**Sexual selection on male vocal fundamental frequency in humans and other anthropoids**

Short title: Sexual selection and primate vocalization frequencies

David A. Putsa,b,1, Alexander K. Hilla, c, Drew H. Baileyd, Robert S. Walkere, Drew Rendallf, John R. Wheatleya, Lisa L. M. Wellingg, Khytam Dawoodh, Rodrigo Cárdenash, Robert. P. Burrissi, Nina G. Jablonskia, Mark D. Shrivera, Daniel Weissh, Adriano R. Lameirajk, Coren L. Apicellal, Michael J. Owrenm, Claudia Barellin, Mary E. Glenno, and Gabriel Ramos-Fernandezp

1. Department of Anthropology, Pennsylvania State University, University Park, PA 16802
2. Center for Brain, Behavior, and Cognition, Pennsylvania State University, University Park, PA 16802
3. Current address: Department of Anthropology, University of Washington, Seattle, WA 98185
4. School of Education, University of California, Irvine, CA 92697
5. Department of Anthropology, University of Missouri, Columbia, MO 65211
6. Department of Psychology, University of Lethbridge, Lethbridge, Alberta, Canada T1K 3M4
7. Department of Psychology, Oakland University, Rochester, MI 48309
8. Department of Psychology, Pennsylvania State University, University Park, PA 16802
9. Department of Psychology, Northumbria University, Newcastle upon Tyne NE1 8ST, UK
10. Department of Anthropology, Durham University, Durham DH1 3LE, UK
11. Pongo Foundation, Papenhoeflaan 91, 3421XN Oudewater, the Netherlands.
12. Department of Psychology, University of Pennsylvania, Philadelphia, PA 19104
13. OSV Acoustical Associates and Emory University, Atlanta, GA 30322
14. Sezione di Biodiversità Tropicale, Museo delle Scienze, Trento, Italy 38122
15. Department of Anthropology, Humboldt State University, Arcata, CA 95521
16. CIIDIR Unidad Oaxaca, Instituto Politecnico Nacional, Mexico and C3-Centro de Ciencias de la Complejidad, Universidad Nacional Autonoma de Mexico, Mexico 04510.

1. Corresponding author. 218 Carpenter Building, Department of Anthropology, Pennsylvania State University, University Park, PA 16802. Phone: 814-867-0354. Email: dap27@psu.edu

**Abstract**

In many primates, including humans, the vocalizations of males and females differ dramatically, with male vocalizations and vocal anatomy often seeming to exaggerate apparent body size. These traits may be favored by sexual selection because low-frequency male vocalizations intimidate rivals and/or attract females, but this hypothesis has not been systematically tested across primates, nor is it clear why competitors and potential mates should attend to vocalization frequencies. Here we show across anthropoids that sexual dimorphism in fundamental frequency (*F*0) increased during evolutionary transitions toward polygyny, and decreased during transitions toward monogamy. Surprisingly, humans exhibit greater *F*0 sexual dimorphism than any other ape. We also show that low-*F*0 vocalizations predict perceptions of men’s dominance and attractiveness, and predict hormone profiles (low cortisol and high testosterone) related to immune function. These results suggest that low male *F*0 signals condition to competitors and mates, and evolved in male anthropoids in response to the intensity of mating competition.

**Introduction**

Explaining why sexual dimorphisms evolve is central to understanding the evolution of primate mating systems and social organization. In many primate species, the vocalizations of males and females differ dramatically, with male vocalizations and vocal anatomy often seeming to exaggerate the appearance of body size (1-7). Among humans, men’s approximately 60% longer vocal folds (8, 9) contribute to an average rate of vocal fold vibration during phonation (fundamental frequency, *F*0) that is about five standard deviations below women’s (5). To human listeners, utterances lower in *F*0 are perceived as being deeper in pitch and as emanating from larger individuals (10, 11). The evolutionary reasons for such apparent size exaggeration have been the subject of speculation since Darwin noted the pubertal enlargement of male vocal structures and their intensified employment during the breeding season in many mammals (12).

Some have suggested that male vocalizations evolve to intimidate male competitors and/or attract mates (6, 13). For example, among orangutans, lower-ranking males avoid long calls given by higher-ranking males (14), suggesting that acoustic cues convey threat-potential to conspecifics. Several studies in humans suggest that *F*0 has relevance under both inter- and intrasexual competition: Experimentally lowering *F*0 increases perceptions of men’s dominance and attractiveness (11, 15), and raising *F*0 increases women’s vocal attractiveness (16, 17). However, little is known about whether these effects persist in unmanipulated speech when *F*0 and other acoustic parameters vary naturally and simultaneously.

Moreover, it is unclear why *F*0 should signal formidability to same-sex competitors or mate value to potential mates; *F*0 is only weakly associated with body size (5, 7, 18-20) and perhaps strength (5, 21) in humans, although *F*0 may be modulated according to relative formidability (22) and mate quality (23, 24). Steroid hormones may provide a link between *F*0 and condition. Growing evidence indicates that glucocorticoids such as the stress hormone cortisol negatively interact with testosterone in affecting both immune function and the expression of secondary sex traits (25-27). Infection stimulates cortisol production (28), which downregulates androgen receptors and inhibits the action of testosterone on target tissues (29-33). Hence, testosterone should be more potent in individuals in good condition with low immune system activation. In humans, positive relationships between testosterone and immune response to a vaccine (34), and between testosterone and both facial attractiveness (34) and dominance (35), were stronger in males with low cortisol. Furthermore, the interactive effect of testosterone and cortisol on attractiveness was mediated by immune function, supporting the stress-linked immunocompetence handicap hypothesis (SL-ICHH) that testosterone-related traits that interact with cortisol are linked to immuno­competence (34). Although previous studies have found negative relationships between men’s testosterone and *F*0 (5, 20), it is unknown whether testosterone and cortisol negatively interact in predicting *F*0, as the SL-ICHH would suggest if *F*0 reflects underlying condition.

More generally, scant evidence exists to support a role for sexual selection in shaping *F*0 and other vocal sexual dimorphisms across primates (6), and there are plausible alternative hypotheses: *F*0 dimorphism may represent a byproduct of selection for greater male size or long-distance transmission of male calls (36), or reflect selection for sex identification. The latter two hypotheses predict relationships between habitat and *F*0 dimorphism. In general, open, terrestrial habitats are poor acoustic environments in which sounds greatly attenuate over distance compared with arboreal habitats. This is particularly true for the heights at which each habitat’s primate residents tend to communicate and for lower frequency sounds (37). Thus, all else equal, selection for long-distance transmission of male calls should tend to produce relatively lower frequency male calls in arboreal environments than in terrestrial ones (for caveats, see Discussion). Likewise, because primates are both more visible and more sexually dimorphic in terrestrial species than in arboreal ones (38), selection for sex identification should favor greater *F*0 dimorphism in arboreal species.

Here, we report the results of three studies designed to clarify the evolution of sexual dimorphism in *F*0. In Study 1, we examined the evolution of *F*0 dimorphism as a function of mating system, as well as body mass dimorphism and habitat, across anthropoid primates. Studies 2 and 3 focused on humans. Not only are humans of special interest, but they are also highly useful as a model organism: There is strong evidence implicating sexual selection in the evolution of human *F*0 (39, 40), as well as a unique richness to the data available for addressing the questions outlined above, as we detail below. In Study 2, we tested the stimulus-response properties of *F*0 on intrasexual competitiveness in humans by examining the independent contributions of *F*0 and other acoustic parameters related to assessments of attractiveness and dominance. In Study 3, we explored the indexical value of *F*0by testing the SL-ICHH prediction that *F*0 will be more strongly linked to testosterone in individuals with low cortisol.

**Study 1: *F*0 across anthropoid primates**

**Methods**

Please refer to *SI Materials and Methods* for additional details.

We obtained recordings of nonhuman primate calls from our own fieldwork and by contacting other primatologists. From these, we selected 1721 files, such that each was without substantial background noise and was produced by a single individual of known species, sex, and adult status. The acoustic properties of primate calls vary across call types and contexts (13). We chose to utilize measurements across all available call types (but see also Supplementary Results) rather than, for example, selecting only calls believed to be analogous across species, or only calls shared between males and females. Our reasoning was that averaging across call types should maximize our ability to capture information about the physical properties of the sound source (e.g., vocal fold length and thickness), especially if some call types may provide more information than others. If *F*0 is driven by underlying anatomy—as it is fundamentally—then any influence on *F*0 should be manifest similarly across call types, even if not specifically adapted to be, because the same anatomy supports the production of all calls. Although it is possible that some other special mechanism of vocal fold action is in play for some calls (e.g., loud calls compared to close calls), this is not a given and certainly cannot be assumed a priori. Moreover, it is unclear whether call types are truly analogous across species, which complicates comparisons of only a particular call type or set of types. In addition, the repertoire shared between males and females can be very limited in some primate species, occasionally down to one call type, as in the orangutan. Finally, we do not know of any reason that our sampling procedures would have systematically biased our sample, and the size of our sample—the largest ever compiled for this type of research—should reduce any bias due to random sampling.

Files were measured as uncompressed .WAV or .AIFF files using the acoustic analysis software Praat version 5.3. *F*0 was measured from each file by identifying in the raw waveform a segment in which cycles were clearly discernible. Cycles were counted along this segment up to 20 cycles, and then divided by the duration of the interval to calculate *F*0. This procedure was repeated for a second segment, if possible (78% of files, *n* = 1343). Mean *F*0 values from each recording were averaged with all other mean *F*0 values per sex to arrive at separate male and female *F*0 averages for each species (Table S2). Between-segment reliability was high for files with two measurable segments (Cronbach’s *α* = 0.973). First segments of a randomly chosen 11% of files (*n* = 184) were re-measured to determine intra-measurer reliability, which was very high (Cronbach’s *α* = 1.000). Body size, habitat, and mating system were obtained from the literature (Table S2).

Mating system was utilized as a proxy for the intensity of sexual selection (41, 42). We categorized mating system as monogamous, promiscuous, or polygynous (43) rather than using an interval-level measure such as socionomic sex ratio, as such measures often vary widely within species and hold uncertain relationships to the intensity of intermale competition (41, 44, 45). Promiscuity differs from monogamy and polygyny in that females far more frequently copulate with multiple males in promiscuous species. Although evidence suggests intermediate levels of male contest competition in promiscuous species, the ability of males to monopolize females varies widely (46), other mechanisms of sexual selection such as sperm competition are more salient (47, 48), and the degree of sexual dimorphism relative to monogamous or polygynous species varies widely by trait (47). Such apparent diversity in the mechanisms and intensity of sexual selection, as well as their influence on sexual dimorphism, precludes straightforward predictions regarding *F*0 dimorphism in promiscuous species. By contrast, predictions regarding monogamous and polygynous species are clearer: In polygynous species, some males are able to monopolize multiple mates, whereas in monogamous species, males do not, leaving fewer males unmated. Hence, sexual selection—particularly, intrasexual selection—tends to be more intense in polygynous than in monogamous primates, which are less dimorphic in size and weaponry (47). Dimorphism in *F*0 was therefore predicted to increase during evolutionary transitions toward polygyny and decease during transitions toward monogamy.

Habitat was categorized as arboreal, terrestrial, or arboreal/terrestrial. We conducted phylogenetically-informed analyses using a consensus phylogeny for all species represented in our sample (49; Fig. 1) and assessed correlated evolution among our variables with phylogenetic generalized least squares regression using the caper package, v.0.5.2, in R (50).

**Results**

Across analyses, *F*0 and *F*0 dimorphism exhibited strong phylogenetic signals (*ʎ* > 0.8). In general, New World primates showed little sexual dimorphism in *F*0, averaging a mean *F*0 dimorphism (male *F*0/female *F*0) of 1.05 across 7 species, while male cercopithecines averaged half of the *F*0 of females (mean *F*0 dimorphism = 0.48 across 10 species). With a similar *F*0 dimorphism of 0.51, humans surprisingly exhibited the greatest dimorphism that we measured in any ape.

We tested whether increases in body size predict decreases in *F*0 across species for each sex. Previous studies relied on published acoustic data measured using varying methodologies and either averaged male and female measurements (51) or included only males (36). In our data, body mass negatively predicted *F*0 (both variables natural log-transformed) in males and females (Table 1). These results suggest that body size constrains the evolution of primate call frequencies in both sexes (51).

We then regressed *F*0 dimorphism on mating system, controlling for body size dimorphism (male mass/female mass). Because diversity in the mechanisms and intensity of sexual selection precludes straightforward predictions regarding *F*0 dimorphism in promiscuous species (see above), we excluded promiscuous species from this analysis; however, promiscuous species appear intermediate in *F*0 dimorphism (Fig. 2a). We found that greater *F*0 dimorphism evolves in transitions to polygyny than in transitions to monogamy (Table 1, Fig. 2c). In this model, changes toward greater *F*0 dimorphism also tended to be accompanied by decreases in body size dimorphism. Humans were treated as polygynous (52) and exhibited *F*0 dimorphism that was outside the range of monogamous species (Fig. 2a); however, we obtained similar results when humans were treated as monogamous, or excluded from the analysis (Table 1).

Finally, we tested the relationship between *F*0 dimorphism and habitat. If male vocalizations are selected primarily to propagate over distance, or if *F*0 dimorphism evolves primarily for sex discrimination, then arboreal species should tend to exhibit relatively lower male *F*0 than terrestrial species. We found the reverse: Arboreal primates showed less *F*0 dimorphism than terrestrial primates when *F*0 dimorphism was regressed on habitat and mass dimorphism (Table 1).

**Study 2: *F*0, dominance, and attractiveness in humans**

**Methods**

Please refer to *SI Materials and Methods* for additional details.

Two hundred fifty-eight female (mean age ± s.d. = 20.0 ± 1.6 y) and 175 male (20.1 ± 1.7 y) students from Michigan State University provided written consent to participate in this study approved by the university’s Institutional Review Board. Participants were recorded reading a standard voice passage (53) in an anechoic, soundproof booth using a Shure SM58 vocal cardioid dynamic microphone (frequency response: 50-15,000 Hz) positioned at approximately 30 degrees and 9.5 cm from the speaker’s mouth, and connected to a desktop computer via a Sound Devices USBPre 2 preamplifier. Voices were recorded in mono at a sampling rate of 44,100 Hz and 16-bit quantization, and saved as uncompressed .WAV files. Recordings were rated by 558 female (19.1 ± 2.4 y) and 568 male (19.4 ± 1.8 y) students from The Pennsylvania State University. Each female recording was rated by 15 men for attractiveness for short- and long-term romantic relationships using 7-point Likert scales. Each male recording was rated by 15 men for dominance (7-point scale) and 15 women for short- and long-term attractiveness. Ratings were averaged to produce composite ratings of short- and long-term attractiveness for each recording, and dominance for each male recording.

Recordings were analyzed using Praat version 5.3 for mean *F*0, standard deviation in *F*0 across the utterance (*F*0-SD), duration, number of voice breaks, harmonics, four measures of jitter (cycle-to-cycle variation in *F*0), and five measures of shimmer (cycle-to-cycle variation in amplitude) using the “voice report” function in Praat (Table S3). Pitch floors were set to 75 Hz and 100 Hz, and pitch ceilings were 300 Hz and 500 Hz, for men and women, respectively. Otherwise, default settings were used. We also measured the first four formant frequencies (*F*1- *F*4, Table S3). Formants were measured at each glottal pulse, averaged across measurements, and then used to compute formant position (*Pf*), the average standardized formant value for the first four formants (5).

Using SPSS v.22, we performed a separate male and female principal components analyses to reduce the number of acoustic control variables (SI Materials and Methods, Table S4), and we utilized multiple regression to examine the effects of acoustic parameters on perceptual variables.

**Results**

*F*0 predicted men’s perceived dominance to heterosexual male listeners (*β* = -0.43, *p* < 0.0001) and attractiveness to heterosexual female listeners for both prospective short-term (*β* = -0.36, *p* < 0.001) and long-term (*β* = -0.32, *p* = 0.001) romantic relationships (Table S5). When perceived dominance and short-term attractiveness were entered into a multiple regression to predict men’s *F*0 (model *F*2,171 = 12.99, *p* < 0.0001, *R*2 = 0.13), dominance negatively predicted *F*0 (*β* = -0.30, *p* = 0.001), but short-term attractiveness did not (*β* = -0.09, *p* = 0.314), suggesting a stronger role for male contests than female choice in shaping men’s *F*0. *F*0 did not predict women’s attractiveness to men for either short- (*β* = 0.03, *p* = 0.695) or long-term (*β* = -0.03, *p* = 0.722) relationships when other acoustic parameters were statistically controlled (Table S5). These results are thus more consistent with sexual selection (primarily intrasexual selection) on males, rather than intersexual selection on females, influencing the evolution of human *F*0 dimorphism.

**Study 3: *F*0 and hormonal profiles in humans**

**Methods**

Please refer to *SI Materials and Methods* for additional details.

Participants from The Pennsylvania State University provided written consent to participate in this study approved by the university’s Institutional Review Board. Fifty-three normally-cycling women (mean age ± s.d. = 19.4 ± 1.6 y) and 62 men (19.9 ± 2.0 y) were recorded in an anechoic recording booth in a quiet room (Sample 1), and58 men (19.9 ± 1.2 y) were recorded in a quiet room (Sample 2) using the equipment and methods in Study 2 above.

Participants rinsed their mouths with water before providing two saliva samples of 1-2 ml each via passive drool approximately 30 (Sample 1) or 20 (Sample 2) min apart. From each sample, 0.5 ml of saliva was aliquotted into a third tube, which was shaken and then frozen at -20°C until analysis by the Johns Hopkins Center for Interdisciplinary Salivary Bioscience Research (Baltimore, MD) using Salimetrics® kits. Samples were analyzed in duplicate via enzyme immunoassay. Duplicates correlated highly for both cortisol and testosterone (all *r* ≥ 0.97, *p* < 0.0001)*,* and were consequently averaged. For cortisol assays, sensitivity is <0.003 μg/dL, and average intra-assay coefficient of variation is 3.5%. For testosterone assays, sensitivity is <1.0 pg/mL, and average intra-assay coefficient of variation is 4.6%.

Hormone data were natural log-transformed to correct skew prior to analysis. We utilized multiple regression to examine the effects of cortisol and testosterone on *F*0 and statistically controlled for diurnal decreases in cortisol and testosterone (54), but results were similar without controlling for these effects (Table S6, Figs. S1, S2).

**Results**

In women, *F*0 was unrelated to cortisol, testosterone, and their interaction (Table S6). However, in both male samples, cortisol and testosterone interacted in predicting *F*0 (Sample 1: *β* = 0.36, *p* = 0.007; Sample 2: *β* = 0.28, *p* = 0.033; Table S6) such that testosterone was negatively related to *F*0 only in low-cortisol men (median split for cortisol; Sample 1: partial *r* = -0.44, *p* = 0.018; Sample 2: partial *r* = -0.40, *p* = 0.034; see also Fig. S2). This pattern of relationships between hormones and a putative sexually selected trait has been found to indicate men’s immune function (34), as well as attractiveness (34) and dominance (35), and is consistent with the SL-ICHH (34).

**Discussion**

Our data supported the sexual selection hypothesis: *F*0 dimorphism increased with evolutionary changes toward polygyny and decreased with transitions toward monogamy across anthropoid primates. Although our primary analyses averaged across call types in order to maximize data, we also examined patterns of *F*0 dimorphism across mating systems using only those calls for which we had examples from both sexes (SI Results, Fig. S3). Patterns were similar across analyses, indicating that the observed pattern in our primary results does not merely reflect sex differences in the use of particular call types, which may differ in *F*0. Our data also indicate that sex differences in *F*0 result mainly from selection on males rather than females: We observed greater *F*0 dimorphism in polygynous species, where male sexual selection is stronger, and *F*0 affected components of men’s but not women’s mating success. These results thus provide a plausible explanation for the prior finding that *F*0 predicted men’s but not women’s reproductive success among Hadza foragers (55). If low *F*0 augments male reproductive success by increasing access to mates in some other anthropoids, as appears to be the case in humans, then we would expect males to evolve relatively lower *F*0 when mating competition intensifies during transitions to polygyny, as we found.

By contrast, *F*0 dimorphism appears unlikely to be a byproduct of greater male size: With mating system controlled, *F*0 dimorphism decreased with relative male size. Although *F*0 dimorphism likely facilitates sex identification, if it evolves primarily for this function, then one might expect it to be greater in arboreal species where visibility is obscured, and in monogamous species where the sexes are otherwise less dimorphic (38), yet in both cases we found the opposite.

Previous work indicates that lower frequency male loud calls are selected for propagation over distance (36), but our finding that male *F*0 is relatively lower in terrestrial species than in arboreal species suggests that selection for long-distance male calls is not the only or primary influence on *F*0 dimorphism across species’ vocal repertoires. Lower frequencies should, *ceteris paribus*, aid long-distance transmission to a greater degree in arboreal habitats than in terrestrial ones for the heights at which primates normally communicate. Although we view our measure of environmental influence on acoustic transmission (arboreal vs. terrestrial) as a safe parsing for such a broad sample encompassing a panoply of subtly different habitat types, it is admittedly crude, and environmental effects on acoustic transmission are complex. Other variables such as type of call, location in an arboreal environment, and ambient noise are likely to play a role (37), and an exploration of these influences should be taken up more fully as more precise data accumulate. However, such possible environmental influences might be expected to weaken any relationship between *F*0 dimorphism and arboreal vs. terrestrial habitat rather than producing an effect that is opposite the predicted direction. Our results thus challenge the hypothesis that overall *F*0 dimorphism primarily evolves when male calls are selected for long-distance transmission, and our findings are more consonant with elevated male-male competition in terrestrial compared with arboreal species (56, 57).

Inspection of Fig. 1 indicates an increase in *F*0 dimorphism from the last common ancestor of the apes to modern humans, culminating in humans exhibiting the greatest *F*0 dimorphism of all apes. These results contrast sharply with moderate human body mass dimorphism and negligible canine length dimorphism, which some have suggested indicate weak sexual selection in ancestral humans (58). However, unlike other primates, in humans, female adiposity greatly exceeds that of males, producing modest overall mass dimorphism despite the fact that males possess 60% more muscle mass, and men fight with handheld weapons and fists rather than teeth in combat (57, 59). These unique features preclude conclusions about the strength of human sexual selection based on overall mass or canine size dimorphism (60, 61). Yet, if mating competition also tends to decrease male *F*0 relative to female *F*0 across primates, then *F*0 dimorphism has the potential to elucidate human sexual selection in ways that comparisons of body mass or canine size cannot. Our results suggest that, despite widespread pair-bonding and contrary to some claims (58, 62), ancestral human mating should not be viewed as fundamentally monogamous (52).

When phylogeny and mating system were statistically controlled, evolutionary changes toward greater *F*0 dimorphism were associated with changes toward less body size dimorphism and vice versa. Perhaps where mates are won mainly through direct fighting, males receive less benefit from exaggerating size acoustically and must instead invest in mass that is useful in contests. Conversely, acoustic threats and displays may be more effective when body mass or fights are especially costly, or where female choice is more important to male fitness. In humans, male *F*0 was indeed important in mate attraction, yet *F*0 more strongly predicted perceptions of men’s dominance, consistent with previous experimental evidence (22, 39). Masculinity in men’s faces is similarly perceived as aggressive across human societies, whereas the influence on attractiveness is more variable in magnitude and direction (63). While such male traits appear better designed to function in male contests than in female choice (57), it remains possible that female choice is relatively more important in humans compared to other polygynous primates, and that stronger female choice tends to favor lower male *F*0 and more modest size dimorphism among polygynous primates. It is also possible that the unprecedented lethality imbued to human fighting with the advent of handheld and projectile weapons elevated the importance of threats and deference in relation to physical attacks (52, 57). Finally, in arboreal primates, large body mass may impose additional energetic costs, as well as increasing the risk of injury from falls. Although the relatively limited number of species in our sample prohibited exploring interactions between mating system, habitat, and body mass dimorphism, we note that the single species in our sample with the greatest *F*0 dimorphism, *Cercopithecus campbelli*, is polygynous, partly arboreal, and exhibits body mass dimorphism comparable to that of humans. In other words, one can speculate that male *C. campbelli* achieve polygyny partly via acoustic threats and/or mate attraction while body mass is constrained by partial arboreality.

In many species, males exaggerate size to intimidate conspecifics, but attention to these exaggerations is likely maintained by a continued association between apparent size and formidability (4). Although *F*0 decreased with increasing body size in both sexes across primate species, body size only weakly predicts *F*0 in adult humans (5, 7). However, our data show that cortisol and testosterone interact to predict men’s *F*0 in a pattern that has previously been found to predict men’s dominance (35), attractiveness (34), and immunocompetence (34), and hence that *F*0 is likely to reveal male condition to same-sex competitors and potential mates.

Our results thus not only demonstrate a likely influence of sexual selection in the origins and maintenance of sexual dimorphism in *F*0 across anthropoids, but also suggest that male contests, and to a lesser degree female mate choice, favor low male *F*0 as a signal of condition, shedding new light on the intensity and mechanisms of sexual selection in humans and other primates. Future research should extend these comparative investigations to vocal tract resonance frequencies, which have been implicated in mating competition and size exaggeration in several mammals (1, 4, 7), including humans (11, 15).

**Ethics.** Participants provided informed written consent. The ethics committees of Michigan State University and The Pennsylvania State University approved these studies, and all procedures adhered to the Declaration of Helsinki.

**Data accessibility.** Data deposited in Dryad Digital Repository: <http://datadryad.org/submit?journalID=RSPB&manu=RSPB-2015-2830>

**Authors’ contributions.** Conception and design of the experiments: DAP, AKH, RAC, RPB; collection, analysis, and interpretation of data: DAP, AKH, DHB, RSW, DR, JRW, LLMW, RPB, ARL, CLA, MJO, CB, MEG, GR-F; drafting the article or revising it for important intellectual content: DAP, AKH. All authors approved the final version of the manuscript.

**Competing interests.** We declare we have no conflict of interest.

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**Figure Legends:**

Fig. 1. Phylogenetic tree of anthropoid primates included in Study 1 for which data were available on at least 2 vocalizations from each sex (mean number of vocalizations: females = 38.6, males = 22.1; max: females 181, males = 155; Table S1). Tree was constructed using a consensus phylogeny for all species in our sample from the 10kTrees website (http://10ktrees.fas.harvard.edu/). Sexual dimorphism (male/female) in *F*0 is shown in the column to the left of species names. Inferred ancestral states were determined using squared change parsimony and are shown at nodes on the tree (colored to highlight evolutionary trends).

Fig. 2. Sexual dimorphism in vocal *F*0 as a function of mating system. In panel (a), data points represent individual species, and horizontal bars represent mean *F*0 dimorphism for each mating system. Sexual dimorphism in *F*0 is most extreme in polygynous anthropoid primates and lowest in monogamous species. This remains true after adjusting for body mass dimorphism (b), and after adjusting for both body mass dimorphism and phylogenic non-independence (c). Panel (c) shows results from phylogenetic generalized least squares regression with statistics for the effect of mating system on *F*0 dimorphism. In panels (b) and (c), least-squares regression lines with 95% CI are plotted, mating system refers to polygyny (coded as -1) vs. monogamy (coded as 1), and adjusted values are residuals after each variable was regressed on body mass dimorphism. Key for independent contrasts in panel (c) is shown in panel (d).