibit little external morphological sexual dimorphism (Novak 1987;
Beavers use extensive scent marking to demarcate and maintain territories (Aleksiuk 1968; Rosell 2002a; Schulte 1993; Wilsson 1971). The number of scent marks is density dependent. Beavers with many close neighbors (highly challenged) scent mark more often to be unambiguously recognized as territory owners. Beavers living in higher densities may reside in higher-quality habitat and incur greater rates of intrusions, stimulating increased marking (Rosell and Nolet 1997). Some behavioral differences in adult males and females have been observed, especially during spring and summer when young are reared (Buech 1995; Hodgdon and Larson 1973; Schulte and Müllar-Schwarze 1999; Wheatley 1997; Wilsson 1972). Two recent studies on mated male and female Eurasian beavers suggest that males have significantly larger kernal home ranges (Herr and Rosell 2004) and that males allocate significantly more time to travel (Sharpe and Rosell 2003). Male Eurasian beavers also perform more scent marking and hence territorial maintenance than do female beavers (Thomsen 2002). As yearlings, female North American beavers act more like adults, whereas males behave more like kids (Busher and Jenkins 1985), although this may not be the case for scent-marking behavior (Hodgdon 1978). Maturational rate may affect morphological characteristics related to behavioral activities such as marking.

Beavers have 2 primary scent structures, anal glands and castor sacs (Rosell et al. 1998; Svendsen 1978). These are located in 2 cavities between the pelvis and base of the tail (Valeur 1988; Walro and Svendsen 1982). The anal gland is a holocrine secretory gland, but the castor sac is only a pocket beyond the normal bunting season. The Norwegian Directorate for Natural Management (59°25'S, 9°3'E), Telemark County, southeastern Norway, during the normal hunting and trapping season of 1 October–30 April 1996–1997, 1997–1998, and 1999–2000 (n = 119). Additional beavers were collected during a 15-day extension from 1 to 15 May during 1997–1999 (n = 32). The Norwegian Directorate for Nature Management and the respective landowners granted permission to hunt beavers beyond the normal hunting season. The Norwegian Directorate for Nature Management approved the hunting process, and we adhered to guidelines of the American Society of Mammalogists (Animal Care and Use Committee 1998).

We delineated 3 seasons as autumn (October), winter (December to March), and spring (April and May). A majority of the beavers (86%) from which a full data set was acquired were captured in the spring. Shortly after death, carcasses were brought to the laboratory where castor sacs and anal glands were dissected and excess tissue was removed. Both left and right sacs and glands were weighed (to the nearest 0.1 g) and measured by a slide caliper (length and width at the middle, to nearest 1.0 mm). Precautions were taken to minimize fluid loss from the sacs and glands. Age, sex, body weight (nearest 0.1 kg), habitat, and date of collection also were recorded for each beaver. Beavers were sexed by color of the anal gland secretion (Rosell and Sun 1999) and by absence or presence of the os penis (Osborn 1955).

Beavers are generally classified into 3 age classes (Hartman 1992; Rosell and Pedersen 1999): 1 year old (<12 months, 0–10 kg), 2 years old (12–24 months, 10–15 kg), and adult (≥24 months, ≥15 kg). Because weights only provide approximate age classes, we used a more specific method of aging to separate developmental periods. For most of the beaver specimens (n = 124), we examined tooth root closure and annual cementum and dentine layers of the 2nd molar (van Nostrand and Stephenson 1964). We used these 124 beavers for the statistical analyses involving different age classes. At age 3 years, beavers are 1st entering adulthood, so we considered this age as part of
the early developmental process. For comparisons to the traditional adult age class delineated by using weights only, we also considered 3-year-old animals as adults but only for separate analyses. Final sample sizes for each age group were as follows: 1 year old (11 females, 7 males), 2 years old (9 females, 18 males), 3 years old (9 females, 8 males), and >3 years old (28 females, 34 males). For analyses of adult age classes (>2 years old), we included 9 additional beavers (5 females, 4 males) not aged by their teeth but clearly adults.

Beavers were collected from 3 different habitats (tarns, rivers (>10 m wide), and creeks (<10 m wide)) that differed in the density of beaver sites. Territorial boundaries were determined based on location of scent mound concentrations (Rosell and Nolet 1997; Rosell et al. 1998), and/or regular sight observations of animals moving up- and downstream of the lodge (Rosell et al. 1998). Number of neighbors (referred to as site density) was calculated as number of territories located together, that is, bordering on each other with no unoccupied stretch of stream area separating them.

Data compilation and analysis.—Weights of left and right anal glands or castor sacs were summed to give a total scent structure weight for each type gland. Length and width were multiplied to give area and then added (left and right anal gland or castor sac) to give a total area. The correlation between weight and area was determined. Because weight was a more reliable and meaningful measure than area, we limited most analyses to weight. All means are presented as mean ± SE. Analysis of variance was performed by using Statistica software (StatSoft, Inc. 1999). Logarithm of weight was used in analyses to normalize the data. Log of body weight was used as a covariate so that differences in scent structure size could not be accounted for simply by differences in body weight. A test of parallelism was used to examine if the slopes of linear regression lines differed.

RESULTS

Seasonal differences.—No significant differences were found in scent structure weights by season (P > 0.15 for anal gland and castor sac). Because of insignificant results and small sample sizes in autumn (n = 5) and winter (n = 12) compared to spring (n = 107), the data sets were combined across season.

Habitat differences.—The number of beaver sites with neighbors differed among the 3 habitats (tarns, rivers, and creeks; F = 13.97, df = 2, 46, P < 0.001), with an average of 1.2 ± 0.16 neighbors per site in the river habitat, 0.4 ± 0.13 neighbors in the tarn, and 0.2 ± 0.11 neighbors in the creek habitat. However, size of scent-producing structures did not differ significantly by habitat for adult beavers of either sex (n = 79, age 3 years and greater). This was true for area of the anal gland (F = 1.72, df = 2, 72, P = 0.19), weight of the anal gland (F = 1.34, df = 2, 72, P = 0.27), area of the castor sac (F = 1.02, df = 2, 74, P = 0.36), and weight of the castor sac (F = 1.88, df = 2, 74, P = 0.16). We made planned comparisons by combining tarns and creeks, which iced over in winter, whereas rivers in our study did not. However, no significant differences were evident (all P > 0.08). The weight and area for each scent structure were highly and significantly correlated (anal gland: r² = 0.54, n = 78, P < 0.001; castor sac: r² = 0.48, n = 81, P < 0.001). To check that beavers from the same site were not overrepresented (and thus to avoid pseudoreplication), we repeated analyses with only 1 adult beaver from each site (5–12 of each sex per habitat type). For the 3 site types (river, creeks, and tarns), no difference in weight of the anal gland (F = 0.20, df = 2, 41, P = 0.82) or castor sac (F = 1.57, df = 2, 41, P = 0.22) by site type was apparent. Age of adult beavers at the 3 sites (n = 11–17) also did not differ (F = 1.54, df = 2, 37, P = 0.23); age averaged 6.2 ± 0.79 years at river sites, 5.0 ± 0.54 years at tarns, and 7.3 ± 1.17 years at creeks.

Ontogeny of the scent structures.—Beavers showed a typical growth curve, with no statistical difference in mean weight between males and females across all ages (F = 2.14, df = 1, 121, P = 0.15), although rate of growth differed (test of parallelism: F = 15.3, df = 1, 120, P < 0.001; Fig. 1a). The weight of the scent-producing structures (anal gland plus castor sac) showed a similar pattern of no significant difference between sexes, but a difference in weight change with age (F = 0.46, df = 1, 121, P = 0.5; test of parallelism: F = 5.1, df = 1, 120, P = 0.025; Fig. 1b).

Male and female beavers showed differential investment in the 2 scent-producing structures. Males exhibited a significantly greater investment than females in anal gland weight relative to body weight (F = 57.8, df = 1, 117, P < 0.0001; test of parallelism: F = 30.2, df = 1, 116, P < 0.0001; Fig. 2a). Castor sacs of females were generally larger than those of males but differences were not significant (Fig. 2b). Early developmental differences were apparent in the growth of both anal glands and castor sacs. Considering only beavers younger than 4 years, a significant difference existed between males and females for weight of the anal gland (F = 10.2, df = 1, 59, P = 0.002) and the castor sacs (F = 7.06, df = 1, 59, P = 0.01; Fig. 3). These young females had average anal gland weights of 25.5 ± 2.24 g compared to 40.4 ± 3.72 g for young males. These same females had average castor sac weights of 95 ± 14.2 g compared to 78.5 ± 8.02 g for males. The slopes differed for the linear fits for anal glands (test of parallelism: F = 17.5, df = 1, 58, P < 0.0001) but not for castor sacs (test of parallelism: F = 0.24, df = 1, 58, P = 0.63). Hence, for anal glands, both the amount and rate of investment differed between the sexes during the 1st few years of life, whereas for castor sacs, the amount of investment but not the rate of investment differed between males and females.

DISCUSSION

The study revealed sexual dimorphism in the development of scent structures in Eurasian beavers. Males and females invested differently in the 2 types of scent structures (anal glands and castor sacs). This investment was most apparent in the first 3 years of life. Beavers from different habitats that also differed in the density of beaver sites had similarly sized scent structures.

Castor sac material serves as a territorial scent mark (Rosell and Sundsdl 2001; Rosell et al. 1998; Schulte 1998). Males spend more time devoted to marking and territory investigation than do females (Herr and Rosell 2004; Thomsen 2002). A study of North American beavers indicated that males and females had similarly sized castor sacs, but the samples came mainly from adult beavers (Bollinger et al. 1983). The current study found that young males had relatively smaller castor sacs than young females, but sizes were not significantly different in adults.
Because males clearly did not have larger castor sacs than females, use and investment in size are not directly related. This contradicts our 1st prediction on a direct relationship between marking rates and castor sac size. Our 2nd prediction stated that castor sacs of males could be smaller because of increased use. Higher flow rates of urine through the sacs could deplete scent material, resulting in lighter castor sacs. This does not explain why young males would have lighter castor sacs than young females, unless young, male Eurasian beavers mark more than do young females. This would be different from 1 population of North American beavers, where young of each sex marked previously created mounds at similarly frequencies (Schulte 1998), possibly to create a family scent. In the current study, smaller observed sizes of castor sacs in young males might occur if castor sac size was influenced by investment in the anal gland.

Previous studies have suggested that anal gland secretion acts as an individual identifier by being released into the water (Gronneberg and Lie 1984; Rosell 2002b; Sun and Müller-Schwarze 1998). Individuals who are most likely to meet other beavers may use anal gland secretion more readily. This would permit intruders an opportunity to match the scent mark to its owner (the scent-matching hypothesis—Gosling 1982).
Because males appear to spend more time patrolling the territory, they are most likely to interact with nonresident beavers. In a study by Bollinger et al. (1983), male North American beavers had slightly larger anal glands than females. In the current study, males had an increased level and rate of investment in anal gland weight compared to females throughout development and into adulthood. Thus, males had smaller castor sacs but larger anal glands than females, but overall investment in scent structures did not differ between the sexes. Size of the scent structures increased with age in an allometric fashion, showing the same relative growth as overall body weight.

Males may require more equitable investment in both the individual identifier scent structure (anal gland) and the territory demarcation scent structure (castor sac) than females. If higher flushing rates reduce castor sac size, then this size reduction may create more space for larger anal glands. Combined, these ideas suggest that castoreum and anal gland secretion are both important signalers from a territory holder. This hypothesis makes testable predictions for future studies. First, males should respond to experimental scent marks on the periphery of the territory sooner or more often than females. Second, males should make or update outlying scent marks (those on extremes
of territory) more than females. Third, males should encounter nonresident beavers more frequently. The 3rd prediction is partially supported by the observation that in a population of North American beavers, adult males had more tail scars and notches than adult females (Müller-Schwarze and Schulte 1999). Such injuries indicate a greater frequency of competitive interactions between beavers from different sites.

Beavers exhibit a monogamous mating system with high levels of parental investment by both sexes. Morphological sexual dimorphism is generally absent in such species (Kleinman 1977) and only slight behavioral dimorphism is apparent, as is typical in such obligate monogamous species (Clutton-Brock 1989; Svendsen 1989; Wilsson 1971). As with many mammals, beavers rely heavily on scent to communicate within the species, especially to communicate territoriality (Rails 1971; Rosell 2002a; Schulte 1998). We suggest that a difference in territorial behavior between male and female beavers has led to differential investment in scent structures. Total allocation to scent structures is apparently constrained, because each sex exhibits a similar combined weight relative to total body weight. However, this similar total investment arises from very different functional ends. We propose that females have larger castor sacs (low flushing rates) and smaller anal glands (infrequent use) because of reduced employment of both glands for territorial purposes, whereas males have just the opposite configuration because of enhanced use of both glands in territorial demarcation. Hence, behavioral dimorphism selects for morphological dimorphism in signal-producing structures, even in an obligate monogamous species, but the extent of the divergence is limited by total developmental allocation to the structures.

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**LITERATURE CITED**


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