

Sexual Selection and Canine Dimorphism in New World Monkeys

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ABSTRACT Social and ecological factors are important in shaping sexual dimorphism in Anthroidea, but there is also a tendency for body-size dimorphism and canine dimorphism to increase with increased body size (Rensch's rule) (Rensch: *Evolution Above the Species Level*. London: Methuen, 1959.) Most ecologists interpret Rensch's rule to be a consequence of social and ecological selective factors that covary with body size, but recent claims have been advanced that dimorphism is principally a consequence of selection for increased body size alone. Here we assess the effects of body size, body-size dimorphism, and social structure on canine dimorphism among platyrrhine monkeys.

Platyrrhine species examined are classified into four behavioral groups reflecting the intensity of intermale competition for access to females or to limiting resources. As canine dimorphism increases, so does the level of intermale competition. Those species with monogamous and polyandrous social structures have the lowest canine dimorphism, while those with dominance rank hierarchies of males have the most canine dimorphism. Species with fission-fusion social structures and transitory intermale breeding-season competition fall between these extremes.

Among platyrrhines there is a significant positive correlation between body size and canine dimorphism. However, within levels of competition, no significant correlation was found between the two. Also, with increased body size, body-size dimorphism tends to increase, and this correlation holds in some cases within competition levels.

In an analysis of covariance, once the level of intermale competition is controlled for, neither molar size nor molar-size dimorphism accounts for a significant part of the variance in canine dimorphism. A similar analysis using body weight as a measure of size and dimorphism yields a less clear-cut picture: body weight contributes significantly to the model when the effects of the other factors are controlled. Finally, in a model using head and body length as a measure of size and dimorphism, all factors and the interactions between them are significant. We conclude that intermale competition among platyrrhine species is the most important factor explaining variations in canine dimorphism. The significant effects of size and size dimorphism in some models may be evidence that natural (as opposed to sexual) selection also plays a role in the evolution of increased canine dimorphism.

It is generally recognized that there are many "causes" for sexual dimorphism in primates and other mammals. Four sorts of mechanisms have been proposed to explain dimorphism: phyletic inertia, correlated response, natural selection, and sexual selection.

Explanations that invoke phyletic inertia (Cheverud et al., 1985a,b) depict di-

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morphism in a species as an adaptation of its ancestors that is not explicable as part of that species' current adaptations. Sexual dimorphism has also been viewed as a nonselected "correlated response" to overall selection favoring either 1) an increase in body size or 2) an increased body-size dimorphism (Leutenegger and Cheverud, 1982, 1985): in this view, selection favoring increased size produces a correlated increase in dimorphism when there is unbalanced phenotypic variability between the sexes or a difference between male and female heritabilities for the dimorphic trait; by the same token body-size dimorphism might "drag" canine dimorphism along with it. The third class of explanations for dimorphism involves natural (as distinct from sexual) selection, e.g., differential resource utilization by the sexes, and different roles of the sexes for defense against predation (Richard, 1974; Clutton-Brock et al., 1977; Leutenegger and Kelly, 1977; Harvey et al., 1978; Clutton-Brock and Harvey, 1978; Smith, 1980; Anderson, 1986).

Finally, ever since Darwin, sexual selection has been invoked to explain sexual dimorphism. In this model, dimorphism is maintained or enhanced because females prefer to mate with males having certain morphological attributes or because dimorphic traits confer an advantage in intermale competition (Darwin, 1871; Trivers, 1972; Brown, 1975; Harcourt et al., 1976).

There are also many different ways that sexual dimorphism is expressed. There may be differences in body size, color patterns, body shape, or canine size, to name a few of these. The degree of dimorphism in these various anatomical features is probably under different selective pressure and this probably explains, for example, why dimorphism in body weight and canine size do not precisely covary in primates. In this paper we use a comparative approach to consider canine sexual dimorphism in relation to the above possible causes. We particularly are interested in the degree to which variation in intermale competition is sufficient to explain dimorphism in platyrrhine canines and the degree to which canine dimorphism and body-size dimorphism respond similarly with changes in social structure and/or size.

MATERIALS AND METHODS

Social structure

To avoid the possibility of bias in the outcome of this study, Glander and Wright,

working independently, and without prior knowledge of the morphological findings of Kay and Plavcan, gathered data from the literature and from personal observations to produce the classification of social structure presented here. In this analysis, four levels of social structure are recognized in platyrrhine primates. At each ascending level there is an increase in the degree of intermale competition. Table 1 summarizes the taxa used in this study according to social structure and the level of competition. In level 1, once breeding units are established, there is little or no competition among males for breeding access to mates. There are two such social structures among platyrrhines. In polyandry, seen in most or all Callitrichinae, two or more males copulate with a single female during one breeding season and cooperate to raise the offspring (e.g., Terborgh and Goldizen, 1985). In monogamy, which occurs in many other platyrrhines, including *Aotus* and *Callicebus*, one adult pair breeds exclusively over several mating seasons (e.g., Wright, 1986).

In levels 2–4, significant breeding competition occurs. Platyrrhines at these levels are characterized by multimale/multifemale troops. In level 2 there is limited intermale competition for breeding access to females. Such a breeding pattern occurs in fission-fusion social groups. For example, in *Ateles* promiscuous mating may occur during the breeding season within nonpermanent subgroups drawn from a larger, more stable community (Klein and Klein, 1975; Symington, personal communication). Alternatively, estrous females may form consort pairs away from the group. Level 2 competition also is found in *Brachyteles*, which lives in multimale/multifemale groups. Male *Brachyteles* show little aggression in competing for sexual access to receptive females (Aguire, 1971; Milton, 1985; Strier, 1986). In the level 3 breeding pattern, exemplified by *Cebus capucinus* and *Saimiri*, competition among males occurs primarily during the mating season (Robinson and Janzen, 1987). At other times of the year, there is less intermale competition. Finally, intermale competition is most extreme in level 4 species. Often, as in *Alouatta*, an established hierarchy among males is maintained throughout the year (Rudran, 1979; Glander, 1980; Clarke, 1983; Crockett and Eisenberg, 1987). This hierarchy may determine not only which males have access to mates during the breeding season but also may confer an advantage on

TABLE 1. Summary of levels of intermale competition for the species used in this study, as defined in the text, and the behavioral data on which the classification was based

Taxon	Social structure	Competition level	References
<i>Callithrix jacchus jacchus</i>	Monogamy/polyandry ^a	1	Alonso (1984)
<i>jacchus penicillata</i>	Monogamy/polyandry ^a	1	Hershkovitz (1977)
<i>Cebuella pygmaea pygmaea</i>	Monogamy/polyandry ^a	1	Hernandez-Camacho & Cooper (1986)
<i>Saguinus fuscicollis nigrifrons</i>	Polyandry ^b	1	Goldizen & Terborgh (1986)
<i>midas niger</i>	? Polyandry	?1	None
<i>oedipus geoffroyi</i>	Polyandry ^b	1	Dawson (1977); Neyman (1977)
<i>Leontopithecus rosalia rosalia</i>	Mongamy/polyandry ^c	1	Deitz (1985)
<i>Alouatta belzebul</i>	Unknown		None
<i>caraya</i>	Dominance rank competition ^d	4	Thorington et al. (1984)
<i>fusca</i>	Dominance rank competition	4	Da Silva (1981)
<i>palliata</i>	Dominance rank competition	4	Milton (1978)
<i>pigra</i>	Dominance rank competition	4	Coelho et al. (1976); Watts et al. (1986)
<i>seniculus insulanus</i>	Dominance rank competition	4	Neville (1972)
<i>seniculus seniculus</i>	Dominance rank competition	4	Klein & Klein (1975)
<i>Ateles fusciceps fusciceps</i>	Fission/fusion ^e	2	Klein & Klein (1975)
<i>fusciceps robustus</i>	? Fission/fusion	?2	None
<i>geoffroyi vellerosus</i>	Fission/fusion	2	Fedigan & Baxter (1984)
<i>paniscus chamek</i>	Fission/fusion	2	White (1986)
<i>Brachyteles arachnoides</i>	Multimale/multifemale low competition or Fis/Fus	2	Aguire (1971); Milton (1985); Strier (1986)
<i>Lagothrix flavicauda</i>	Multimale/multifemale low competition	2	Leo Luna (1980)
<i>lagotricha lagotricha</i>	Unknown	?	None
<i>Aotus trivirgatus griseimembra</i>	Monogamy	1	Wright (1978)
<i>trivirgatus lemurinus</i>	?Monogamy	?1	None
<i>trivirgatus trivirgatus</i>	?Monogamy	?1	None
<i>Callicebus moloch discolor</i>	Monogamy	1	Kinzey et al. (1977)
<i>torquatus lugens</i>	Monogamy	1	Defler (1983)
<i>Cebus apella libinosus</i>	Dominance rank competition	4	Janson (1984)
<i>capucinus capucinus</i>	Mating season competition	3	Freese & Oppenheimer (1981)
<i>Saimiri sciureus boliviensis</i>	Mating season competition ^f	3	Hershkovitz (1977)
<i>sciureus macrodon</i>	Mating season competition ^g	3	Klein & Klein (1975)
<i>oerstedii oerstedii</i>	Mating season competition ^h	3	Baldwin & Baldwin (1981); Boinski (1987)
<i>Cacajao calvus</i>	Mating season competition	3	Ayres (1986)
<i>Chiropotes satanas chiropotes</i>	Monogamy ⁱ	1	Ayres (1981)
<i>Pithecia pithecia</i>	Multimale/multifemale low competition	2	Buchanon et al. (1981); Oliveira et al. (1985)

¹Notes: a, lack of agreement as to the type of social organization; b, reference population is *S. f. weddelli* from Peru; c, reference population is from Bolivia; d, reference population is from Argentina; e, reference population is *A. belzebul* from Colombia; f, at Manu study site in Peru, extreme breeding season competition occurs involving the death of some males (C. Janzen and C. Mitchell, personal communication); g, reference population is from Columbia; h, breeding season described as being at a comparatively low level; i, this species found in multimale/multifemale groups, but reported to be monogamous within these groups.

TABLE 2. Sample sizes of males and females, canine dimorphism, Canine Dimorphism Index (CDI), \ln female M^1 area, and M^1 area dimorphism for each of the species studied¹

Taxon	M/F	Canine dimorphism				CDI	\ln female M^1 area	M^1 area dimorphism
		C_1 max	C_1 min	C^1 max	C^1 min			
<i>Callithrix jacchus jacchus</i>	5/3	1.05	.99	1.04	1.06	1.035	1.980	.991
<i>Callithrix jacchus penicillata</i>	3/5	1.03	1.02	.99	.97	1.003	1.972	1.054
<i>Cebuella pygmaea pygmaea</i>	14/23	.99	1.01	.99	1.01	1.000	1.455	.988
<i>Saguinus fuscicollis nigrifrons</i>	19/15	.97	.97	1.00	.98	.980	2.055	.924
<i>Saguinus midas niger</i>	31/25	.98	1.01	1.03	.99	1.002	2.110	.986
<i>Saguinus oedipus geoffroyi</i>	21/21	1.02	1.01	1.04	1.00	1.017	2.277	.993
<i>Leontopithecus rosalia rosalia</i>	4/6	1.12	1.02	1.05	1.01	1.050	2.596	1.045
<i>Alouatta belzebul</i>	10/10	1.46	1.35	1.37	1.52	1.426	3.910	1.175
<i>Alouatta caraya</i>	10/10	1.36	1.31	1.30	1.38	1.338	3.815	1.201
<i>Alouatta fusca</i>	8/10	1.48	1.36	1.37	1.41	1.405	3.916	1.133
<i>Alouatta palliata aequatorialis</i>	14/13	1.32	1.29	1.26	1.26	1.283	3.945	1.114
<i>Alouatta pigra</i>	7/10	1.24	1.25	1.28	1.25	1.255	3.964	1.057
<i>Alouatta seniculus insulanus</i>	12/10	1.36	1.25	1.40	1.32	1.331	3.868	1.235
<i>Alouatta seniculus seniculus</i>	11/10	1.35	1.20	1.39	1.30	1.309	3.939	1.176
<i>Ateles fusciceps fusciceps</i>	1/2	1.17	1.11	1.15	1.11	1.135	3.430	1.137
<i>Ateles fusciceps robustus</i>	1/3	1.17	1.05	1.04	1.21	1.118	3.332	.943
<i>Ateles geoffroyi vellerosus</i>	22/19	1.19	1.14	1.00	1.12	1.113	3.391	1.008
<i>Ateles paniscus chamek</i>	10/15	1.15	1.07	1.07	1.11	1.100	3.379	1.000
<i>Brachyteles arachnoides</i>	4/4	1.24	1.21	1.08	1.10	1.158	4.039	1.184
<i>Lagothrix flavicauda</i>	2/1	1.08	1.08	1.00	1.14	1.075	3.754	.810
<i>Lagothrix lagotricha poepigii</i>	27/26	1.24	1.15	1.21	1.29	1.223	3.463	.970
<i>Aotus trivirgatus griseimembra</i>	2/4	1.04	.97	.97	.93	.978	2.532	.908
<i>Aotus trivirgatus lemurinus</i>	13/13	1.07	1.02	.99	1.02	1.025	2.505	.999
<i>Aotus trivirgatus trivirgatus</i>	10/19	1.05	1.04	1.01	1.07	1.043	2.528	1.028
<i>Callicebus moloch discolor</i>	12/7	.98	1.03	.98	.98	.993	2.700	1.000
<i>Callicebus torquatus lugens</i>	10/10	1.01	.99	1.03	1.01	1.010	2.733	.983
<i>Cebus apella libinosus</i>	18/21	1.29	1.23	1.17	1.18	1.218	3.267	1.036
<i>Cebus capucinus capucinus</i>	28/25	1.18	1.16	1.08	1.22	1.161	3.196	1.049
<i>Saimiri oerstedii oerstedii</i>	26/19	1.42	1.13	1.12	1.33	1.251	2.205	.974
<i>Saimiri sciureus macrodon</i>	24/11	1.22	1.10	1.15	1.21	1.170	2.314	1.015
<i>Saimiri sciureus boliviensis</i>	6/3	1.28	1.13	1.03	1.23	1.169	2.346	1.022
<i>Cacajao calvus</i>	11/8	1.22	1.15	1.24	1.15	1.190	3.074	1.020
<i>Chiropotes satanas chiropotes</i>	15/16	1.06	1.05	1.10	1.06	1.067	2.683	1.030
<i>Pithecia pithecia</i>	14/11	1.16	1.17	1.20	1.17	1.175	2.777	.988

¹Each dimorphism value is the ratio of mean male dimension to the mean female dimension (dimensions were not \ln transformed prior to calculating sexual dimorphism). Dimensions are as follows: C_1 max: maximum diameter of the lower canine at its base; C_1 min: minimum diameter at right angles to C_1 max; C^1 max: maximum diameter of the upper canine at its base; C^1 min: maximum diameter at right angles to C^1 max. Calculation of CDI is explained in the text.

dominant males for access to resources throughout the year. If so, then canine dimorphism in species with level 4 social structure could be selected in response to both sexual and natural selection.

Morphological data

The morphological data used in this study document the amount of sexual dimorphism in the canines of platyrrhines. Estimates also were made of body size based on the natural logarithms of 1) the area of the upper first molars of females (M^1 area was calculated as the product of M^1 mesiodistal length times buccolingual breadth), 2) the head and body length of adult females, and 3) the body

weight of adult females. Estimates of body-size dimorphism were based on 1) the ratio of male M^1 area to female M^1 area, 2) the ratio of male head and body length to female head and body length, and 3) the ratio of male body weight to female body weight. Tables 2 and 3 summarize these data.

All dental measurements were made on original specimens by using a Wild M5 stereomicroscope fitted with a calibrated reticle. Measurement consistency was plus or minus 3.2% on average; 34 species or subspecies of platyrrhines, representing 15 of 16 extant genera, were sampled (see Table 2 for sample sizes). Each dental sample comes from a small area of the overall range of the species to hold interdeme variation to a minimum.

TABLE 3. Female head and body lengths and body weights (natural logs), head and body length dimorphism, and body-weight dimorphism¹

Species	ln female		Dimorphism	
	H&B length	Body weight	H&B length	Body weight
<i>Callithrix jacchus</i>	5.366	5.943	.981	.950
<i>Cebuella pygmaea</i>	4.970	4.890	.924	.872
<i>Saguinus fuscicollis</i>	5.394	—	.968	—
<i>Saguinus midas</i>	5.529	—	.976	—
<i>Saguinus oedipus</i>	5.606	6.286	1.022	1.039
<i>Leontopithecus rosalia</i>	—	6.174	—	1.152
<i>Alouatta belzebul</i>	6.230	—	1.161	—
<i>Alouatta caraya</i>	—	8.496	—	1.557
<i>Alouatta fusca</i>	6.148	—	1.194	—
<i>Alouatta palliata</i>	6.254	8.695	1.079	1.240
<i>Alouatta seniculus</i>	6.267	8.770	1.102	1.252
<i>Ateles geoffroyi</i>	—	8.973	—	.937
<i>Ateles paniscus</i>	6.292	—	1.009	—
<i>Brachyteles arachnoides</i>	6.351	9.159	1.038	—
<i>Lagothrix flavicauda</i>	6.254	—	1.000	—
<i>Lagothrix lagotricha</i>	6.194	8.678	1.073	1.163
<i>Aotus trivirgatus</i>	5.832	6.802	1.015	1.024
<i>Callicebus moloch</i>	5.884	6.242	1.009	1.325
<i>Callicebus torquatus</i>	6.064	—	.972	—
<i>Cebus apella</i>	5.966	7.693	1.138	1.207
<i>Cebus capucinus</i>	5.999	7.888	1.149	1.430
<i>Saimiri oerstedii</i>	5.652	6.603	1.098	1.212
<i>Saimiri sciureus</i>	5.743	6.430	1.019	1.266
<i>Cacajao calvus</i>	6.319	8.175	.991	—
<i>Chiropotes satanas</i>	5.984	7.972	1.113	—
<i>Pithecia pithecia</i>	5.852	6.965	1.184	1.223

¹Head and body length (H&B) data from Napier (1976). Body-weight data from Hershkovitz (1977); Kay (1973); Leutenegger and Cheverud (1982); van Roosmalen et al. (1981); and Fontaine (1981).

As measures of canine dimorphism, we used the greatest canine diameter and the diameter at right angles to it in the occlusal plane on both upper and lower canines of males and females. In Table 2, four estimates of canine dimorphism are presented: 1) mean male lower canine occlusal length divided by mean female lower canine occlusal length, 2) mean male lower canine occlusal breadth divided by mean female lower canine occlusal breadth, 3) mean male upper canine occlusal length divided by mean female upper canine occlusal length, and 4) mean male upper canine occlusal breadth divided by mean female upper canine occlusal breadth.

The four initial measures of canine dimorphism described above were found to be highly intercorrelated (Table 4). To derive a single measure of dimorphism, a principal-components analysis was undertaken on the correlation matrix of the four male/female measurement ratios of each species. The first principal component (PCI) of this analysis accounts for 92% of the total variance, and

the factor loadings for the four initial variables are very similar (Table 5). A species having male/female canine size of 1.00 for each of the four initial measures of dimorphism will have a PCI value of -2.00 . Therefore we use $PCI/-2.00$ as a canine dimorphism index (CDI). This index is 1.00 if there is no canine dimorphism and theoretically would be 2.00 if male canines were twice as large as those of females. CDIs are presented in Table 2.

To avoid the possibility that the data points are not statistically or biologically independent (e.g., Clutton-Brock and Harvey, 1978; Ridley, 1983), in tests for correlation between dimorphism and body size (i.e., molar area, head and body length, and body weight) we considered two subsets of the data: 1) where any species was represented by more than one subspecies or population, we selected the largest sample to represent that species for the analysis; 2) in a second analysis, we selected the largest sample of one species for each genus.

TABLE 4. Product-moment correlations among the four canine dimorphism estimates (measurements defined in Table 2; all correlations significantly different from zero at the 0.001 level)

	Lower canine		Upper canine	
	Max	Min	Max	Min
Lower canine				
Max	1.00	—	—	—
Min	0.90	1.00	—	—
Upper canine				
Max	0.82	0.86	1.00	—
Min	0.94	0.88	0.84	1.00

TABLE 5. Eigenvalues for the first principal component of the analysis of dimorphism

Lower canine maximum	-0.506
Lower canine minimum	-0.503
Upper canine maximum	-0.485
Upper canine minimum	-0.506

RESULTS

Canine dimorphism and body size

The effects of body size on canine dimorphism are considerable but appear to be more a consequence of the fact that large platyrrhine species tend to have more breeding competition than small species. To assess these effects we calculated the correlations between CDI and \ln female molar area, \ln female head and body length, and \ln female body weight. Irrespective of which measure of body size is chosen, canine dimorphism increases as body size increases in platyrrhines. The product-moment correlation coefficients between CDI and molar area, head and body length, and body weight for species are 0.76, 0.61, and 0.67, respectively (Table 6); when the sample is composed of just one species per genus the correlations between the variables are 0.62, 0.57, and 0.64 (Table 8). All these correlations are significantly different from zero at the 0.01 level or better.

TABLE 6. Correlations between CDI and three measures of body size in platyrrhine species: 1) \ln female M^1 area, 2) \ln female head and body length, and 3) \ln female body weight¹

	R	DF	F-test	P
CDI vs. \ln female M^1 area				
1. All platyrrhines	0.76	25	33.21	.0001
2. Monogamous/polyandrous	0.40	9	1.55	NS
3. Multimale/multifemale low competition or fission/fusion	0.17	3	0.06	NS
4. Multimale/multifemale breeding season competition	0.62	3	1.25	NS
5. Single male or multimale/ multifemale dominance rank competition	0.51	6	1.80	NS
CDI vs \ln female head and body length				
1. All platyrrhines	0.61	21	11.67	.003
2. Monogamous/polyandrous	0.41	8	1.40	NS
3. Multimale/multifemale low competition or fission/fusion	0.59	2	0.53	NS
4. Multimale/multifemale breeding season competition	0.42	3	0.44	NS
5. Single male or multimale/ multifemale dominance rank competition	0.49	4	0.94	NS
CDI vs. \ln female body weight				
1. All Platyrrhines	0.67	18	13.51	.002
2. Monogamous/polyandrous	0.72	6	5.39	NS
3. Multimale/multifemale low competition or fission/fusion	0.66	2	0.75	NS
4. Multimale/multifemale breeding season competition	0.40	3	0.37	NS
5. Single male or multimale/ multifemale dominance rank competition	0.80	3	3.54	NS

¹Just one subspecies is selected to represent each species. For canine and molar dimorphism, samples are used only when each species is presented by three or more specimens of each sex. NS = correlation not significant at $P = .05$.

TABLE 7. Correlations between CDI and three measures of body-size dimorphism in platyrrhine species: 1) dimorphism in female M^1 area (ratio of male molar area to female molar area), 2) dimorphism in head and body length (ratio of male head and body length to female head and body length), and 3) dimorphism in body weight (ratio of male body weight to female body weight)¹

	R	DF	F-test	P
CDI vs. M^1 area dimorphism				
1. All platyrrhines	0.72	25	25.17	.0001
2. Monogamous/polyandrous	0.83	9	17.18	.003
3. Multimale/multifemale low competition or fission/fusion	0.33	3	0.25	NS
4. Multimale/multifemale breeding season competition	0.96	3	26.01	.04
5. Single male or multimale/multifemale dominance rank competition	0.72	6	5.44	NS
CDI vs head and body length dimorphism				
1. All platyrrhines	0.77	21	28.21	.0001
2. Monogamous/polyandrous	0.74	8	8.44	.023
3. Multimale/multifemale low competition or fission/fusion	0.78	2	1.56	NS
4. Multimale/multifemale breeding season competition	0.06	3	0.008	NS
5. Single male or multimale/multifemale dominance rank competition	0.62	4	1.91	NS
CDI vs body weight dimorphism				
1. All platyrrhines	0.62	15	8.83	.01
2. Monogamous/polyandrous	0.17	5	0.12	NS
3. Multimale/multifemale low competition or fission/fusion	Insufficient sample (n=2)			
4. Multimale/multifemale breeding season competition	0.71	2	1.34	NS
5. Single male or multimale/multifemale dominance rank competition	0.74	3	2.48	NS

¹Just one subspecies is selected to represent each species. For canine and molar dimorphism, samples are used only when species is represented by three or more specimens of each sex. NS= correlation not significant at $P = .05$.

Depending on the model chosen, between 32 and 60% of the variance in canine dimorphism is explained by body size.

However, a very different pattern emerges when we attempt to subtract crudely the effects of breeding pattern. Within each level of breeding competition, no significant inter-taxon correlations with body size are found. With the all-species data set (Table 6) there are 12 possible within-group comparisons; there is no significant correlation (i.e., $P < .05$) between CDI and body size in any case.

Canine dimorphism and body-size dimorphism

If we examine the association between canine dimorphism and body-size dimorphism

there are indications that the two are not "produced" by the same selective factors. To assess the effects of body-size dimorphism on canine dimorphism we calculated the correlations between CDI and 1) molar area dimorphism, 2) head and body length dimorphism, and 3) body-weight dimorphism. Irrespective of which measure of body-size dimorphism is chosen, canine dimorphism increases as body-size dimorphism increases in platyrrhines. The product-moment correlation coefficients are 0.72, 0.77, and 0.62, respectively (Table 7). When the sample is composed of just one species per genus, significant correlations are found only between head and body length dimorphism and CDI (Table 8). Thus, it appears that the selective factors influencing body-size dimorphism

TABLE 8. Correlations between CDI and various measures of body size and body-size dimorphism (see Tables 6 and 7 for details)¹

CDI versus	R	DF	F-test	P
Size measures				
ln female molar area	0.62	14	8.29	.01
ln female head and body length	0.57	13	5.66	.04
ln female body weight	0.64	14	9.16	.01
Dimorphism measures				
Molar sexual dimorphism	0.24	14	0.81	NS
Head and body length sexual dimorphism	0.60	13	6.71	.02
Body weight sexual dimorphism	0.49	11	3.17	NS

¹Just one species is selected to represent each genus.

may not be the same ones controlling canine dimorphism, and this emerges most clearly if the effects of phyletic inertia are muted by using only one species per genus.

The same pattern generally emerges when we attempt to subtract crudely the effects of breeding pattern. With the all-species data set (Table 7) of 11 possible within-group comparisons, there is a significant correlation (i.e., $P < .05$) between CDI and body size in just two cases involving monogamous/polyandrous species.

Canine dimorphism and breeding competition

When CDIs of platyrrhine species are segregated by the level of breeding competition, we observe a clear trend toward increased dimorphism with increased intermale competition. CDIs in our sample of platyrrhines range from a low of 0.98 in *Aotus trivirgatus* to the high of 1.42 in *Alouatta belzebul*. Histograms of canine dimorphism are shown in Figure 1 for each level of intermale competition in our scheme. Here all species and subspecies are plotted and no statistical tests for the significance of differences among the groups are calculated.

Level 1 taxa, including species of *Callithrix*, *Saguinus*, *Aotus*, *Callicebus*, *Leontopithecus*, and *Chiropotes*, range from 0.98 (*Aotus trivirgatus griseimembra*) to 1.07 (*Chiropotes satanas*). The 13 lowest-ranking CDI species are all either monogamous or polyandrous.

Level 2 taxa (three *Ateles* species, *Lagothrix flavicauda*, *Brachyteles arachnoides*, and *Pithecia pithecia*) have CDIs between 1.08 and 1.18. Higher CDIs are found in level 3 taxa than in those of level 2 although there is a slight overlap between sampled taxa of

the two levels. Four of five level 3 taxa (two *Saimiri sciureus* subspecies, *Cebus capucinus*, and *Cacajao calvus*) have CDIs between 1.15 and 1.20. Another level 3 taxon, *Saimiri oerstedii*, exhibits considerably more dimorphism (CDI = 1.25) than would be expected given its reportedly low level of intermale competition during the breeding season (Baldwin and Baldwin, 1981; Boinski, 1987). One possible explanation could be that intermale competition is extreme but occurs only at the time when the males reach sexual maturity and is followed by relative stability in adult social interactions. This explanation is supported by the fact that *Saimiri oerstedii* groups contain many more adult females than adult males (Baldwin and Baldwin, 1972).

Level 4 taxa, including *Cebus apella* and all species of *Alouatta* for which behavioral data are available, range from 1.22 (*Cebus apella*) to 1.40 (*Alouatta fusca*). The eight highest-ranking CDI species have social systems in which there is severe and/or protracted intermale competition for dominance. Thus, one might speculate that not only is there competition for access to mates (and perhaps female choice) during the mating season, but there is also competition among males at other times of the year for other limiting resources such as food.

The social structures of two taxa for which we have dimorphism data remain unstudied. Dimorphism in *Lagothrix lagotricha* falls at the lower end of level 4 species in terms of dimorphism—a sharp contrast to *Lagothrix flavicauda*, a level 2 species with low dimorphism. *Alouatta belzebul* has the highest CDI of any platyrrhine in this study, although perhaps not significantly higher than *Alouatta fusca*, a level 4 taxon.

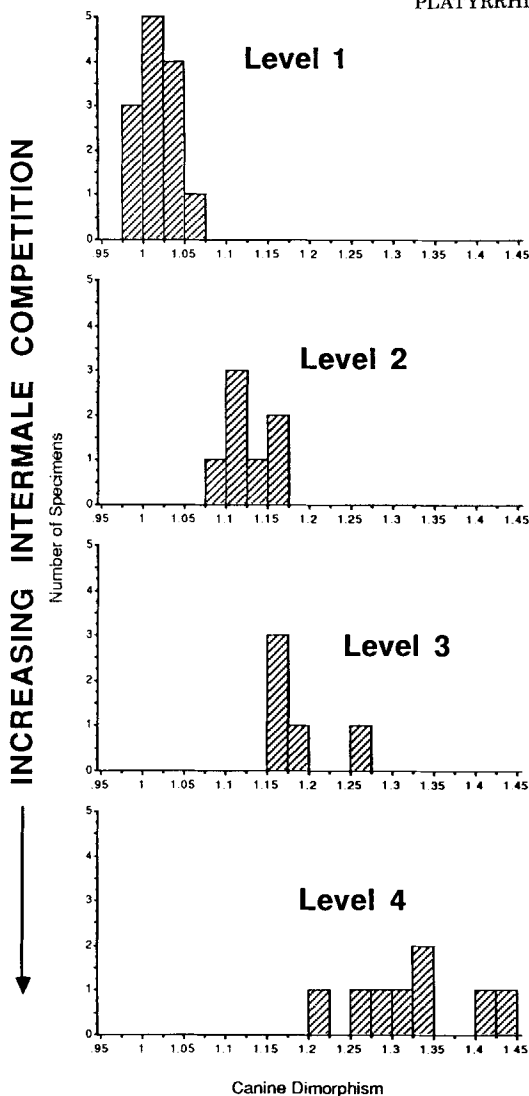


Fig. 1. Histograms of the canine dimorphism index (CDI) for species or subspecies of platyrrhines in the study sample. Separate histograms are provided for species falling within each of the four levels of breeding competition described in the text. Species are not plotted unless their behavior is known. A clear trend is noted for increased dimorphism with increased intermale competition. For individual CDIs, see Table 2.

Joint assessment of the effects of size, size dimorphism, and breeding competition on canine dimorphism

As indicated above, analyzed separately, social structure appears to have the largest effect on canine dimorphism, and body-size dimorphism appears to be a significant but less important factor. Analysis of covariance

(using the general linear model procedure in the Statistical Package for the Social Sciences) was used to assess how much of the variance in CDI is explained by each of these factors and the interactions among them when other factors are controlled for. In this analysis, social structure was treated as an ordered class variable with four states. Three separate analyses were made using each of the three size and size-dimorphism variables and their interaction terms.¹ In model 1 (Table 9), in which 98.7% of the variance in canine dimorphism is explained, the independent variables were 1) competition level, 2) ln female weight, 3) weight dimorphism, 4) an interaction term for competition with weight and 5) an interaction term for competition with weight dimorphism. Of the independent variables examined, competition explains the most variance but body weight also explains a significant proportion of the variance ($P = .04$). Taking the .05 level as the cutoff for significant variance, neither weight dimorphism nor either of the interaction terms has further significant explanatory power once the effects of the other independent variables are controlled. Models 2 and 3 (Table 9) are similar but employ head and body length and molar area as size and dimorphism variables. These analyses give similar results, explaining 97.9 and 95.9% of the variance, respectively. In both cases, competition is a very important explanatory variable. However, in model 2, ln head and body length and head and body length dimorphism and the interaction terms all contribute significantly to explaining the variance in canine dimorphism, whereas in model 3, once the effect of social structure is removed, none of the other factors is significant. Thus, several conclusions are possible: 1) Competition seems to be the most important predictor of canine dimorphism in platyrrhines in all three models. 2) Body size is also a significant but less important predictor in all three models. 3) Body size dimorphism is generally least important but is significant in the case of head and body length. 4) Interaction terms are only significant in the second model, which employs head and body length.

¹One assumption of the analysis of covariance is the identity of slopes of regression equations between the dependent variable and each of the independent variables. In this case, this assumption appears to be violated. Therefore, we used a model with interaction terms for competition with size and competition with dimorphism.

TABLE 9. Assessment of the contributions of three independent variables: 1) sexual competition (a four-level class variable), 2) body size, and 3) body-size dimorphism, to the dependent variable canine dimorphism (CDI)¹

Source	DF	Sum of squares	Mean square	F-value	P
Model 1²					
Model	10	0.2038	0.0204	32.36	.0022
Error	4	0.0025	0.0006		
Corrected total	14	0.2063			
Competition	3	0.0115		9.14	.0323
Body-weight dimorphism	1	0.0029		4.57	.0992 (NS)
Female Body weight	1	0.0055		8.67	.0422
Competition times body-weight dimorphism	3	0.0060		4.75	.0878 (NS)
Competition times body weight	2	0.0028		2.21	.2255 (NS)
Model 2^a					
Model	11	0.2912	0.2647	34.26	.0001
Error	8	0.0062	0.0008		
Corrected total	19	0.2974			
Competition	3	0.0177		7.65	.0098
H & B length dimorphism	1	0.0090		11.63	.0092
Female H & B length	1	0.0053		6.85	.0308
Competition times H & B length dimorphism	3	0.0115		4.96	.0312
Competition times H & B length	2	0.0158		6.841	.0134
Model 3⁴					
Model	11	0.4018	0.0365	38.03	.0001
Error	18	0.0173	0.0010		
Corrected total	29	0.4191			
Competition	3	0.0108		3.75	.0298
Molar dimorphism	1	0.0004		0.44	.5173 (NS)
Female molar size	1	0.0006		0.64	.4355 (NS)
Competition times molar size dimorphism	3	0.0062		2.14	.1301 (NS)
Competition times molar size	2	0.0035		1.21	.3334 (NS)

¹Three separate analyses of covariance (type III) are summarized in which "body size" and dimorphism are based on molar area, head and body (H&B) length, and body weight. Each model also takes into account the effects of interactions between CDI and body size and CDI and size dimorphism. In each model, CDI is the dependent variable.

²Independent variables: competition level, ln female body weight, body-weight dimorphism. Interaction terms: competition with ln female body weight, and competition with body-weight dimorphism. The model explains 98.8% of the variance in CDI.

³Independent variables: Competition level, female head and body length, head and body length dimorphism. Interaction terms: competition with ln female head and body length, and competition with head and body length dimorphism. The model explains 97.9% of the variance in CDI.

⁴Independent variables: Competition level, ln female molar area, molar-area dimorphism. Interaction terms: competition with ln female molar area, and competition with molar area dimorphism. The model explains 95.9% of the variance in CDI.

DISCUSSION

Platyrrhine primates follow the general predictions of Rensch's rule of increased dimorphism with increased body size, and this is correlated at least in part with a tendency for larger species to exhibit more intermale breeding competition than smaller ones. However, this is not to say that selection for

body-size increase by itself selects for a change in social structure, for within taxonomic groups with similar levels of competition no such correlation was found. Among medium and large-sized platyrrhines, there is almost as much variation in canine dimorphism and breeding systems as there is in platyrrhines as a whole. *Saimiri*, *Aotus*, and *Callicebus* are similar in size but range

in the canine dimorphism index from 0.98 to 1.25 and in intermale competition from level 1 through level 3. Likewise, among large platyrrhines, *Ateles*, *Brachyteles*, *Lagothrix*, and *Alouatta* CDI ranges from 1.10 to 1.43 and breeding competition from level 2 to level 4.

These analyses suggest that sexual selection is the primary factor responsible for the observed pattern of canine dimorphism in platyrrhines. Thus, canine dimorphism increases in parallel with increased male-male competition for access to mates.

However, there is also some evidence that natural selection plays an important role in the evolution of increased canine dimorphism. The difference between intermale competition in level 3 and level 4 is essentially a temporal one; males of level 4 species maintain a dominance hierarchy over a protracted period of time whereas those in level 3 do so only during the mating season. Competition for access to females in the two groups may be equally intense and result in the same successful outcome for favored males. However, level 4 dominant males may also be increasing their fitness relative to low-ranking males by greater access to resources such as better feeding localities and sleeping sites or less exposure to predation. Note that we are not invoking feeding niche separation between the sexes—rather only a difference in relative competition among conspecifics for access to these resources. Consideration of natural selection as a cause of sexual dimorphism has focused on the hypothesis that such selection will be reflected by differences in the diets of the sexes (e.g., Clutton-Brock et al., 1977; Clutton-Brock and Harvey, 1978). As far as we are aware, this analysis presents the first actual evidence among primates that natural selection via differential competition for limiting resources can effect the degree of sexual dimorphism expressed in a character. Thus we might predict that while competition for access to females should produce the same amount of dimorphism between level 3 and level 4 species, the additive effect of natural selection produces even greater dimorphism in level 4 species.

Other selective factors are thought to produce canine sexual dimorphism in mammals (see references above) but are unlikely to do so in platyrrhines. For example, all platyrrhine taxa except *Aotus* are diurnal. Therefore, an explanation for variation in dimorphism based on the tendency for diurnal

mammals to have more complex social structures and increased selection for visually mediated secondary sexual characteristics could not account for different degrees of dimorphism within platyrrhines. Again, all platyrrhines are essentially arboreal so increased predation pressures of a terrestrial way of life, where there might be an increased role of male weapons (canines) in protecting the social unit, cannot be invoked to explain increased dimorphism in this group. Finally, feeding niche separation between the sexes is uncommon in platyrrhines and does not provide a general explanation for their canine dimorphism.

Nor does phylogenetic inertia play an important role in the differences observed in canine dimorphism. The considerable variation in dimorphism and breeding system that occurs within platyrrhine subfamilies (e.g., Pitheciinae and Atelinae) and even among species of a single genus (*Lagothrix*, *Alouatta*, *Cebus*, and *Saimiri* for example) attests the evolutionary lability of canine dimorphism. If phylogenetic inertia really were important in platyrrhine canine dimorphism, we would expect to find instances where canine dimorphism was unexpectedly higher or lower than would be predicted for the observed level of intermale competition. We find no evidence for such a situation.

This study of platyrrhine primates demonstrates a very strong link between canine dimorphism and breeding system. Level 1 taxa all have lower CDIs than taxa of any other level. Limited overlap occurs between level 2 and 3 breeding systems but level 3 species tend to be more dimorphic than those of level 2. Finally, level 4 taxa have higher CDIs than the other taxa. Natural selection via differential competition for resources, at least in level 4 taxa, also seems to play a role in the evolution of dimorphism. Other proposed explanations for sexual dimorphism (phyletic inertia, correlated response, and feeding niche separation) do not seem to be particularly important determinants of platyrrhine canine sexual dimorphism. As more is learned about the breeding systems of platyrrhines it will be desirable to further refine the system used here for measuring breeding competition. For example, the great variation among level 4 taxa, including *Cebus* and *Alouatta*, begs the question of whether there might be unappreciated variations in the breeding competition of species in this assemblage. Also, it should become possible to bring in other factors we have neglected be-

cause the data are wanting: some of the variation within the levels of our scheme may result from intermale competition among juveniles and young adults as stable breeding systems are initially established.

These results are also notable in illustrating that there is generally a very low correlation between canine dimorphism and body-size dimorphism. This suggests that different factors are controlling the evolution of dimorphism in body size and canine size, a phenomenon previously reported by Oxnard and colleagues (Oxnard 1983, 1987; Oxnard et al., 1985).

In this analysis we have dealt only with dimorphism—the relative size of male and female canines—not with the absolute or relative size of either male or female canines itself. For example, this analysis does not distinguish between one species in which both males and females have large canines for their body size and another in which both males and females have small canines. The absolute sizes of female and male canines will need to be studied before we can gain a more balanced understanding of the selective factors in canine evolution. Suppose, hypothetically, that the larger-canined males of a species are more successful in agonistic encounters with other males over access to mates. Suppose also that the females of our hypothetical species have their own dominance hierarchy established by agonistic encounters with other females in relation to some other resource like food. In this case, large canines might be selected in both males and females. One wonders whether a selective pattern of this nature might be operating in diurnal prosimians. Many of these animals have polygynous social systems and large, sharp canines in both sexes.

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