No evidence of short-term exchange of meat for sex among chimpanzees

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Abstract

The meat-for-sex hypothesis posits that male chimpanzees (Pan troglodytes) trade meat with estrous females in exchange for short-term mating access. This notion is widely cited in the anthropological literature and has been used to construct scenarios about human evolution. Here we review the theoretical and empirical basis for the meat-for-sex hypothesis. We argue that chimpanzee behavioral ecology does not favor the evolution of such exchanges because 1) female chimpanzees show low mate selectivity and require little or no material incentive to mate, violating existing models of commodity exchange; and 2) meat-for-sex exchanges are unlikely to provide reproductive benefits to either partner. We also present new analyses of 28 years of data from two East African chimpanzee study sites (Gombe National Park, Tanzania; Kanyawara, Kibale National Park, Uganda) and discuss the results of previously published studies. In at least three chimpanzee communities, 1) the presence of sexually receptive females did not increase hunting probability, 2) males did not share preferentially with sexually receptive females, and 3) sharing with females did not increase a male's short-term mating success. We acknowledge that systematic meat sharing by male chimpanzees in expectation of, or in return for, immediate copulations might be discovered in future studies. However, current data indicate that such exchanges are so rare, and so different in nature from exchanges among humans, that with respect to chimpanzees, sexual bartering in humans should be regarded as a derived trait with no known antecedents in the behavior of wild chimpanzees.

Introduction

In this paper we review published articles and provide new data on the influence of female sexual state on hunting and meat sharing among chimpanzees (Pan troglodytes) to review the ‘meat-for-sex’ hypothesis. The meat-for-sex hypothesis proposes that male chimpanzees share meat with females in exchange for short-term mating access. Specifically, the meat-for-sex hypothesis states that, “... the hunting performance of chimpanzees may be under sexual selection for capture and/or control of meat as a way to obtain additional copulations from females with estrous swellings in the hunting party” (Stanford, 1996: 101). Meat-for-sex exchanges among chimpanzees are often treated as a well-established, species-typical phenomenon (e.g., Radetsky, 1995; Bird, 1999; Hawkes and Bird, 2002; Shlain, 2003; Bunn, 2007; Lovejoy, 2009), and they provide the basis for various speculations about the evolution of human sexual behavior (e.g., Lovejoy, 2009).

Among humans the trading of meat for sex can be explicit. Siriono wives who are more sexually active with their husbands reportedly get more to eat (Holmberg, 1969: 126), while men use meat to obtain extramarital sex (Holmberg, 1969: 166). Among the Mehinaku, women are said to, “use their sexuality to secure food and support in exchange for intercourse” (Gregor, 1985: 33), and men secretly give fish to their lovers as a symbol of their productivity and sexuality (Gregor, 1985: 75). The Kulina practice a public ritual called ‘order to get meat’ (Pollock, 2002: 52) which begins when each woman raps on a man’s house at dawn with a stick. If that man hunts successfully that day, she will have sex with him. After the men hunt, women form a semi-circle around them, requesting meat while singing erotically provocative songs. Men hurl the meat in a pile, and the women cook and eat it before retiring with the selected partner.

Clear examples of food-for-sex exchanges also occur in many non-primate species, especially insects (Thornhill, 1984; Noé and Hammerstein, 1994; Arnvist and Nilsson, 2000; Fedorka and Mousseau, 2002). However, the claim that chimpanzees trade meat for copulations is more important for human evolution because it raises the question of whether human meat-for-sex...
exchanges can be traced to a pre-human ancestry. In this paper we review the full theoretical and empirical base underlying the meat-for-sex hypothesis in chimpanzees and present new data from Gombe National Park, Tanzania and the Kanyawara community in Kibale National Park, Uganda. We find that the behavioral observations which provided the initial impulse for the meat-for-sex hypothesis have never been replicated and can be explained by alternative mechanisms. In four study communities (Gombe, Kanyawara, Ngogo [Kibale National Park], and Tai National Park, Côte d’Ivoire), controlled examinations of multiple predictions of the hypothesis find no evidence that chimpanzees systematically exchange meat for copulations on a short-term basis.

Although short-term exchanges of meat for sex are not supported, Gomes and Boesch (2009) reported that there was a long-term correlation across male-female dyads between meat sharing and copulations at Tai. Specifically, during a 22-month study period, copulations were more likely to occur in dyads that shared meat at least once. Gomes and Boesch (2009) interpreted this result as indicating that meat was exchanged for sex. However, despite careful analysis they failed to find any evidence that meat sharing was correlated with copulations over the short term. Their result is therefore mysterious, since it implies that the motivation of estrous females to copulate was not affected in the short term by receiving meat, whereas it was affected in the long term. In this paper our data do not address the possibility that chimpanzees exchange meat for sex over a period of days, weeks, or months. Instead, we are concerned only with the question of whether they perform these exchanges within the immediate context of meat-eating.

Chimpanzee natural history and meat sharing

Chimpanzees typically live in communities of approximately 40–50 individuals, although some communities can reach ~150 (e.g., Ngogo: Mitani and Watts, 1999). Within a community, fission-fusion grouping generates ‘parties’ of varying size, composition, and temporal stability (Nishida, 1968; Wrangham and Smuts, 1980; Goodall, 1986). Thus, while male and female members of a community maintain long-term social relationships, the strength of these relationships varies as does the frequency of contact between any two individuals (e.g., Gilby and Wrangham, 2008). Reproductive rates are very slow, with a mean female inter-birth interval of 5–7 years (data from 6 sites summarized by Emery Thompson et al. [2007a]). Cycling females exhibit a large anogenital estrus swelling for approximately 10–12 days per 35-day cycle (Walls, 1997). With very few exceptions, females mate only when they are maximally swollen (Goodall, 1986; Walls, 1997). We refer to females with a maximal sexual swelling as ‘estrus’ females. Females usually experience multiple cycles and mate with many males before each conception (Tutin, 1979; Wrangham, 2002; Deschner and Boesch, 2007; Watts, 2007; Emery Thompson and Wrangham, 2008a).

Chimpanzees primarily eat ripe fruit, leaves, and other plant materials. However, they also prey upon small- and medium-sized vertebrates. Where available, the most common prey is the red colobus monkey (Procolobus spp.; Uehara, 1997). Meat consumption varies considerably across populations. At one extreme, the Ngogo chimpanzees killed and consumed at least 292 prey items during a 34-month study period (~103 prey per year; Watts and Mitani, 2002b). At the other extreme, chimpanzees in the Budongo Forest, Uganda were observed to eat meat only 17 times in more than 10 years (Newton-Fisher et al., 2002; Reynolds, 2005). Even where hunting occurs with some frequency, some individuals rarely if ever gain access to meat (Gombe: Goodall, 1986; Tai: Boesch and Boesch-Achermann, 2000; Ngogo: Mitani and Watts, 1999). These observations suggest that meat is not an essential component of chimpanzee diet (Boesch and Boesch-Achermann, 2000). However, prey animals offer a concentrated source of protein, fat, and micronutrients (Teleki, 1973; Takahata et al., 1984; Goodall, 1986; Boesch, 1994; Stanford, 1996; Mitani and Watts, 2001; Milton, 2003a; b; Tennie et al., 2009) and are clearly a highly preferred food item for males and females alike.

While females occasionally hunt (Goodall, 1986; Boesch, 1994; Watts and Mitani, 2002b), males are responsible for most kills and readily seize fresh prey caught by others, and are therefore most often in a position to share meat. When a male obtains prey, several adult male and female chimpanzees typically surround him, each begging for a piece of the carcass (Teleki, 1973; Wrangham, 1975; Takahata et al., 1984; Goodall, 1986; Boesch and Boesch-Achermann, 2000; Gilby, 2006). Begging behavior includes pulling on the carcass, attempting to co-feed, and reaching toward the possessor’s mouth (Teleki, 1973; Goodall, 1986; Gilby, 2006). Possessors often attempt to reduce such harassment by sneaking away, sitting on terminal branches, and/or physically retaliating against particularly persistent beggars (Wrangham, 1975; Goodall, 1986; Gilby, 2004).

Despite these attempts to avoid beggars, meat possessors at Gombe and Ngogo share in approximately 50% of all begging bouts (Mitani and Watts, 2001; Watts and Mitani, 2002a; Gilby, 2006). Most often, possessors share ‘passively’ by allowing an individual to take pieces or feed from the carcass (Boesch and Boesch, 1989; Gilby, 2006). However, sharing may also be ‘active,’ when the possessor facilitates transfer by purposefully dividing the carcass, handing a piece to a beggar, or depositing a mouthful of meat into a waiting hand (Teleki, 1973; Goodall, 1986; Boesch and Boesch, 1989; Gilby, 2006). At Gombe, approximately 20% of sharing events are active (Gilby, unpublished data).

Study sites and data collection

Our new data analysis comes from Gombe and Kanyawara two of the longest continuously studied free-ranging chimpanzee communities. Systematic data collection has been ongoing since 1974 at Gombe (Goodall, 1986) and 1987 at Kanyawara (Wrangham et al., 1992). For the present study, we analyzed data collected from 1995 to 2005 at Gombe, and from 1990 to 2006 at Kanyawara. During these study periods, teams of observers, which included local field assistants and experienced scientists, followed chimpanzees from dawn until dusk on most days at both sites. At Kanyawara, the team followed and recorded data on a party of chimpanzees, and at Gombe, they followed a focal adult. At both sites, one observer recorded party composition, feeding behavior, and location at 15-minute intervals. Another observer made detailed notes ad libitum of all social behavior, including aggression, mating, and hunting. After a successful hunt, the observers wrote detailed accounts of all meat eating, including the identity of meat possessors and beggars and the occurrence of sharing and mating. At Gombe, the observers concentrated their sampling efforts on the focal individual.

Confidence in the accuracy of these data comes from the presence of several observers dedicated to obtaining detailed descriptions of all behavior through immediate collaborative discussion. At Kanyawara, observers average over 95% in inter-observer reliability tests (Kibale Chimpanzee Project, unpublished data). For more detail on long-term data collection at Kanyawara, see Gilby et al. (2008), Emery Thompson et al. (2007b), and Muller et al. (2007). Data collection at Gombe was described by Goodall (1986), Williams et al. (2002), and Gilby et al. (2006). In this study, we analyzed data from 260 successful hunts that occurred during 11 years at Gombe, and 126 successful hunts that
occurred during 17 years at Kanyawara. For simplicity, we provide the specific details, including sample sizes and analytical criteria, when reporting each result.

In addition to long-term data, we also used targeted data on meat sharing collected at Gombe by I. Gilby over 12 months of field work during peak hunting periods between 1999 and 2002. Gilby videotaped 65 meat-eating bouts (defined as the total time an individual possessed meat following a successful hunt) by five focal males, and subsequently extracted detailed begging, sharing, and mating data. Further details are described by Gilby (2006). In the present article, we provide new analyses of these data (the “Gilby data set” hereafter). We used SAS 9.1 (SAS Institute, Cary, North Carolina) for all analyses, and set statistical significance at alpha = 0.05.

We take the following approach. First, we discuss two critical assumptions that must be satisfied for short-term meat-for-sex exchanges to evolve. We argue that as a consequence of certain fundamental aspects of chimpanzee behavioral ecology, these assumptions are unrealistic for this species. Therefore, we conclude that chimpanzees should not be expected to trade meat for sex. We then refer to published data and use our own new analyses to test four predictions generated by the meat-for-sex hypothesis.

**Assumptions of the meat-for-sex hypothesis**

The meat-for-sex hypothesis hinges upon the assumptions that 1) females need incentive to overcome a reluctance to mate, and 2) the change in female behavior following meat transfer can enhance the reproductive success of the male donor. While previous studies of meat sharing among chimpanzees are rarely this explicit, these assumptions are fundamental to the evolutionary hypotheses that are most often invoked to explain such exchanges. Reciprocal Altruism theory (Trivers, 1971) posits that individuals take turns engaging in costly acts that benefit the other, resulting in a net benefit for both participants over time. Therefore, if we view meat-for-sex as a reciprocal exchange, we must assume that for the female, mating with certain males incurs a greater cost than withholding sex, but that this cost is offset by the benefits of obtaining meat. Biological markets theory (Noë et al., 1991; Noë and Hammerstein, 1995) is based upon similar principles, noting specifically that partners exchange commodities over which each has differential control. However, as we demonstrate below, both theory and available empirical evidence suggest that under most circumstances female chimpanzees pay low costs by mating and even gain benefits through promiscuity. Furthermore, a number of characteristics of the mating system of chimpanzees make it unlikely that meat-for-sex exchanges would significantly alter male or female reproductive success. This obviates the evolutionary significance of meat sharing as a mating strategy for chimpanzees, even if it occasionally occurs. Thus, with specific reference to the evolution of mating markets and “nuptial gifts,” we argue below that current models entail assumptions and predictions that are unrealistic for chimpanzees.

We begin by challenging the notion that female chimpanzees should be reluctant to mate in general. Females in species that form multi-male, multi-female social groups can benefit in several ways by mating promiscuously. Mating with many males may 1) confuse paternity, and thus discourage infanticide (Hrdy, 1979; van Schaik et al., 2000); 2) promote sperm competition, thereby increasing the chances of acquiring high quality genes (Clutton-Brock and Harvey, 1976); and 3) ensure fertilization (Milton, 1985; Small, 1986). Indeed, in chimpanzees, a single conception typically entails hundreds of copulations with most males in the community (Tutin, 1979; Hasegawa and Hiraïwa-Hasegawa, 1990; Wrangham, 2002). Females sometimes mate extensively during pregnancy (Wallis, 1982; Kibale Chimpanzee Project, unpublished data). Furthermore, mating patterns indicate that females engage in a high number of copulations in order to mate with more males rather than to mate more frequently with a particular male (Wrangham, 2002; Watts, 2007). Consistent with the argument that mating is a low-cost activity for females (thereby violating assumption #1), rates of female resistance to male copulation solicitations are low. Goodall (1986) notes that at Gombe, females failed to approach for copulation (within one minute) after only 61 (4%) of 1475 male solicitations. Forty-one (67%) of these refusals involved a single sterile female or the male’s maternal sister. The pattern at Kanyawara is similar. We identified all male copulation attempts (defined as copulatory mounts or near-monts following male approach or solicitation) in the long-term data between 1996 and 2006. During this period, only 55 (3%) of 1894 male copulation attempts failed due to female resistance (rather than male interference), usually because the female ran away screaming or was aggressive to the male. Even in these rare events, it was often noted that a female fled a copulation in response to threats from other males, rather than in direct resistance to her partner. At Tai, resistance has been defined more broadly, as when a female ignores or moves away from a male in her first response to his solicitation, regardless of whether she subsequently mates with him. Females were resistant to only 265 of 938 (28%) male copulation attempts at Tai (Stumpf and Boesch, 2006; cf. 8%, Boesch and Boesch-Achermann, 2000). This was only marginally higher than the 21% of female solicitations that were resisted by males (Stumpf and Boesch, 2006).

Thus, neither theoretical expectations nor empirical evidence support the assumption that female chimpanzees are generally reluctant to mate. However, it has been suggested that females display subtle preferences for certain males when conception probability is highest (Tai: Stumpf and Boesch, 2005; Kanyawara: Pieta, 2008; Mahale Mountains National Park, Tanzania: Matsumoto-Oda, 1999), raising the possibility that meat sharing could skew these preferences. Under existing models of mating markets (e.g., Noë and Hammerstein, 1994, 1995; Gwynne, 2008), two additional conditions must be met for this to be a feasible argument for explaining the evolution of meat sharing in chimpanzees. First, female mate selectivity must be an important predictor of male reproductive success, as in insects that provide “nuptial gifts” (Gwynne, 2008). In other words, female choice must influence male mating outcomes. If females cannot achieve their mating preferences, then meat will not affect the outcome of any conflict of interest. Second, given the high cost of reproduction for females, preference for a male that offers meat should benefit the female in one of three ways: (a) the energetic benefit should help subsidize the costs of reproduction; (b) the willingness to share meat should be an indication of a male’s future investment in offspring care; or (c) the ability of the male to acquire meat should be indicative of his genetic quality. However, male chimpanzees neither provision infants nor provide obvious forms of paternal care. As noted previously, any caloric benefits gained from shared meat seem to be small, although meat may provide micronutrients that could positively affect a female’s reproductive success (Tennie et al., 2009). It remains possible that meat sharing provides an advertisement of male fitness. However, as we argue below, the preponderance of various male competitive strategies suggest a limited role for female selectivity in determining male reproductive success.

Three alternative male strategies, which are strongly supported as determinants of reproductive success, suggest that female mate choice plays a minor role in chimpanzee communities, although this may vary across populations. First, as in many multi-male species, male chimpanzees fight intensely over mating opportunities and powerful males often use possessive tactics to monopolize
patterns occur at Gombe, where aggression towards a particular female predicted his relative (POP), when conception probability was highest, and a male’s Kanyawara, such aggression increased in the peri-ovulatory period Mahale (attractive or fecund females (Mahale: Thompson and Wrangham, 2008b increases when females are most likely to conceive (2006 Emery Thompson and Wrangham, 2008b: the majority of copulations (Gombe: Goodall, 1986; Kanyawara: Emery Thompson and Wrangham, 2008b; Tai: Stumpf and Boesch, 2006), and the intensity of male mating effort and competition increases when females are most likely to conceive (Emery Thompson and Wrangham, 2008b). Thus, high-ranking males achieve the highest mating success, particularly with the most attractive or fecund females (Mahale: Nishida, 1997; Budongo: Newton-Fisher, 2004; Kanyawara: Muller et al., 2006; Duffy et al., 2007). Additionally, paternity results from Gombe (Constable et al., 2001; Wroblewski et al., 2009), Tai (Boesch et al., 2006), Mahale (Inoue et al., 2008), and Budongo (Newton-Fisher et al., in press) indicate that high rank and the use of competitive mating strategies is a strong, although not perfect, predictor of male reproductive success in chimpanzees.

Second, while it is extremely rare for male chimpanzees to force a female to copulate (0.001% of matings at Tai [Stumpf and Boesch, 2006]; 0.002% at Gombe, [Goodall, 1986]), males, who are approximately 25% heavier than females (Pusey et al., 2005), frequently use aggression against females (Muller, 2002). At Kanyawara, such aggression increased in the peri-ovulatory period (POP), when conception probability was highest, and a male’s aggressive behavior towards a particular female predicted his relative mating success with that female (Muller et al., 2007). Similar patterns occur at Gombe, where Goodall (1986) notes that, ‘By attacking his female...[a] possessive male not only stops her from mating with his rival, but also delivers a warning that she should avoid the sexual advances of other males.’ In fact, at Kanyawara, conditioning aggression by high-ranking males is sufficient to explain inter-individual and temporal variation in female solicitation patterns (Muller et al., 2009, in review) that are often used as measures of preference (Stumpf and Boesch, 2005, 2006; Pieta, 2008). It is feasible, however, that the impact of coercion on the expression of female preferences varies in chimpanzees, as suggested by West African chimpanzee (P. verus) females at Tai, who are reported to exert choice more effectively (Stumpf and Boesch, 2005, 2006).

In sum, based on the available evidence, it is likely that female mate preferences have a weak overall influence on male reproductive outcomes in chimpanzees in comparison with the forces of male aggressive competition, sexual coercion, and sperm competition, suggesting that sharing meat with the occasional reluctant female would do little to increase a male’s reproductive success. This conclusion is reinforced by the fact that male meat possessors are usually high ranking (e.g., Gilby, 2006). These males already hold a significant advantage in securing matings by virtue of aggressive competition with other males and their ability to effectively use mate guarding and coercive aggression to increase their relative mating success. The competitive success of high-ranking male chimpanzees also suggests high genetic quality, making it likely that they are desirable mates regardless of any sharing behavior (Tutin, 1979; Goodall, 1986; Matsumoto-Oda, 1999; Stumpf and Boesch, 2005).

Meat-for-sex exchanges are therefore generally implausible among chimpanzees. By contrast they have obvious potential significance in insects and humans because females in those species restrict their copulations to one or a few males and obtain large nutritional benefits from substantial portions of food provided by males. Nevertheless, when low- and mid-ranking male chimpanzees do control shareable portions of meat (Goodall, 1986; Gilby, 2006), such as when multiple monkeys are captured, or when high-ranking males have eaten their fill, they could be in a position to trade meat for sex as an alternative strategy to competition. We therefore address the empirical basis for meat-for-sex exchanges among chimpanzees.

Predictions of the meat-for-sex hypothesis

Figure 1. Probability of hunting vs. number of adult male and estrous female chimpanzees in a party that encounters red colobus monkeys. From Gilby et al. (2006). Regression lines are from a logistic regression of 25 years of data from Gombe. Error bars represent 95% confidence intervals. At all party sizes, the presence of an estrous female lowered the odds of at least one male hunting.
to choose between hunting and mating. Males frequently guard estrous females to prevent them from mating with other males. Solo mate guarding is less likely to succeed in a large party (Watts, 1998), and male chimpanzees cannot simultaneously mate guard and hunt. Therefore even if males are more interested in hunting when estrous females are present (Stanford et al., 1994), they might forgo the opportunity to hunt if doing so would compromise their ability to monopolize a sexually receptive female.

The same analysis has been conducted at two other sites. Gilby et al. (2008) found a similar trend at Kanyawara, suggesting that hunting was less likely when fully estrous females were present (p = 0.08; 610 encounters, 99 hunts). Mitani and Watts (2001) showed that the presence of swollen females had no effect on male hunting probability at Ngogo (164 encounters, 61 hunts). Therefore, at the three sites where Prediction 1 has been tested with data on hunting behavior in the presence of fully estrous females, it has been rejected (Table 1).

Prediction 2: males share preferentially with estrous females

Even though sexual interest is not known to drive hunting decisions at Gombe, Ngogo, and Kanyawara, it could still influence the motivation to share once meat is acquired. If meat were being exchanged for sex, one would expect preferential sharing with estrous females. Teleki's (1973) study of meat eating among the Gombe chimpanzees is frequently cited as providing support for this prediction. Teleki (1973) stated that estrous females that 'took and requested' meat from male possessors were more likely to obtain meat (91/132 ‘interactions’ = 69% success) than anestrous females (42/104 = 40% success). Critically, however, neither statistical tests nor data on individuals were reported (Teleki, 1973: 162–3). Furthermore, Teleki’s (1973) report does not differentiate between begging (‘requesting’) and sharing (‘taking’), and it is therefore unclear whether estrous females were actually more likely to obtain meat than anestrous females.

At Tai, there was a tendency for males to share meat more frequently with estrous females than with anestrous females after controlling for their proportional representation in the party (p = 0.06, n = 5 males; Gomes and Boesch, 2009), and at Ngogo, males shared with estrous females more often than expected by chance (Watts and Mitani, 2002a). However, neither of these analyses reported the relative frequency with which females approached male meat possessors to beg for meat, nor did they take differences in begging behavior into account. Plausibly, therefore, estrous females were more likely to obtain meat because they were more likely to approach meat possessors (Goodall, 1986). Indeed, ‘...estrous females stay closer to adult males during predation and are more persistent in their efforts to obtain meat’ (Teleki, 1973: 163). The strength of a female's interest in obtaining meat is thus a vital component of any test of the prediction that males preferentially share with estrous females.

To illustrate this important point, when Watts and Mitani (2002a) analyzed their Ngogo data using each begging bout as the unit of analysis, estrous and anestrous females obtained meat at equal rates. Specifically, estrous female beggars obtained meat in 45% of 82 begging bouts; anestrous female beggars in 48% of 139 begging bouts (Watts and Mitani, 2002a). Similarly, at Gombe, there was no statistically significant association between female reproductive state and the probability of active sharing, passive sharing, or the amount of meat a female obtained (n = 217 begging bouts, 20 females, 8 adult males; Gilby, 2006).

In this study, we identified 86 instances in 260 successful hunts at Kanyawara in which an adult female was observed in proximity to an unrelated adult male (neither son nor maternal brother) who was eating meat. We found that 5 different estrous females obtained meat in 7 (41.2%) of 17 of these occasions (from 4 different males), while 11 anestrous females had 43.5% success (30/69 with 11 different males). This difference was not statistically significant (GEE logistic regression, $\chi^2_1 = 0.03$, p = 0.86, repeated measures = possession/beggar), further corroborating the results from Gombe and Ngogo.

Table 1

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<thead>
<tr>
<th>Site</th>
<th>Study details</th>
<th>Predictions</th>
<th>Level of analysis</th>
<th>Study</th>
<th>Predictions 2: Preferential sharing with estrous females</th>
<th>Prediction 3: Share with particularly fecund females</th>
<th>Prediction 4: Sharing correlated with male short-term mating success</th>
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$^a$ Blanks indicate that a given prediction was not tested. Each study is discussed in detail in the text. 'Sampling unit' indicates the smallest unit of analysis in a given study. Analyses focusing on events (e.g., bouts or encounters) are generally considered to be most informative. 'Level of analysis' indicates whether data analysis was conducted at the focal, dyad, or group level. Group-level analyses often do not allow for appropriate tests of the predictions. There is considerable evidence against short-term meat-for-sex exchanges.

$^b$ In the Stanford et al. (1994) study, 'hunts' included instances of mere hunting interest (without actual pursuit), and females with partial (3/4) sexual swellings were classified as ‘estrous.’ See text for details.
Thus the only analyses that have taken into account the rate at which females approached male meat possessors reach the same conclusion: male chimpanzees do not take the sexual state of female beggars into account when making sharing decisions (Table 1). Whether there are any circumstances under which males do respond to a female’s sexual state by being more generous with meat is unknown.

Prediction 3: males share preferentially with particularly fecund females

There is growing evidence that chimpanzee mating behavior is sensitive to a female’s probability of conception. Most notably, males solicit more copulations and invest more in mate guarding, intersexual competition, and sexual coercion when a female is most likely to conceive (Mueller et al., 2007; Emery Thompson and Wrangham, 2008b). High-ranking males are typically responsible for the majority of a female’s copulations when conception risk is highest, such as during the peri-ovulatory period (POP; Nishida, 1997; Emery Thompson and Wrangham, 2008b). Males at Kanyawara also prefer older, parous females as mating partners (Mueller et al., 2006), and direct more sexual coercion toward them (Mueller et al., 2007). As a result, it is possible that by simply classifying females as “estrous” or “anestrous”, hence including many ‘low-fecundity’ females in our sharing analyses, we masked an important effect. Therefore, we tested whether males preferentially shared with ‘high-fecundity’ females.

Over 16 years at Kanyawara, during which 520 meat-eating bouts were observed, an estrous female approached an unrelated male meat possessor on only 17 occasions. Estrous parous females obtained meat in 5/8 of these begging bouts, all with the same male. Estrous nulliparous females obtained meat in 2/9 begging bouts with 5 different males. These data show no significant difference (GEE logistic regression, \( \chi^2_1 = 2.70, p = 0.10 \), repeated measure = possessor/beggar). The small sample size serves to highlight the rarity of meat-for-sex opportunities at Kanyawara.

Our new analyses of Gombe data show that estrous females with known parity begged from focal males on 65 occasions (combining long-term data and the Gilby data set). Parous females in estrous (\( n = 9 \)) obtained meat in 72% of 44 bouts, while nulliparous females in estrous (\( n = 10 \)) obtained meat in 43% of 21 bouts, a difference that approached statistical significance (GEE logistic regression, \( \chi^2_1 = 3.06, p = 0.08 \), repeated measure = possessor/beggar; Fig. 2a).

To account for the possibility that this trend could be independent of sexual state, we examined whether parity also affected sharing with anestrous females. This was indeed the case. At Gombe, anestrous parous beggars were significantly more likely to obtain meat (60% of 62 begging bouts) than anestrous nulliparous beggars (30% of 7 begging bouts, GEE logistic regression, \( \chi^2_1 = 4.41, p = 0.04 \), repeated measure = possessor/beggar; Fig. 2b). The same trend existed at Kanyawara but was not statistically significant (GEE logistic regression, \( \chi^2_1 = 2.99, p = 0.08 \), repeated measure = possessor/beggar). One likely explanation for this pattern is that parous females (which tend to be older and higher-ranking [Kahlenberg et al., 2008]) may beg for meat more persistently than nulliparous females (regardless of sexual state) thus increasing their chances of obtaining meat.

Due to the small number of begging bouts (\( n = 5 \)) we were unable to test whether males at Kanyawara preferentially shared with females in POP. At Gombe however, we identified 70 instances in the long-term and Gilby data sets in which estrous females begged from focal males. In 51 of these bouts there were sufficient data to determine with high confidence whether the female was in POP (between 2 and 7 days prior to detumescence: Deschner et al., 2003; Emery Thompson, 2005; Emery Thompson and Wrangham, 2008b). Females in POP were equally likely to obtain meat as estrous females outside of POP (GEE logistic regression, \( \chi^2_1 = 0.20, p = 0.65 \), repeated measure = possessor/beggar).

In sum, we found no evidence that males at Kanyawara and Gombe were more likely to share meat with estrous females with a high probability of conception (Table 1).

Prediction 4: sharing meat is correlated with a male’s short-term mating success

The meat-for-sex hypothesis predicts that sharing with a sexually receptive female should be closely associated with copulation (Gilby, 2006), and, in turn, mating rates during or closely following meat-eating bouts should be higher when sharing occurred than when it did not (Mitani and Watts, 2001; Gilby, 2006). We cannot test this prediction with much statistical power since the rate of copulations during meat eating is low. At Gombe, only 26 (0.6%) of 4401 copulations recorded between 1995 and 2005 occurred during a meat-eating bout. Likewise at Kanyawara, only 6 matings (0.1%) of the 4576 that were observed between 1996 and 2006 occurred when the copulating male possessed meat. However, the available data do not support the hypothesis.

Stanford (1998) describes 5 instances in which a male at Gombe withheld meat from a female until she mated with him. Additionally, mating occurred after 33% of successful hunts at which swollen females were present (Stanford, 1998). However, there was no evidence that the copulation rate was higher than expected. Furthermore, it was not reported whether the females who mated were those who received meat.

More recent data are contrary to the prediction that mating frequency is positively affected by sharing. At Ngogo, mating occurred in 43% of 37 sharing episodes with estrous females, a probability that was not statistically different from 0.5 (Watts and Mitani, 2002a). At Gombe, among begging bouts by estrous females, mating occurred in 35% of 23 bouts in which sharing occurred vs. 15% of 13 bouts without sharing (Gilby, 2006). While these data suggest that there might be a positive short-term effect of sharing on mating, the difference was not statistically significant after controlling for repeated observations of dyads (\( p = 0.2 \)). To examine this question further, we identified all instances (between 1995 and 2005) at Gombe (\( n = 24 \)) when a focal estrous female received meat from a male and asked whether the male was more likely to mate with that female after sharing occurred than before. We limited our analysis to all instances in which both the female and the male were present in the same party for 3 hours before and 3 hours after sharing. Contrary to the prediction, the male meat donor was equally likely to mate with the female within the 3-hour period after sharing (\( n = 4 \)) as he was in the 3-hour period before sharing (\( n = 3 \); Sign test, \( p = 1 \)). On most (10/17) occasions, no mating was observed either before or after sharing. Similarly, at Kanyawara (1990–2006), we recorded only 7 instances of sharing with an estrous female when both the male and female were present in the party for 3 hours before and 3 hours after sharing. Mating occurred on only two of these occasions—once immediately before sharing, and once immediately following sharing. Thus, data from Gombe, Ngogo, and Kanyawara do not support the prediction that short-term male mating success is correlated with sharing (Table 1).

Discussion

Based on our review of meat sharing and sexual behavior in chimpanzees, the widely-cited hypothesis that male chimpanzees exchange meat with females to increase their short-term mating success is not supported. We challenge the fundamental
assumptions of the hypothesis in light of existing models of exchange in mating markets. We note that female chimpanzees rarely require material incentives to mate, and that amid the competitive mating system of chimpanzees, the potential effects of such incentives on the reproductive success of either partner are insufficient to drive the evolution of meat sharing, particularly at the low rates reported from East African sites.

Our review challenges the evidence that has been cited in favor of the meat-for-sex hypothesis in two ways. First, we note that support for the hypothesis has only ever come from a single statistically significant result (Stanford et al., 1994). Second, that analysis included instances of hunting interest, rather than only unequivocal hunting behavior, and did not consistently distinguish females that were motivated to copulate from those that were not.

Our statistical tests of four key predictions of the meat-for-sex hypothesis lead to consistent results. Collectively, these studies reveal that 1) the presence of sexually receptive females does not increase hunting probability, 2) males do not share preferentially with sexually receptive females, 3) males do not share preferentially with particularly fecund females, and 4) sharing with females does not increase a male’s short-term probability of mating. Our analyses included two attempts to replicate early claims in favor of the meat-for-sex hypothesis, both of which failed.

If not for sex, then why do male chimpanzees share meat with females? We suggest that males share with females for the same reasons that they share with other males. Several studies have suggested that possessors share meat as a means of avoiding the costs associated with defending meat against persistent beggars.
sharing events at Tai can be explained by long-term meat-for-sex exchanges? Answers to such questions will indicate whether chimpanzees can maintain a long-term exchange of meat for sex, even in the absence of any short-term exchanges.

While the data from other study sites are sparse, the Tai result does not seem to be generalizable to other chimpanzee populations. At Ngogo, a given male was equally likely to mate with a female during sexual cycles in which he shared with her as during those in which he did not \((n = 12; \text{Mitani and Watts, 2001})\). Similarly, at Burgers’ zoo (Arnhem, The Netherlands), males neither copulated more \((\text{Hemelrijk et al., 1992})\) nor sired more offspring \((\text{Hemelrijk et al., 1999})\) with those females with whom they most frequently shared food. It is possible, therefore, that chimpanzee social dynamics are different in Tai from elsewhere. For example, the rate of meat transfers observed at Tai \((262 \text{ male-female meat transfers in 22 months [Gomes and Boesch, 2009]})\) was much greater than at the East African sites \((\text{eg., Ngogo: 103 male-female meat transfers in 37 months [Watts and Mitani, 2002a]}\)), while the rate of copulations observed was much lower \((0.144 \text{ copulations/hour [Gomes and Boesch, 2009]}\). These points suggest a different mating dynamic that may confer more agency to females at Tai.

Such inter-site variation raises the possibility that even short-term meat-for-sex exchanges among chimpanzees may one day be found. They have not been found yet, however.

Conclusions

The notion that chimpanzees trade meat for sex has been frequently cited by anthropologists seeking to understand the evolutionary roots of human behavior. Food sharing is commonly heralded as an important milestone in human evolutionary history \((\text{Isaac, 1978})\), linked with such quintessential human traits as central place foraging, large, metabolically expensive brains \((\text{Aiello and Wheeler, 1995})\), and even the concepts of fairness, equity, and punishment \((\text{Fehr and Schmidt, 1999; Gintis, 2000})\). Arguably, one of the most important putative consequences of food sharing for human evolution was the development of the sexual division of labor and pair bonds. Males concentrated their foraging efforts on hunting game, a high-quality, high-variance food source, while females collected low-quality, low-variance foods such as roots and tubers. Sharing allowed both sexes to have a high quality diet \((\text{i.e., more meat})\) while minimizing starvation risk. This scenario would have created mating competition for the most successful hunters, who could advertise their prowess by sharing meat widely and exchanging meat for sex, thus providing the foundation for pair bonds. While meat-for-sex anecdotes in chimpanzees are often used as evidence for the evolutionary origin of these exchanges, the link between food sharing, pair bonding, and the derived nature of the hominin sexual division of labor would actually be made more tenuous if our promiscuously-mating ancestors conducted similar exchanges of meat for sex. However, we conclude that such scenarios are inappropriate. Instead, we find that meat transfers in chimpanzees are rarely sexually motivated. Short-term sexual bartering in humans appears to be a unique and derived trait.

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