Grandmothering life histories and human pair bonding

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The evolution of distinctively human life history and social organization is generally attributed to paternal provisioning based on pair bonds. Here we develop an alternative argument that connects the evolution of human pair bonds to the male-biased mating sex ratios that accompanied the evolution of human life history. We simulate an agent-based model of the grandmother hypothesis, compare simulated sex ratios to data on great apes and human hunter-gatherers, and note associations between a preponderance of males and mate guarding across taxa. Then we explore a recent model that highlights the importance of mating sex ratios for differences between birds and mammals and conclude that lessons for human evolution cannot ignore mammalian reproductive constraints. In contradiction to our claim that male-biased sex ratios are characteristically human, female-biased sex ratios are reported in some populations. We consider the likelihood that fertile men are undercounted and conclude that the mate-guarding hypothesis for human pair bonds gains strength from explicit links with our grandmothering life history.

We call attention to evidence that connects the evolution of human pair bonds to the male-biased sex ratios in fertile ages that characterize human populations. As in mammals generally, age-specific mortality is higher in males than in females (e.g., refs. 1–3). However, this difference is overshadowed by a distinctive feature of human life history: Oldest ages at parturition are about the same in humans as in other living hominids, the great apes (4, 5), whereas longevity is substantially greater and male fertility continues to older ages (6). Exceptional longevity with a distinctive postmenopausal life stage (7–9) may have evolved in our lineage when grandmothers’ subsides for weaned dependents allowed mothers to have next babies sooner. According to this grandmother hypothesis (10–16), longevity increased as longer-lived grandmothers could help more and so left more longer-lived descendants of both sexes. Women’s postfertile life stage (7) produces a bias in the sex ratio of fertile adults with repercussions for mate strategies. As longevity increased, older-aged males expanded the pool of competitors for the still-fertile females. With more competitors for each fertility, males’ average success in finding new mates inevitably declined until defending a current mate became the better option. Our distinctive life history thus supplies previously unrecognized support for a mate-guarding hypothesis for the evolution of human pair bonds.

Here we simulate hominid mating sex ratios with an agent-based model of the evolution of human longevity via grandmothering (13, 15). We then compare simulated sex ratios to demographic data from both great apes and human hunter-gatherers. Having identified the human bias, we connect it to increased male payoffs for mate guarding, noting some broad patterns in humans, the tradeoffs observed in other taxa, and a history of modeling in which increased guarding is the likely outcome of more competing males.

We then consider a recent model (17) of the evolution of sex roles built to show that anisogamy, the primary sex difference of large (female) and small (male) gametes (18), is insufficient by itself to explain why females care for offspring more often whereas males more often compete for mates. Mating sex ratios are decisive for mating strategies in this model. Although it does not include guarding as an option, it does highlight a connection between mating sex ratios and broad differences between mammals and birds. We elaborate key phylogenetic constraints, consider recent work on birds, and underline features of mammalian offspring production that temper direct application of the model to mammals, including hominids.

An important challenge to our claim that human life history entails male-biased mating sex ratios comes from reports of female bias in some human populations. We identify common measurement problems and link men’s age-specific fertilities to status hierarchies, concluding that the mate-guarding hypothesis remains both promising and directly relevant to explaining the long-recognized importance of male alliances in human societies.

Grandmothering Simulations

We track mating sex ratios through simulations of an agent-based model first built to investigate the evolution of human longevity via grandmothering (13, 15). For reasons elaborated below, we follow both the adult sex ratio (ASR), defined as the ratio of males to females in the fertile ages, and the operational sex ratio (OSR), which counts only the subset of adults currently capable of a conception (19) (see Supporting Information for model parameters and definitions).

Elsewhere we have shown that simulations of this model result in two life history equilibria: a great ape-like one with no grandmothering and a human hunter–gatherer–one like one when grandmothers’ subsidies allow mothers to have next babies sooner (15). Not surprisingly, each equilibrium is associated with distinct, characteristic sex ratios in the fertile ages (Fig. 1). Of 30 simulations without grandmothering run over a million years, the ASR (Fig. 1A) fluctuates around an average of 0.77 males for every female whereas OSR (Fig. 1C) remains at about 50.

Significance

Pair bonds are universal in human societies and distinguish us from our closest living relatives. They characteristically involve men’s proprietary claims over women—mate guarding—which in animals generally is both predicted and observed to be more frequent when sex ratios in the fertile ages are male biased. A marked male bias in the fertile ages evolved in our lineage as longevity increased without an extension of female fertility. We compare the sex-ratio shift in simulations of the grandmother hypothesis to demographic data from chimpanzees and human hunter–gatherers then connect the expanded proportions of older men to benefits for mate guarding, the evolution of pair bonds, and the long recognized importance of male alliances in human social life.

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To evaluate the effect of grandmothering on the mating sex ratios, we use the end point of a simulation chosen at random to start 30 new simulations with grandmothering. All 30 simulations evolve away from the previously stable ape-like mating sex ratios toward values roughly twice higher. ASR (Fig. 1B), initially female-biased, increases with grandmothering to about 1.56 fertile males for every female in the fertile ages, whereas OSR (Fig. 1D) rises to ~111 available males for every female ready to conceive.

Life Tables
To see whether the simulated values are empirically plausible we compare them to the same sex ratios calculated using life tables for living populations. Our empirical comparison for the simulated ape-like equilibrium is limited to chimpanzees because they are the only apes for which life tables based on multiple, long-studied populations have been published. Bonobo population dynamics are still poorly known. We expect, based on arguments here, ASRs to be higher in mate-guarding gorillas than in chimpanzees. Demography of the Virunga population (20) indicates that, unlike chimpanzees, mortality is not higher in males until the oldest ages. Sexes are not always distinguished in more recent gorilla censuses (21, 22), so we await more data. For orangutans (23), relatively low male mortality is also indicated, but it is difficult to know which males to include among the competitors for mates.

We estimate chimpanzee mating sex ratios using demographic data from the five-study site synthesis of Hill et al. (24) and also include Muller and Wrangham’s (25) more recent (though very small) sample for one of those sites where overall mortality is lowest.

For humans, age structure data come from four well-known hunter-gatherer groups: Howell’s (26, 27) life tables for Dobe !Kung, Hill and Hurtado’s (28) for forest Ache, Hill et al.’s (29) for Hiwi, and Blurton Jones’s (30) for Hadza. We chose these groups because of their reliance on wild foods and the quality of demographic data collected among them. Although contemporary foragers are not living fossils, their mortality regimes make demographic patterns observed among them the best representation of human demography during our evolutionary past.

Table 1 shows that the empirical values computed from chimpanzee sources (ASRs of 0.47 and 0.70; OSRs of 41 and 61) are in general accord with the simulated values without grandmothering. Among the hunter–gatherers (Table 2) ASRs average 1.64, whereas OSRs average 83 (range 69–92), mirroring the equilibrium ASR and OSR values reached in our simulations with grandmothering. Two attributes of these results are especially important. First, every empirical human ASR and OSR is encompassed, or nearly so, in every simulation. Second, in both the real world and simulations with grandmothering the human ratios are distinctly higher than those of the great apes.

Mating Sex Ratios and Mating Strategies
Marlowe and Berbesque (37) pointed to the effects of a strongly male-biased OSR on male competition and marriage resulting from women’s long postfertile life stage. In general, male responses to such a situation should depend on the costs and benefits of available alternatives. Parker (38) was the first to suggest a positive relationship between the relative number of males in a population and the fitness benefits of defending females, or mate guarding. Predictions about the evolution of mate guarding have been elaborated by a number of subsequent
Assuming stationary populations, the mortality curve mirrors the age structure. To model age structures we used probability of survival to each age in the published life tables, summing the calculated number of survivors for males and females to each of the fertile ages, then dividing the sum for each sex by their combined total to get the fraction fertile adults by sex (columns 2 and 3). We included males older than 15 years and females between ages 10 and 45 years (31). Data on chimpanzee birth intervals come from averaging reports in Knott (32).

For OSR we used the formula for nonseasonal breeders derived by Mitani et al. (33), where

\[ OSR = \frac{m \cdot B \cdot 365}{f \cdot \sum_{i=1}^{8} i} \]

In this equation, \( m \) and \( f \) are the fraction of fertile adults that are male and female, respectively; \( B \) is the average birth interval; and 365 is the days per year that males can compete for a paternity. The summation in the denominator is the fecundable days per birth interval for fertile females. It depends on the number of conception risk days per estrous cycle (\( s \)) and the number of cycles per conception (\( n \)). As in the simulations we use observations from humans (34, 35) and fix \( s = 6 \) and \( n = 4 \).

treatments (e.g., refs. 39–44). The hypothesized relationship between biased sex ratios and mate guarding has also been supported by observational and experimental studies conducted across a range of species (45–50).

Although mate guarding is not synonymous with pair bonding, similar tradeoffs are involved. Guarding a mate (or potential mate) is beneficial for members of the more numerous sex when it leads to greater reproductive success (RS) than continually seeking out new mates (38). The logic underlying pair bonding is the same: individuals of the more prevalent sex should guard continuously (i.e., form pairs) when doing so leads to greater RS than the alternatives (44).

Parker and Stuart’s (39) model assumed that males acquire females through either guarding a current mate (G) or searching for another one (S). Clutton-Brock and Parker (51) retained this S vs. G notation, but they posed the alternatives differently (ref. 51, p. 447), defining “time in” the mating pool (S) to include both searching and guarding, with guarding also part of “time out” (G).

Of special relevance here, Kokko and Jennions (17) model has been read by some (59) as undermining OSR as a predictor of mating patterns, warranting focus on ASR instead.

If the tradeoffs for hominid males were the ones assumed in Kokko and Jennions’ (17) basic model, the evolution of increasingly male-biased OSRs would be unstable. Where OSR is male-biased, males take longer to find another mate and so do better to choose an alternative to searching for them. While emphasizing this important effect, the only alternative considered in their basic model is supply parental care, with each additional unit of care (from either sex) earning equivalent marginal gains in offspring survival. Given those assumptions, male bias in the mating pool must decline as males stay out to parent longer.

### Table 1. Demographic parameters for chimpanzees

| Population          | Males age 15+ years | Females age 10-45 years | Birth interval, year | Male paternity, days/year | Females fecundable days per cycle | Cycles to conception | ASR M/F | OSR M/F |
|---------------------|---------------------|-------------------------|----------------------|---------------------------|----------------------------------|----------------------|---------|---------|
| Synthesis of five   | 0.321               | 0.679                   | 5.72                 | 365                       | 6                                | 4                    | 0.47    | 40.13   |
| Kanyawara (25)      | 0.411               | 0.589                   | 5.36                 | 365                       | 6                                | 4                    | 0.70    | 60.70   |

### Table 2. Demographic parameters for human hunter-gatherers

| Population          | Males age 20–65 years | Females age 20–40 years | Birth interval, year | Male paternity, days/year | Female fecundable days per cycle | Cycles to conception | ASR M/F | OSR M/F |
|---------------------|-----------------------|-------------------------|----------------------|---------------------------|----------------------------------|----------------------|---------|---------|
| Dobe Kung (26, 27)  | 0.593                 | 0.407                   | 4.17                 | 365                       | 6                                | 4                    | 1.46    | 92.40   |
| Ache forest (28)    | 0.652                 | 0.348                   | 2.44                 | 365                       | 6                                | 4                    | 1.87    | 69.52   |
| Hwi (29)            | 0.618                 | 0.382                   | 3.70                 | 365                       | 6                                | 4                    | 1.62    | 91.04   |
| Hadza (30)          | 0.616                 | 0.384                   | 3.23                 | 365                       | 6                                | 4                    | 1.60    | 78.80   |

Assuming stationary populations, the mortality curve mirrors the age structure. To model age structures we used probability of survival to each age in the published life tables, summing the calculated number of survivors for men and women to each of the fertile ages, then dividing the sum for each sex by their combined total to get the fraction fertile adults by sex (columns 2 and 3). We included men from 20 and 65 years based on reported age ranges of fertilities from the ethnographers and those reported by Tuljapurkar et al. (36). Women from 20 to 40 years are included based on average ages of first and last birth (4). Since populations are younger when increasing, and the growth rates for the human populations range from 0.26% to 2.5%, our figures overestimate the ASRs by 2–12%.

We computed birth intervals by dividing years of female fertility by the total fertility rate reported in the ethnographic sources. OSR was calculated as in Table 1.
Measuring Human Sex Ratios: ASRs or Elder Advantage?
In our simulations mating sex ratios become increasingly male-biased with a growing fraction of elders as a human-like life history evolves from a great ape-like one. This reflects the approximate doubling of longevity while female fertility maintains its ancestral decline to near zero by 45. Although mortality is generally higher in men than in women (2, 3), the increased number of older men who are not yet frail makes human ASRs inevitably male-biased.

How, then, can some explorations of human mating sex ratios report so many to be female-biased? Using variance in RS reported by Brown et al. (72), Kokko and Jennions (ref. 73, pp. 113–114) used the Fisher condition and the RSs reported to infer ASRs for 18 human populations, estimating them to be about even for the societies classified as monogamous or serially monogamous and significantly female-biased for the groups classified as polygynous. For nine cases Brown et al. (72) classified as polygynous, the mean RS for males averages 31% higher than the female mean—possibly due, as Kokko and Jennions (73) conclude, to female-biased ASRs. But Brown et al. (ref. 72, p. 300) note another possibility: missing men who were less successful in the mating competition.

An earlier study by Ember (74) also found an association between polygyny and female-biased sex ratios using data from the Human Relations Area Files. While high enough male mortality could make ASRs female-biased, Ember’s sex ratios were not ASRs but sex ratios of whole populations—immatures and postfertile women included. Schacht et al. (ref. 59, p. 218) note this difficulty with “which sex ratio.” They review social science research into sex-ratio effects listing 20 studies in their first table; none of them used fertile ages.

Figure 2 in Schacht et al. (ref. 59, p. 217) is the most serious challenge to our characterization of human mating sex ratios as male-biased. It plots results from the authors’ queries to ethnographers about 15 traditional populations. Here, they say, “Sex ratio is determined from the ethnographers’ data on the number of individuals of mating age in their population, hence it approximates ASR.” Eight of these societies—more than half—have female-biased ASRs, a pattern that our model and arguments suggest is “not human.”

At least three factors could lead ethnographers to underestimate sex ratios in the fertile ages. First, physiologically fertile young men may be excluded because local conventions class them as not yet of mating age. Székely et al. (ref. 75, p. 1501), reviewing broader cross-species issues of ASR and OSR associations with breeding systems, define an adult as “an animal (male or female) that is physiologically capable of producing offspring.”

Second, counting unmarried men is difficult if they move more often than others, a common pattern among hunter–gatherers. Such high mobility gave Hawkes et al. (ref. 76, p. 683) an insufficient sample of unmarried Hadza men to compare their time allocation patterns with those of married men. Székely et al. (ref. 75, p. 1501) note similar ascertainment biases with other animals, because “nonbreeding adults (e.g. floaters) often…remain unnoticed.”

A third, closely related possibility is that ethnographers chose not to include unmarried men. For example, polygynous Kipsigis have a notably low ASR. Reporting that Kipsigis men have much higher average RS compared with women (which, given the Fisher condition, would require extremely high male mortality), Borgerhoff Mulder (ref. 77, p. 433) said that her tabulation excluded “poor men, who leave the community to become plantation laborers and marry late in life, if at all.”

Marlowe and Berbesque (37) showed the advantage that elder men can have in ethnographically known hunter–gatherers by plotting the relationship between age of first marriage and the extent of polygyny across 130 hunter–gatherer societies. Their figure 1 (ref. 37, p. 836) shows that men’s age at first marriage
rises from an average near 20 y to more than 30 y as the fraction of polygynously married men rises from few to more than 50%. We tentatively conclude that apparent female bias in some human ASRs may actually index the shape of the male status hierarchy. Exclusion of some men, young and old, from paternities occurs not because they are infertile but because they are outcompeted by other men. As Darwin said in developing his theory of sexual selection, it favors features that “serve only to give one male an advantage over another male, for the less well endowed males, if time were allowed them, would succeed in pairing with the females...” (ref. 78, p. 257).

Unmarried men are not only a part of the ASR, their relationships with married men have long been reported by cultural anthropologists to shape much of community life (71, 79). Estimates of fertile ages for men confront entirely different measurement challenges than do estimates for women. As Vinicius and colleagues (ref. 80, pp. 4–5) concluded, noting variation in paternities attributed to older men across a sample of traditional societies, “Since there is no evidence of widespread male mid- or late-life sterility, variation in late-life reproduction in men must therefore reflect differences in opportunities to reproduce at old age.” This and the likelihood of undercounting men less successful in the competition provide grounds for skepticism about the reported female biases. Age structures themselves support, if indirectly, our characterization of human mating sex ratios as characteristically male-biased.

Conclusion

Our hypothesis is that human pair bonds evolved with increasing payoffs for mate guarding, which resulted from the evolution of our grandmothering life history. This mate-guarding hypothesis is an alternative to long-favored arguments that pairing evolved in our lineage as a consequence of the benefits of cooperative parenting (e.g., refs. 81–86). We are far from the first to connect human pair bonds with mate guarding. Wilson and Daly (87) assembled a wide range of cross-cultural evidence and argument nearly 25 y ago. Nor is the proposal that human pair bonds more likely began with mate guarding than with paternal care a novel suggestion (e.g., refs. 88–90). As Chapais (90) notes, the group living patterns of humans differ from the territorial monogamy of other pair-living mammals, but where parental cooperation did evolve in mammals or more narrowly primates, it followed the prior establishment of pair bonds (91–93).

We have focused on changes in the mating sex ratio that accompany the evolution of our grandmothering life history and raise the net benefits to males for mate guarding and pair bonding—patterns that distinguish humans from our closest living relatives. This emphasizes an aspect of human pair bonds ignored by models that assume the only reproductive options are parental care or competition for another mate. An early attempt to distinguish mate guarding as an allocation of reproductive effort that trades off with other kinds of mating competition—as well as with offspring care—found unexpectedly wide conditions under which guarding displaced the other options even without varying mating sex ratios (41). Although all observed human societies feature pair bonds, their character and stability is notably variable. Across our hunter–gatherer sample pair-bond stability is greater where mating sex ratios are more male-biased (94). Guttentag and Secord (95) showed the same correlation over time and across socioeconomic classes in state societies. Ethnographic explorations of the tension between a man’s conjugal bonds and his alliances with other men have a rich history in anthropology (71, 96–98). The argument here begins to link that tension to the evolution of our grandmothering life history.

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1. Trivers R (1972) Parental investment and sexual selection. Sexual Selection and the Descent of Man 1871–1971, ed Campbell B (Aldine, Chicago), pp 139–179.
2. Kruger DJ, Nesse RM (2006) An evolutionary life-history framework for understanding sex differences in human mortality rates. Hum Nat 17(1):74–91.
3. Oksuzyan A, Joel K, Vaupel JW, Christensen K (2008) Men: Good health and high mortality. Sex differences in health and aging. Aging Clin Exp Res 20(2):91–102.
4. Robson SL, van Schaik CP, Hawkes K (2006) The derived features of human life history. The Evolution of Human Life History, eds Hawkes K, Payne R (SAR, Santa Fe, NM), pp 17–44.
5. Robbins AM, Robbins MM, Gerald-Steklis N, Steklis HD (2006) Age-related patterns of infertility. J Evol Biol 19(6):1235–1245.
6. Hawkes K, O’Connell JF, Blanton Jones NG, Alvarez H, Charnov EL (1998) Grandmothering, menopause, and the evolution of human life histories. Proc Nat Acad Sci USA 95(3):1336–1339.
7. O’Connell JF, Hawkes K, Blanton Jones NG (1999) Grandmothering and the evolution of Homo erectus. J Hum Evol 36(2):461–485.
8. Hawkes K (2003) Grandmothers and the evolution of human longevity. Am J Hum Biol 15(3):380–400.
9. Kim PS, Coxworth JE, Hawkes K (2012) Increased longevity evolves from grandmothering. Proc Biol Sci 279(1749):4880–4884.
10. Coxworth JE, Hawkes K (2014) Primate sociality to human cooperation. Why us and not them? Hum Nat 25(1):28–48.
11. Kim PS, McQueen JS, Coxworth JE, Hawkes K (2012) Grandmothering drives the evolution of longevity in a probabilistic model. J Theor Biol 353:84–94.
12. Hawkes K, Coxworth JE (2013) Grandmothers and the evolution of human longevity: A review of findings and future directions. Evol Anthropol 22(4):294–302.
13. Kokko H, Jennings MD (2008) Parental investment, sex selection and sex ratios. J Evol Biol 21(4):919–948.
14. Parker GA, Baker RR, Smith VGF (1972) The origin and evolution of gamete dimorphism and the male-female phenomenon. J Theor Biol 36(3):529–553.
15. Emelin ST, Oring LW (1977) Ecology, sexual selection, and the evolution of mating systems. Science 197(4248):215–223.
16. Gerald CN (1995) Demography of the Virunga mountain gorilla (Gorilla gorilla beringei). Master’s thesis (Princeton Univ, Princeton).
17. Robbins MM, Robbins AM (2004) Simulation of the population dynamics and social structure of the Virunga mountain gorillas. Am J Primatol 63(6):201–223.
18. McNeilage A, et al. (2006) Census of the mountain gorilla Gorilla beringei beringei population in Bwindi Impenetrable National Park, Uganda. Orang 40(4):419–427.
19. Wich SA, et al. (2006) Life history of wild Sumatran orangutans (Pongo abelii). J Hum Evol 47(5):385–398.
20. Hill K, et al. (2001) Mortality rates among wild chimpanzees. J Hum Evol 40(5):437–450.
21. Muller MN, Wrangham RW (2014) Mortality rates among Kanyawara chimpanzees. J Hum Evol 66:107–114.
22. Coale AJ, Demeny P (1966) Regional Model Life Tables and Stable Populations (Princeton Univ Press, Princeton).
23. Hill K, Hurtado AM (1996) Ache Life History: The Ecology and Demography of a Foraging People (Al dia de Gruyter, New York).
24. Hill K, Hurtado AM, Walker RS (2007) High adult mortality among !Kung hunter-gatherers: Implications for human evolution. J Hum Evol 52(4):443–454.
25. Blanton Jones NG (2015) Demography and Evolutionary Ecology of Hadza Hunter-Gatherers (Cambridge Univ Press, Cambridge, MA).
26. Emery Thompson M, et al. (2007) Aging and fertility patterns in wild chimpanzees provide insights into the evolution of menopause. Curr Biol 17(24):2150–2156.
27. Knoott C (2001) Female reproductive ecology of the apes: Implications for human evolution. Reproductive Ecology and Human Evolution, ed Ellison PT (Al dia de Gruyter, New York), pp 429–463.
28. Mitani JC, Gros-Louis J, Richards AF (1996) Sexual dimorphism, the operational sex ratio, and the intensity of male competition in polygynous primates. Am Nat 147:966–980.
29. Wilcox AJ, Weinberg CR, Baird DD (1995) Timing of sexual intercourse in relation to ovulation. Effects on the probability of conception, survival of the pregnancy, and sex of the baby. N Engl J Med 333(23):1517–1521.
30. Gensburg G, Godderitz D, Ehrhardt E, Frank-Herrmann P, Freundl G (2003) Time to pregnancy: Results of the German prospective study and impact on the management of infertility. Hum Reprod 18(9):1959–1966.
