Moral Sentiments and Material Interests
The Foundations of Cooperation in Economic Life

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The Natural History of Human Food Sharing and Cooperation: A Review and a New Multi-Individual Approach to the Negotiation of Norms

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3.1 Introduction

Humans share food unlike any other organism does. Many other animals, including eusocial insects (bees, ants, termites); social carnivores (lions, wolves, wild dogs); some species of birds (e.g., ravens) and vampire bats actively share food. However, the patterning and complexity of food sharing among humans is truly unique. Unlike other mammals, for which food sharing between mothers and offspring is limited largely to lactation during infancy, human parents provision their children until adulthood. Moreover, the sharing of food between human parents and their children continues bidirectionally until death in most traditional non-market societies. Additionally, marriage is universal among human societies, and husbands and wives regularly share food with one another throughout their marriage. Food sharing within human families is based upon a division of labor in subsistence effort by age and sex, where tasks are divided and the proceeds of work are shared. In fact, within-family transfers of food are so universal among humans that they are largely taken for granted and have rarely been systematically studied. This gap in anthropological research is ironic, since the sexual division of labor and the concomitant sharing of food between spouses and between parents and offspring have figured prominently in several models of hominid evolution (e.g., Isaac 1978; Lancaster and Lancaster 1983).

In addition to within-family food transfers, food sharing sometimes extends beyond the nuclear family in many societies. Most recent research on food sharing has focused on food transfers among adults living in different households. The majority of this research has been conducted in small-scale societies, particularly hunter-gatherers and groups that combine simple horticulture with hunting and gathering.
(forager-horticulturalists). There are two reasons for this focus. First, intragenerational food sharing is pervasive among hunter-gatherers and many forager-horticulturalists; they are often referred to as "egalitarian societies." Second, hominids lived as hunter-gatherers for the vast majority of their evolutionary history (which has lasted over two million years). Agriculture originated only about 10,000 years ago and has been practiced by the majority of the world's population for only two or three millennia. Since most hunter-gatherers share food on a daily basis, studies of food transfers among foragers may provide important insights into the evolutionary basis of human food sharing and more generally, about the origins of human hyper-sociality.

This chapter has three principal objectives. The first is to provide a brief overview of existing theory and research about food sharing in small-scale societies for nonspecialists. In the first part of the chapter, we outline the principal hypotheses proposed to account for variation in food sharing and evaluate available evidence pertaining to these hypotheses. The second objective is to present evidence regarding why we consider it necessary to rethink existing approaches to food sharing. In this second part of the chapter, we argue that intragenerational resource flows are critical to the understanding of intragenerational sharing and that neither the human life course nor human intelligence could have evolved without long-term imbalances in flows of food between families. We suggest that future research on this topic should consider small-group decision processes and the emergence of institutionalized sharing norms. In the third part of the chapter, we review several case studies of food sharing in different societies and across contexts within societies as a preliminary step towards building a theory of how these norms may correspond to local ecological conditions. The chapter concludes with a discussion of new directions for research and some major unresolved questions that should be addressed.

3.2 Part I: Theories and Empirical Evidence

Six different theories have been proposed to explain the existence and patterning of intragroup food sharing.

1. Reciprocal altruism
Several investigators have proposed that reciprocal altruism (Trivers 1971), where food is exchanged at one point in time for food at some later time, may explain many or most cases of human food sharing (Kaplan and Hill 1985; Winterhalder 1986; Smith 1988). The perverseness of reciprocal food sharing among humans is commonly explained in terms of the kinds of foods they acquire and their inherently "risky" nature (Kaplan and Hill 1985; Winterhalder 1986; Smith 1988). Human hunter-gatherers tend to specialize on the largest, highest-quality, most nutrient-dense foods available in their environments (Kaplan et al. 2000), and as a result, they experience high variance in foraging luck due to the difficulty in acquiring these items. For example, individual Ache hunters return empty-handed on 40 percent of the days they hunt, but on some days return with several hundred thousand calories of meat (Hill and Hawkes 1983). Hunting success is even more sporadic among large-game hunters, such as the Hadza, who only acquire meat on about 3 percent of their hunting days (Hawkes, O'Connell, and Blerton Jones 1991).

Since there are diminishing returns to consumption of large quantities of food (especially in environments where spoilage is a problem) and because food portions are very valuable to hungry individuals, reciprocal sharing can significantly reduce variation in day-to-day consumption and maximize the intertemporal utility of food. Reciprocal altruism therefore allows people to devote time and energy to the pursuit of large, asynchronously acquired, high-quality packages. Trade is a special form of reciprocal altruism where the return benefits of giving are in another currency, such as meat for sex, food for labor, or fish for yams. However, when the return benefit is a non-food currency, such as increased mate access, such sharing does not serve the goal of risk-reduction.

II. Cooperative acquisition and byproduct mutualism
Sharing may also enable individuals to achieve gains from cooperative pursuits of food. The acquisition of difficult-to-acquire foods, especially wild game, often requires the coordinated efforts of several individuals. However, usually only a single individual is identified as the owner of the acquired resource, determined by cultural-specific norms of ownership (e.g., the hunter who makes first lethal shot, the finder, the killer [Dowling 1968]). In many groups, sharing among task group members occurs as an initial wave of sharing (e.g., Pygmies [Bailey 1991; Harako 1976]). Owners may reward nonowners for their current cooperation by giving them shares of the resource, but this sharing may also act as a means of insuring future cooperation in similar food production activities. Thus, sharing is a form of trade-based reciprocal altruism, where labor is rewarded with food.
An alternative interpretation of the same phenomenon is that engaging in group production when there is sharing provides participants with higher per capita returns than if they produced food by themselves. Thus, group production may represent a form of byproduct mutualism (Clements and Stephens 1995; Dugatkin 1997; Alvard and Nolin 2002). Once rigid sharing norms exist in a population, the decision to participate in, say, a group whale hunt or cooperative monkey hunt should depend only on the higher per capita return rates relative to those that could be gained in solitary subsistence activities (see Alvard 2002). Thus, an advantage of strong sharing norms is that they act to transform the payoff structure of group food production strategies from that of a Prisoner’s Dilemma to that of mutualism.

III. Tolerated theft or scrounging

Building on the same insights regarding large asynchronously acquired food packages and diminishing returns to consumption of large food quantities, others have proposed that much apparent voluntary sharing may actually be “theft” or scrounging of food from food acquirers by individuals who have little or none (Blurton Jones 1984, 1987). This hypothesis is based upon the assumption that asymmetries between individuals in the marginal value of additional food can lead to contests over packages. The hungry person is more motivated to fight, while the person with more should relinquish some food because the lost food value is not worth the fight (Blurton Jones 1987; Winterhalder 1996). When power or “resource holding potential” is equal among contestants, a simple prediction of tolerated scrounging is that distributions will be highly egalitarian—such that any additional food portions have the same marginal value for each contestant (Winterhalder 1996).

Proponents of this view have pointed out that tolerated theft in the context of large, highly variable foods raises a secondary problem. Why do people spend time foraging for large packages if they know that much of what they acquire will be taken from them? Scrounging of large packages may effectively reduce their per capita consumption return rate below several other food production options in the environment, especially the pursuit of small packages (Hawkes 1993). To answer this question, Hawkes (1991, 1992, 1993) proposed that the pursuit of large packages, particularly hunted foods, is very sex-biased and that men acquire large packages to “show off” and garner attention. Men focus their efforts on acquiring large packages, precisely because others will scrounge from them. As a result, these men will gain the attention and support of scroungers, many of whom will be women. The payoffs to this attention presumably come in the form of increased access to mates and an increased number of offspring.

IV. Costly signaling

Costly signaling is an extension of the show-off hypothesis that may explain why certain individuals (usually men) pursue difficult-to-acquire foods that often yield suboptimal caloric return rates (Smith and Biege Bird 2000; Gurven, Allen-Arave et al. 2000; Biege Bird, Smith, and Bird 2001). The costliness of the signal insures the honesty of the information being advertised (Zahavi and Zahavi 1997; Grafen 1990; Johnstone 1997). The signal might provide information about phenotypic quality (such as disease-resistance) or about intentions to cooperate in the future. Recipients of the signal confer benefits on the generous donor not as payback for food given, but because information about the donor’s phenotypic quality makes that donor a desirable partner, mate, or ally. Costly signaling differs from showing off because it does not rely on tolerated theft to explain food transfers. Additionally, because the honesty of the signal makes the signaler an attractive partner, costly signaling avoids the second-order–collective action problem of who should pay prestige back to good hunters.

V. Nepotism based on kin selection

Because biologically related individuals share genes by descent, any behavior that sufficiently benefits kin can be favored by natural selection. According to models developed by Hamilton (1964), natural selection will favor altruism to kin when the benefits to the recipient, discounted by Wright’s coefficient of genetic relatedness between donor and recipient, outweigh the costs of giving. A simple prediction is that, all else being equal, close kin should receive shares either more frequently or in greater quantities than distantly related and unrelated individuals (Feinman 1979). It has also been argued we should expect to find greater imbalances in quantities given and received among close kin than among nonkin or distant kin (Hames 1987; Feinman 1979), who, presumably, would only share reciprocally. However, this might not be true if close kin are also reciprocity partners and if reciprocal altruism is an important factor influencing food transfers among kin (Gurven, Hill et al. 2000).

VI. Trait-group selection

Selection among groups has also been proposed in order to explain cooperation and food sharing within human groups (Wilson 1998; Boyd and Richerson in press; Boehm 1993). In group selection models, the
relative fitness of altruists is lower than that of selfish individuals within groups, but the average fitness of individuals in groups containing more altruistic members is greater than those in groups containing fewer altruists. Group selection could favor costly food sharing if the increased absolute fitness of altruists among groups in a metapopulation outweighs the decrease in relative fitness within groups, where “group” refers to any congregation of individuals (Wilson 1990, 1998). While the conditions favoring trait-group selection are much less stringent than those of older group selection models, its overall influence is still limited by grouping patterns and migration, and ultimately may be no more revealing than egoistic models (Harpending 1998; Krebs 1987). However, given the conflict between group and individual interests, cultural means of encouraging individuals to share food may increase the frequency of giving within groups (Simon 1990; Boyd and Richerson in press), leading to socially enforced egalitarian behavior (Boehm 1993).

3.2.1 Cross-cultural Evidence
There is a great deal of cross-cultural evidence that sharing is most common for large packaged resources characterized by high acquisition variance, especially wild game. Widespread pooling of large game animals is common among the Hadza (Hawkes 1993; Marlowe n.d.), Dobe !Kung (Lee 1979; Marshall 1976), G/wi (Silberbauer 1981), Ifaluk (Sosis, Feldstein, and Hill 1998), Ache (Kaplan and Hill 1985), Yanomamo (Hames 1990), and Gunwinggu (Altman 1987). While such pooling can significantly reduce variation in daily meat consumption, the outcome of risk reduction is consistent with all six models, even though risk reduction is only explicitly incorporated as a goal within the reciprocal altruism and group selection frameworks. This is because widespread sharing of relatively large sized game items, characterized by high within- and across-individual variability in acquisition, can be explained by future reciprocation (reciprocal altruism), demands and threats of hungry individuals (tolerated theft), honest signals of phenotypic quality (costly signaling), and the Pareto-optimal distribution solution maximizing group benefit (trait-group selection). Because the costs of sharing decrease with increased package size of the resource (assuming diminishing returns), it is not surprising that large package size is a significant predictor of sharing for meat and other food items such as fruits, cultigens, and market foods (Hames 1990; Gurven, Hill et al. 2000; Gurven, Hill, and Kaplan 2002; Kitanishi 1998).

Thus the observation that the sharing of large packages is widespread does not help distinguish between the models discussed in section 3.2. The empirical findings relevant to understanding which models are most appropriate for explaining much of the variance in sharing within and across groups are generally concerned with three issues: contingency of giving upon receiving, producer control over distributions, and imbalances between families in what is given and received.

3.2.1.1 Contingency  Contingency is the feature of sharing relationships that is critical for distinguishing between reciprocal altruism and other models (Rothstein and Pierroti 1988; Hill and Kaplan 1993). General contingency requires that all giving be balanced by all receiving, while specific contingency requires that giving to specific others be balanced by receipts from those same individuals (Hames 2000; Gurven, Hill et al. 2000). Specific contingency is usually estimated as the correlation between the percentage or quantity of food given by A to B with the percentage or quantity given by B to A over some appropriate sample period. Contingency can be measured within food categories (such as meat for meat, roots for roots, and so forth) or for all food, which includes exchange across food types. In order for reciprocal altruism to be beneficial to a donor, donors should preferentially give to those who are likely to share with them in the future (specific contingency). Costly signaling, on the other hand, requires that the prestige-related benefits from signaling outweigh the costs of producing food that is widely shared (general contingency). Neither tolerated theft, kin selection, nor trait-group selection predict that food donations will be conditional upon expected return.

Specific contingency has only been measured in four groups, all of which are in South America—the Yanomamo (Hames 2000), Hiwi (Gurven, Hill et al. 2000), Ache (Gurven, Allen-Arave et al. 2000), and Pilaga (Henry 1951). Correlations which describe specific contingency for all foods are significantly greater than zero, but not very high (between 0.2 and 0.5 [Gurven in press], see discussion of imbalances in section 3.2.1.3), while within resource categories, contingency is often highest for cultivated and collected foods. These results are most consistent with reciprocal altruism and least consistent with tolerated theft, because tolerated theft predicts that there should be no relationship between giving and receiving. Among the Ache, however, there is no evidence of specific contingency for wild game over the duration of single foraging trips, nor for game items brought back to the
permanent colony. This is inconsistent with reciprocal altruism, unless sharing among the Ache rewards group work effort (cooperative acquisition and mutualism, discussed in part III). Nevertheless, further research is needed to determine whether these significant positive correlations imply that the time-discounted value of food returns is sufficient to offset the present costs of giving.

Anecdotal evidence that giving is balanced by future receiving and that those who do not give do not receive is found in many traditional societies. As one Maimande explained, "If one doesn't give, one doesn't get in return ... Some people are specifically excluded from most distributions because they never or only rarely give any of their products to us" (Aspelin 1979, 317). Similar anecdotes exist among the Agta (Peterson 1978; Ivon Griffin personal communication), Pintupi (Myers 1988), Sirionó (Holmberg 1969, 45), and G/wi (Silberbauer 1981, 463). Although there is an emphasis on peoples' expectations for future receipt in these and other ethnographies, the extent of time depth separating episodes of giving and receiving is often unclear. Indeed, Sahlin's (1972) use of the term "generalized reciprocity" was meant to reflect short-term imbalances, especially among kin, that are eventually balanced over the span of peoples' lives.

General contingency or balance has been measured in five societies—the Ache (Gurven, Hill, and Kaplan 2002), Hiwi (Gurven, Hill et al. 2000), Meriam (Biege Bird and Bird 1997), Pilaga (our analysis of Henry 1951), and Yanomamo (Hames 2000). These studies showed mixed support for general balance. While the lack of specific 'balance contradicts reciprocal altruism, the presence of general balance is consistent with indirect reciprocity (Alexander 1987; Boyd and Richerson 1989) or costly signaling, where the return benefit to the donor is food. If the return benefit is in another currency, such as increased mating opportunities, then a lack of general balance is not inconsistent with costly signaling.

3.2.1.2 Producer Control Reciprocal altruism and kin selection require that producers maintain some control over the distribution of foods they acquire, whereas tolerated theft assumes no producer "rights." If producers have no control over the distribution of certain items, then those food items may act as partial public goods (Dowling 1968; Hawkes 1993). Despite observations of widespread game distributions in some groups (e.g., Ache [Kaplan and Hill 1985], G/wi [Silberbauer 1981], Hadza [Hawkes 1993], Western Desert Aborigines [Gould 1980]), several lines of evidence indicate that producers often maintain significant control over distributions in many—if not most—cultures. First, as shown in the previous section, there is often a bias in sharing towards those who shared with the producer. Additionally, there are clear biases in distributions towards close kin living in other families at the expense of distant kin and unrelated families (Gunwinggu [Altman 1987], Copper and Netsilik Eskimo [Damas 1972], Pilaga [Henry 1951], Hiwi [Gurven, Hill et al. 2000], Kaingang [Henry 1941], Batek [Endicott 1988], Pintupi [Myers 1988], Washo [Price 1975], Yanomamo [Hames 1989], Machiguenga [Kaplan 1994], Ache [Kaplan and Hill 1985; Gurven, Hill, and Kaplan 2002], Iatuk [Sosis 1997], Basarwa [Cashdan 1985]), and to those participating in the hunting party (Netsilik Eskimo [Damas 1972], Nyae Nyae Ikung [Marshall 1976], Iatuk [Sosis 1997], Pintupi [Myers 1988], Washo [Price 1975], Mbuti [Ichikawa 1983], Aka [Bahuchet 1990; Kitaniishi 1998], Efe [Bailey 1991], Lamalera [Alvard 2002], Northwest Coast Indians [Gould 1980]). Among the Hiwi and Ache at the settlement, there are clear kin biases in sharing, even when controlling for residential distance. Finally, several ethnographies are explicit about the ownership of shares after an initial distribution, even if others still have not received any meat (Efe [Bailey 1991, 100]; Nyae Nyae Ikung [Marshall 1976, 363]).

Second, hunters frequently consume portions of kills (e.g., internal organs and marrow) at the kill site and usually no one begrudges them this right (Batek [Endicott 1988]; Hadza [Marlowe n.d.]; Nyae Nyae Ikung [Marshall 1975]; G/wi [Silberbauer 1981]). Third, producers often keep significantly more than 1/4 of the game packages they acquire, even though others in the camp or village may not possess any meat (Gunwinggu [Altman 1987]; Yoruba [Hill and Kaplan 1989]; Yuqui [Stearman 1989]; Yanomamo [Hames 2000]; Hadza [Hawkes et al. 2001]; Ache [Gurven, Allee-Arave et al. 2001]; Hiwi [Gurven, Hill et al. 2000]). Finally, the frequent observations of incessant demands for food in many foraging societies (Peterson 1993; Chagnon 1983) does not mean that producers are powerless to ignore or reject requests for food made by other group members. There is evidence that nonproducers do not possess automatic claims to shares among the Pintupi (Myers 1988), the Aka (Bahuchet 1990, 38), Agta (Griffin 1984), Pilaga (Henry 1951), and Sirionó (Holmberg 1969, 88).

3.2.1.3 Imbalances: Relative Need, Bargaining, and Signaling Several ethnographies have reported large short-term between-family
imbalances, but long-term balance in food transfers, consistent with
genralized reciprocity (e.g., Batek [Endicott 1988, 118]; Kaingang
[Henry 1941, 101]). While short-term imbalances are relatively easy to
measure, the existence of long-term balances in terms of lifelong sym-
biosis is much more difficult to confirm. Nepotistic food sharing based
on kin selection can predict imbalances in food sharing, even though
kinship is a symmetrical relationship. Food given to dependent and
unskilled offspring and relatives can have a large benefit at a relatively
small cost to a skilled producer (B < C in Hamilton’s Rule). Similarly,
the downward flow of food provides useful calories to younger kin of
higher reproductive value (Rogers 1993). Over the course of an
individual’s lifetime, current evidence suggests that among the Ache, Piro,
Machiguenga, and perhaps other traditional groups, children are net
losses to parents, and thus children’s debt is never repaid directly, but
is instead redirected to grandchildren (Kaplan 1994; see part II of this
chapter). We should expect to find smaller short-term imbalances in
transfers among kin of similar age. Allen-Arave et al. (n.d.) find that
among the Ache, imbalances over a four-month sample period are
smaller among kin of similar ages (e.g., siblings) than among kin of
disparate ages (e.g., between older parents and their adult offspring).

Apart from kinship, there is good evidence that large short-term
and long-term exchange imbalances among individuals and families
occur among foragers and forager-horticulturists. The highest food
producers among the Ache, Efe, Pilaga, and Yuqui consistently gave
away more than they received as compared to low producers (Kaplan
1941; Stearn 1989). Indeed, observations that high producing Ache
and Hadza hunters often do not receive in-kind compensation for their
generosity initially led to the proposition that reciprocal altruism was

Imbalances in quantities transferred across individuals or families
can be interpreted in several ways. First, the short-term nature of most
field studies places an arbitrary horizon on the delayed time for recipro-
cration, and the sampling bias associated with any brief series of
snapshots of interfamly exchange relations is likely to result in some
degree of imbalance. Hames (2000) argues that meticulous score-
keeping meant to ensure balance should be found across pairs of
distantly interacting individuals, where trust is weak (e.g., market
transactions), while imbalances might be quite common among indi-
nividuals who interact over extensive periods of time.

Second, an imbalance may be intentional if sharing is based on
the signaling of phenotypic or genotypic quality. Imbalances in turtle
meat exchanges, where hunters expend a great deal of energy to pro-
vide community feasts, are most likely due to costly signaling (Bliege
Bird and Bird 1997; Smith and Bliege Bird 2000). Similarly, an imbal-
cance is expected if the return benefits of reciprocal altruism or costly
signaling are in other currencies. High-producing Ache hunters may
give away more than they receive, but they obtain greater mating
opportunities and higher offspring survivorship (Hill and Hurtado
1996). Yuqui and Tsimane hunters sometimes trade portions of their
kills for garden products (Stearman 1989; Chichón 1992), while Kukuyu
with unsuccessful gardens will trade labor for access to a neigh-
bors’s manioc patches (Carneiro 1983).

As mentioned above, contingency estimates well below 1.0 suggest a
relatively high occurrence of exchange imbalances among pairs of
families. It is important to mention that all measures of contingency are
based on quantities of food shared among families. Bargaining theory,
however, can lead to outcomes consistent with reciprocal altruism but
inconsistent with strong balance in food quantities (Stähler 1972; Hill
and Kaplan 1993; Sosis, Feldstein, and Hill 1998; Gurven, Hill et al.
2000). If donors continue giving portions to others as long as the
expected future benefits outweigh the current costs of giving relative
to other options, there is no reason to expect the exchange of equal
quantities over time. The extent of imbalance should be a function of
differential wealth holdings, influence, status, and need. One calcula-
tion of contingency that measured the balance in “value” transferred
across families—by incorporating the frequency and sizes of shares—
found a slightly higher level of balance among Hiwi and Ache foragers
(Gurven 2004).

There has been much written on the importance of “need” and the
direction of food flows (Woodburn 1982; Barnard and Woodburn
1988; Winterhalter 1996), supporting the notion that “if there is
hunger, it is commonly shared” (Marshall 1976, 357) and possibly the
group selection hypothesis. Among the Ache, Maimande, G/wi, and
Hiwi, shares are often given in proportion to the number of consumers
within the recipient family (Gurven, Hill, and Kaplan 2002; Aspinel
1979; Silberbauer 1981; Gurven, Hill et al. 2000). Batek families with
high dependency tend be net consumers while those with low depend-
dency are net producers (Cadelina 1982). There is additional evidence
that older men, with larger families, preferentially benefit from sharing
networks at the expense of younger men's labor, especially if one considers bridesservice payments (Efe [Bailey 1991], Gunwinggu [Altman 1987], Kutsie [Kent 1993], Yanomamo [Ritchie 1996], and Agta [Bin Griffin 1984]). Differential need among families leads to different costs and benefits of giving across families, and should therefore influence bargaining outcomes and observed levels of balance. We explore this issue further in part II of this chapter.

Although certain levels of imbalance may be due to differential need, there is much evidence to suggest that such imbalances are sometimes tolerated only within limits. Those who do not produce or share enough are often subject to criticism, either directly or through gossip, and social ostracism. Anecdotes of shirkers being excluded from distributions until they either boosted their production or sharing levels are found among the Maimande (Aspinel 1979), Piila (Henry 1951, 199), Gunwinggu (Altman 1987, 147), Wasbo (Price 1975, 16), Machiguenga (Baksh and Johnson 1990), Agta (Griffin 1984, 20), and Netsilik Eskimo (Balicki 1970, 177). However, other ethnographies report the persistence of long-term imbalances without any obvious punishment, exclusion, or ostracism (Chácobo [Prost 1980, 52]; Kaingang [Henry 1941, 101]; Batek [Endicott 1988, 191]), although these anecdotes suggest that such imbalances are due to a small number of low producers within the group.

In summary, there is substantial cross-cultural evidence supporting the view that reciprocal altruism of some sort underlies much food-sharing behavior. First, in many societies producers appear to exert control over the distribution of resources. Second, although specific contingency of giving upon receiving has been measured in only a few cases, there is evidence over the short term that people form preferential food-sharing partnerships with high rates of giving and receiving and share less with those who give less (meat sharing in the forest among the Ache is one exception, however) (Kaplan and Hill 1985). There are also a plethora of qualitative reports suggesting that giving and receiving are contingent in many or most cultures and in different contexts.

At the same time, persistent imbalances in amounts given and received between families suggest that strict reciprocal altruism cannot account for all food sharing between families. Some of those imbalances may be due to kin selection, costly signaling, tolerated theft, trait-group selection or some combination of these four forces. In the next section, we sketch the importance of food sharing in the evolution of human life. We show that the evolved life history of humans required long-term imbalances in food flows. We also present a new way to understand imbalances in terms of multi-individual decision processes and long-term mutual benefit.

3.3 Part II: Human Life History and Food Sharing

3.3.1 Features of Our Human Life History

The distinctive life history of humans is related to their unique foraging niche relative to that of other mammals (and even primates). Five distinctive features of the human life course are noteworthy.

1) an exceptionally long lifespan
2) an extended period of juvenile dependence
3) support of reproduction by older post-reproductive individuals
4) male support of reproduction through the provisioning of females and their offspring
5) a large brain and its associated capacities for learning, cognition, and insight

Humans have a very flexible foraging strategy, consuming different foods in different environments, and this flexibility has allowed us to survive successfully in all of the world's terrestrial environments. In another sense, however, the human foraging niche is very specialized. In every environment, human foragers consume the largest, most nutrient-dense, highest-quality, and most difficult-to-acquire foods, using techniques that often take years to learn (Kaplan et al. 2000, Kaplan 1997).

This foraging niche is related to human life history because high levels of knowledge, skill, coordination, and strength are required to exploit the suite of high-quality, difficult-to-acquire resources humans consume. The attainment of those abilities requires time, a significant commitment to development, and a large brain to support the learning, information processing, and planning underlying those skills. This extended learning phase during which productivity is low can be compensated by higher productivity during the adult period and subsidized by an intergenerational flow of food from old to young. Since productivity increases with age, the time investment in skill acquisition and knowledge leads to selection for lowered mortality rates and greater longevity, because the returns on the investments in development occur at older ages.
There are three foraging groups (the Ache, Hadza, and Hiwi) and two groups of forager-horticulturalists (the Machiguenga and Piro) for whom quantitative data are available regarding age-profiles of food consumption and production. All of these groups display similar age-profiles of net food production. Children are largely supported by their parents until about age eighteen (when food production approximately equals consumption), after which productivity rises steeply through the twenties until the mid-thirties. The more skill-intensive the task, the greater the delay to peak performance and the greater the increase in productivity with “on-the-job-training” (Bock 2002). High productivity is maintained until the mid-sixties when the deleterious effects of senescence become significant. This pattern of development and aging bears a striking resemblance to modern societies, where wages depend on education-based capital and the ages eighteen and sixty-five have similar significance.

Figure 3.1 shows survival probabilities and net production by age for wild chimpanzees, our closest living relatives, and modern human hunter-gatherers living under conditions similar to our evolutionary past (see Kaplan et al. 2000, 2001 for details on data sources). It is evident that the chimpanzee net production curve shows three distinct phases. The first phase, to about age five, is the period of complete to partial dependence upon mother’s milk and of negative net production. The second phase is independent juvenile growth, lasting until adulthood, during which net production is zero. The third phase is reproductive, during which females, but not males, produce a surplus of calories that they allocate to nursing. Humans, in contrast, produce less than they consume for about twenty years. Net production becomes increasingly negative until about age fourteen and then begins to climb. Net production in adulthood in humans is much higher than in chimpanzees and peaks at a much older age, reflecting the payoff of long dependency. More precisely, human peak net production is about 1,750 calories per day, reached at about age forty-five. Among chimpanzee females, peak net production is only about 250 calories per day and, since fertility decreases with age, net productivity probably decreases throughout adulthood. By age fifteen, chimpanzees have consumed 43 percent and produced 40 percent of their expected lifetime calories, respectively; in contrast, humans have consumed 22 percent and produced only 4 percent of their expected lifetime calories! In fact, the human production profile requires a long lifespan and would not be viable with chimpanzee survival rates, since expected lifetime net production would be negative (Kaplan et al. 2000).

These results imply a highly structured life course in which physiological and behavioral processes are coordinated. The greater proliferation of neurons in early fetal development among humans, as compared to monkeys and apes, has cascading effects, extending other phases of brain development and ultimately resulting in a larger, more complex, and effective brain. From a behavioral point of view, although cognitive development is largely complete among chimpanzees by about eight years of age, formal abstract logical reasoning does not emerge in humans until age sixteen to eighteen. This is the age when productivity begins to increase dramatically among human foragers.

3.3.2 The Evolutionary Role of Sharing
A central thesis of this chapter is that the human life course could not have evolved without long-term imbalances in food transfer within and among families. First, it is clear from the above figures that if children are eating more than they produce for some twenty years, those deficits must be subsidized. Surplus food provided by older people
and those with few dependents can be utilized to finance this long developmental period.

Second, those data represent average production and consumption by age, combining data from both sexes. Men and women, however, specialize in different forms of skill acquisition with correspondingly different foraging niches and activity budgets and then share the fruits of their labor. The specialization generates two forms of complementarity. Hunted foods acquired by men complement gathered foods acquired by women, because protein, fat, and carbohydrates complement one another with respect to their nutritional functions (see Hill 1988 and Hames 1989 for a review) and because most gathered foods, such as roots, palm fiber, and fruits are low in fat and protein (nuts are an exception). The fact that male specialization in hunting produces high delivery rates of large, shareable packages of food leads to another form of complementarity. The meat inputs of men shift the optimal mix of activities for women, increasing time spent in childcare and decreasing time spent in food acquisition. They also shift women’s time to foraging and productive activities that are compatible with childcare and away from activities that are dangerous to them and their children.

There are data on the productivity of adults for ten foraging societies (see Kaplan et al. 2000 for details). On average, men acquired 68 percent of the calories and almost 88 percent of the protein; women acquired the remaining 32 percent of calories and 12 percent of protein. After subtracting their own consumption (31 percent of total calories), women supply only 3 percent of their offspring’s caloric deficit (i.e., children’s consumption minus their production), while men provide the remaining 97 percent! Men not only supply all of the protein to offspring, but also the bulk of the protein consumed by women. This contrasts sharply with most mammalian species (>97 percent), where the female supports all of the energetic needs of the offspring until it begins eating solid foods (Clutton-Brock 1991) and the male provides little or no investment.

In addition to specialization among men and women, specialization in productive activities by age is equally important. Foragers and forager-horticulturalists typically assign low skill/low strength activities (such as collecting fruits or fetching water) to children, high strength/high skill activities (such as hunting and extractive foraging) to prime-aged adults, and low strength/high skill activities (such as child care and craft manufacture) to elderly people (Bock 2002; Gurven and Kaplan n.d.; Kramer 1998). In this way, family returns from labor are maximized. It also appears to be the case that there is some specialization in activities within age-sex classes. Although this is less documented, anecdotal reports suggest that some men spend more time gathering or farming and others more time hunting (Tsimane [Chichón 1992], Yuqui [Stearman 1989]), and there is even specialization in hunting roles and in prey types pursued.

There are also imbalances between families that support this intensive mothering characteristic of human life histories. Figure 3.2 (adapted from Lancaster et al. 2000) compares the acquisition of calories and reproductive status of baboons (Allmann 1980) with Ache foragers (Hill and Hurtado 1996). Time spent foraging during the day is presented in relation to reproductive status for female baboons, where foraging time includes both travel and feeding time. Baboon mothers are hard pressed to meet the demands of lactation. When they must produce energy beyond their own maintenance needs, their daily time budget is stretched to the limit. They cannot afford to increase their.

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**Figure 3.2** Production and reproductive status. Adapted from Lancaster et al. 2000.
travel time, which would be energetically costly, especially since they must carry their infants. Instead they increase their feeding time by reducing resting and socializing to about 15 percent of the day. Lactating baboons thus work harder. In fact, female baboons have higher mortality rates when lactating than when cycling or pregnant (Altmann 1980).

In contrast, when lactating and even when they have dependent juveniles to be fed, Ache, Efe, and Hadza women reduce their time spent foraging for food (Hurtado et al. 1985; Ivey 2000; Hawkes, O’Connell, and Blarton Jones 1997). It appears that human females are able to reduce time spent in energy production when they are nursing, even though their caloric consumption must increase to support lactation. Among the Ache, most of women’s food production is derived from pounding the pulp of palms to produce starch. About 60 percent of the starch that women produce on extended foraging trips is shared outside the nuclear family (Kaplan and Hill 1985) with no bias towards close kin. Since lactating women produce much less palm starch than women without a baby, this pattern of sharing means that there are net food transfers from women to other women over periods of several months to several years.

Third, and most important for the present discussion, even with such extensive cooperation within families, additional flows of food between families are necessary to support this life history pattern. The fact that parental provisioning does not cease when children are weaned means that the caloric burden on parents increases as they produce more children. The diamonds show how the net demand on Ache parents changes with age as they produce additional children (viewed in terms of the man’s age). Demands peak between forty and fifty years of age and remain significant until age sixty. Even though food production increases with age to about age thirty-five or forty and remains high, demands increase faster than food production. The triangles show how net family food production (calories produced minus calories consumed by self and offspring) changes with the age of the man. These data show that there must be net transfers from the families of younger men to the families of older men! Moreover, there is a great deal of variance among men in both family size and productivity. Family size is inherently stochastic, due to both infant and child mortality and to individual differences in fecundity (see Hill and Hurtado 1996). There are also large differences in hunting ability among men. For example, there is a five-fold difference in the long-

![Figure 3.3](image)

**Figure 3.3**

Family demands and net family production.

term average hunting returns between the best and worst hunter in the sample of Ache men (Hill et al. 1987). Similar discrepancies in hunting ability across men have been found among the !Kung (Lee 1979), Hadza (Hawkes, O’Connell, and Blarton Jones 2001), Hiwi (Gurven, Hill et al. 2000), Gunwinggu (Altman 1987), Agta (Bion Griffin 1984), and Machiguenga (Kaplan unpublished data). Therefore, even among men of the same age, there must be net transfers over the long term from families producing a surplus to families producing a deficit.

These food transfers provide great reproductive benefits. The ability to “borrow” and “lend” across the life course is necessary for subsidizing the juvenile learning period (see Kaplan and Robson 2002 and Robson and Kaplan 2003 for theoretical models of the evolution of such age transfers and their relationship to learning). If families had to “balance their budget” at every period, they would either have had to lower their fertility or force their older children to fend for themselves. This would most likely increase childhood and adolescent mortality and lower rates of skills acquisition. Adolescent males could not afford to hunt, because their returns are so low during the learning period. Moreover, there would be no way to buffer the risks associated with
the stochasticity of family size and child demands. If families needed to support all of their individual food needs, regardless of whether few or many children survived, they would be forced to lower fertility or reduce child subsidies. Similarly, the ability of women to shift production across time without changing consumption probably increases infant survival and decreases the length of interbirth intervals, thereby increasing the total reproductive success of women. When the opportunity costs of food acquisition are high due to the need to care for infants, women may produce less when they have infants and then work harder when those opportunity costs are low (i.e., when they have no infant to nurse and protect).

3.3.3 The Problem with Dyadic Reciprocal Altruism

It is unlikely that such a system of sharing would be stable with strictly dyadic reciprocal altruism. Reciprocal altruism will only emerge among self-interested actors if there is repeated interaction that rewards cooperation and punishes defection. In terms of reciprocal altruism, there is no incentive for young or older adults with small families to support older adults with large families. Those older adults with large families will never produce a surplus to "pay back" those subsidies, because they are likely to die before the young adults reach the age in which they need assistance to support their families. While it might be argued that there is intergenerational reciprocity where the children of the older adults, in turn, support the families of those who helped them, the long time periods between changes in directional flows would make such arrangements inherently risky. There is a great deal of mobility between hunter-gatherer bands and residential arrangements are not stable over long periods. There is no guarantee that children who are helped when they are young will live in the same band as those who helped them. The same argument applies to sharing between non-nursing and nursing women. Additionally, time discounting of benefits received in the distant future (relative to the present consumption payoffs from not sharing) also makes intergenerational reciprocity unstable (Hawkes 1992).

Similarly, reciprocal arrangements regarding the stochasticity of family size are unlikely to emerge with dyadic relationships. If family size variation is due primarily to random luck, it may be beneficial for two individuals to agree at the start of their reproductive careers to an arrangement in which the individual who ends up with fewer surviving offspring agrees to support the one with more surviving offspring. However, once the outcome is known, there is no incentive for the one with fewer children to provide the support, since this family will never need the payback and there is no way to enforce the bargain.

Our thesis is that humans have found ways to take advantage of the gains from such trades, even though these gains would not emerge through dyadic reciprocal altruism. We propose that multi-individual negotiations result in the emergence of social norms that are collectively enforced. We base this proposal on a result obtained by Boyd and Richerson (1992), and treated more recently by Bowles and Gintis (2000), in which cooperation is modeled with punishment. These four researchers found that cooperation can be stable in large groups, if noncooperators are punished and if those who do not punish noncooperators are also punished. In fact, they found that any social norm could be stable as long as both those who disobey and those who fail to punish those who disobey are punished. However, we suggest that self-interested actors also negotiate these norms, weighing the individual costs and benefits of different social norms.

3.3.4 Two Thought Experiments

Imagine the following scenario. A woman returns from collecting berries and pounding palm fiber with her bawling infant. A wingless wasp stung her baby while she had put him down to pound the fiber, and the baby is in great pain. She is frustrated and says to the other women in camp, "This is crazy for me to go out and pound fiber when I have such a young baby. I would gladly work twice as hard when he is a little older if I could concentrate on watching him now." A few days later when the baby's wound is infected and the child has a fever, another woman, remembering a similar incident she experienced a few years ago, says, "You know, Singing Deer is right. We should work hard when we have no baby on the breast and allow those with a young one to care for it well." Another woman, who has not had a child in the last ten years, says, "Why should we work to feed other people's babies? If you have a baby, you must feed it." Other men and women consider their own situation (as well as the situation of their children) and present their opinions. Eventually a consensus (or at least, an agreement) is reached, with those in the minority either agreeing to go along with the new norm or leaving to live with "less foolish" people. However, one woman, who is not nursing, hardly pounds fiber at all. Other women begin to gossip about her, remarking upon how lazy she is, because she has no child to care for. She notices that the
shares she receives in food distributions start to become less generous and begins to suspect that others are talking about her behind her back. She leaves and pounds a large quantity of fiber, which she generously shares with the rest of the group. She can feel the warmth of others return and has learned her lesson.

We consider another similar scenario. A fifty-year old man turns to another older man and exclaims, “Look at these lazy young men! They come back to camp at midday and play around. Yet you and I have lots of children to feed and no food to give them. What will those boys do when they have big families to feed?” The other older man agrees, adding, “How do I know if that lazy one is good enough for my daughter? How do I know if he will get enough food to keep her children healthy? He should come to my fire and bring me lots of meat—then I will know.”

The young men are not very enthusiastic, because they do not like hunting all day long, but they are reluctant to anger the men whose daughters they favor. One young man, who is a good hunter for his age, realizes that he could take advantage of such a system and starts to hunt longer hours, giving the older men generous shares from his hunt. The other young men, afraid of being outdone, also begin to hunt longer hours and share the fruits of their labor more generously.

While admittedly hackneyed, these scenarios are meant to reflect the ongoing discussions and commentaries about sharing, work effort, and laziness that are so pervasive in foraging societies. We do not mean to suggest that all social norms are explicitly negotiated with words or that norms solidify over a short period as a result of a few conversations. In some circumstances, lack of compliance and “voting with one’s feet” are almost surely involved in those negotiations. In fact, we know virtually nothing about how standards for appropriate behavior emerge and change in small-scale societies without official means of enforcement. It is likely that majority-rule voting arrangements are not adhered to in a strict sense, since some individuals exercise undue influence (e.g., kombeti among Aka, kapita among Efe [Hewlett and Walker 1990], and chiefs among Yuqui [Stearman 1989]). Nevertheless, we propose that such multi-individual negotiations, partly verbal and partly nonverbal, do result in social norms and that the weight of collective opinion, based upon the individual costs and benefits of norms in given contexts, determines accepted patterns of behavior. In the next section, we develop a preliminary framework for explaining variation in norms regarding cooperation.

3.4 Part III: A Preliminary Framework for Explaining Sharing Norms

We propose that social norms of sharing reflect the relative strengths of two opposing forces: gains from cooperation and possibilities for free-riding. Sociocultural variation in potential benefits of cooperation and possibilities to free-ride on cooperative behavior determine cultural variability in norms of sharing and cooperative labor.

We also propose that in the course of our evolutionary history, natural selection has shaped our psychology to possess certain traits.

1) perceptual sensitivity to potential gains from cooperation
2) motivation to take advantage of those gains
3) perceptual sensitivity to opportunities for free-riding
4) motivation to avoid being a victim of free-riding
5) motivation to take advantage of opportunities for free-riding
6) perceptual sensitivity to the short- and long-term personal costs and benefits of social norms regarding cooperative behavior (from the perspectives of both the self and others)
7) motivation to negotiate social norms so that one’s own personal benefits from cooperation and free-riding are maximized
8) motivation to obey and enforce social norms so that punishment is avoided, and those who disobey norms or fail to enforce them are punished

Our proposal is that this psychology, the complex analytical brain, and the extended life history coevolved in the hominin line—all because of the dietary shift towards large, high-quality food packages and hunting. It is this feeding adaptation that generates the gains from group cooperation. The large size of the packages and the difficulty of their acquisition through hunting

a) facilitate sharing (imagine sharing blades of grass back and forth)
b) increase short-term variation in acquisition luck, since large packages are not abundant
c) require significant learning and experience
d) increase the disparity between production and consumption at the individual and family levels over the medium and long term
e) increase the benefits of collective action and cooperative pursuits, especially in hunting
f) generate economies of scale, since foods are often distributed in larges patches distant from residential locations.

These qualities generate large gains from intertemporal substitution in consumption and production over the short, medium, and long term; gains from specialization by age, sex, and perhaps individual qualities; gains from joint production and cooperative acquisition; and gains from turn-taking in acquisition of patchily distributed foods. The distribution and relative importance of each of those gains is likely to vary with local ecology and the foods exploited.

Possibilities for, and gains from, free-riding act against cooperation. Three factors are likely to influence the threat of free-riding. First, a larger number of individuals in cooperative networks is likely to increase the threat, because the ability to detect and punish free-riders probably diminishes with partner number. As group size increases, the probability that more than one individual free-riders may also increase (Boyd 1988). As the number of free-riders increases, costs of punishment increase and the incentive to cooperate decreases. Second, the quality of information about behavior is also likely to affect opportunities for free-riding. If work effort is difficult to monitor and if it is difficult to determine whether variance in productivity is due to acquisition luck or work effort, opportunities for free-riding may increase (Cosmides and Tooby 1992). Third, gains from free-riding are also likely to vary according to kinship relationships between participants. As overall relatedness decreases, the differences among optimal allocations of work and distribution across individuals are likely to increase.

Those opposing forces may have led to the evolution of some general moral sentiments—supported both by the motivational psychology of individuals and common cultural norms. Variation in need and production among individuals due to stochasticity should engender generosity and cultural norms emphasizing the value of generosity—perhaps mediated through costly signaling and reciprocal altruism. Sharing sentiments and norms would favor those who were unfortunate over the short or long run and require generosity from the more fortunate. Virtually every investigator who seeks to establish friendships with members of traditional subsistence populations, who are much poorer, feels the pressures associated with those sentiments. Similarly, temporary states affecting production or need—such as illness, nursing, and high dependency ratios—would also promote generosity. As mentioned above, the rule that larger families deserve and receive larger shares is very widespread. Conversely, variation due to lack of effort or laziness would not generate generosity and perhaps invoke ridicule or punishment. Indeed, laziness and stinginess are constant themes for gossip in traditional societies. Other things being equal, people should feel more generous towards (and trusting of) close kin, because of the reduced scope of conflicting interests.

At the same time that moral systems are likely to have such general guiding principles, there is scope for considerable variation in the norms of cooperation and sharing, depending upon the specific constellation of gains from cooperation and possibilities for free-riding. Of critical importance is the relationship between the size and composition of residential groups and the optimal size of cooperating units. In general, people will tend to organize residential groups so that they can take maximal advantage of the gains from cooperation and reduce risks of free-riding. Thus, many forager-horticulturalists in South America—such as the Machiguenga, Piro, and Tsimane—settle in extended family units, characterized by an older couple, their adult sons and/or daughters, and the founding couple’s grandchildren. Labor is divided by age and sex, and food is eaten communally. This system of communal production and consumption maximizes gains from specialization and from spreading consumption and production needs through the entire age-structure, while kinship and shared genetic interests in the young children minimize conflicts of interest.

Several factors may cause residential groups to differ in size and composition from their optimal structure for cooperation. Aggregation of larger groups is common, because of threats of violence (e.g., Yanomamo [Chagnon 1983]), lack of resources such as water or groves of trees (e.g., Dobe !Kung [Lee 1979]), and now schools and delivery points of social services (e.g., Chácobo [Prost 1980]). In these cases, restricted sharing—where some or all foods are shared with only a subset of the residential group—is the norm.

Restricted sharing systems appear to be particularly common when the primary gains from sharing derive from variance reduction in consumption and when gains from cooperative pursuits are small or restricted to only some resources or times of the year. A common principle evidenced in restricted sharing systems is that the breadth and depth of resource sharing depends on the size of food packages available. When food packages are small, they are shared with a few special
partners, with whom reciprocal sharing is very common. As package size increases, the size of sharing networks grows (increased breadth) and the percentage of the food kept by the acquirer’s family is reduced (increased depth).

Figures 3.4 and 3.5 illustrate features of this system. Figure 3.4 shows the percentage of sharing events by resource package size, in which specific Ache families receive shares at their permanent horticultural settlement. For each individual, sharing partners were rank-ordered from those who received most often to those who received least often. The x-axis displays the rank order and the y-axis gives the average percentage of occasions in which partners of each rank received shares. The data show that small packages are repeatedly shared with few individuals and that the size of sharing networks expands with large packages. Figure 3.5 (derived from data collected among the Hiwi and adapted from Gurven, Hill et al. 2000) is a path analysis predicting the total accumulation of food transferred between families over a six-month sample period, giving additional information about how partners are selected and about the size of shares given. Kinship predicts the spatial proximity between givers and receivers, which, in turn, predicts both how much was received in the past and the amount given in the present. In addition to kinship and proximity, the past history of sharing also predicts the amount given, suggesting that giving is contingent upon past receipts when controlling for these other factors.
Larger families also receive larger shares, as would be expected if need is being taken into account. Qualitative and quantitative reports from other societies suggest that similar patterns—kin bias, differential rules for sharing different resources (with increased breadth and depth of sharing with increased package size), contingency of sharing on the basis of past receipts, and larger shares to larger families—are found in other societies (Ifaluk [Sosis 1997], Eskimo [Damas 1972], Batak [Cadeiína 1982], Yora [Hill and Kaplan 1989]). It is not always the case, however, that the residential group is larger than the optimal sharing network for all resources acquired. In cases where very large packages are sometimes acquired (e.g., giraffe among the !Kung), it is sometimes necessary to inform members of neighboring groups about kills because the optimal sharing group size is larger than the optimal residential unit (Lee 1979).

Such systems tend to take advantage of the gains from cooperation while minimizing risks of free-riding. Reducing daily variation in the consumption of small packages requires fewer partners than in the case of larger packages. Thus, a small circle of trusted partners, frequently kin and neighbors, is most efficient. As package size increases, the benefits of a greater number of partners increase, but so too do the costs of free-riding.

Another important principle of restricted sharing systems is that work effort in cooperative activities is rewarded. Thus, when cooperative task groups form, food is often shared equally among the participants. When those task groups do not include members from all the families in the residential group, a system of primary and secondary sharing is very common. In the primary distribution, all participants in the cooperative activity receive approximately equal shares of the total catch (see part I of this chapter for a list of groups engaging in this practice). In secondary distributions, each individual that received shares redistributes his or her share to families that did not participate. Those shares are smaller and tend to be shared according to the size of the packages acquired in the manner discussed earlier in this section. Figure 3.6 from the Yora illustrates this pattern (see Hill and Kaplan 1989). The first two bars show the primary distribution to members of the foraging party and the second two bars show the secondary distribution. This is an “incentive compatible” system in which work effort is rewarded in the primary distribution and the other benefits of sharing (e.g., intertemporal substitution in consumption and production) are handled in the secondary distribution. In cases where representa-
and wage labor in larger settlements after their establishment of permanent peaceful contact with the larger Paraguayan society. For the first five or so years following settlement, agricultural fields were cleared and planted communally. All able-bodied men were expected to contribute labor in large work parties. This pattern resembled the cooperative economy of the past. However, within a few years, it became apparent that some people were often absent from work parties and resentments began to build. Some men tired of this system and cleared their own personal gardens. Communal fields became smaller and a system of private fields, with fewer friends helping each other, came to predominate. Similarly, even with hunted and gathered foods, the system changed from communal sharing of all game to a restricted pattern resembling the Hiwi one shown in figure 3.3. It is interesting to note that the Ache still retain the traditional sharing pattern when trekking in the forest, even though they revert to the new pattern when residing in the settlement. Similarly, the !Kung San appear to have undergone major changes in their system of food distribution, since becoming involved in a mixed economy and the larger state society. Again, the trend seems to be from more communal distributions towards more restricted sharing, with a great deal of bickering and strife during the transition (see Hostak 1981 and the associated N/ai film for qualitative accounts).

The transition to horticulture among the Ache and !Kung was very rapid, and encouraged through missionary assistance. As mentioned above, the establishment of private fields was quickly advocated and voted upon in local Ache meetings. This contrasts with the pattern in other groups such as the Hadza (Woodburn 1982) and the Batek (Myers 1988), where initial attempts at horticulture by a minority of the population met with abrupt failure. The first harvests of the few transitional farmers were exploited by incessant demands from those who did not farm, ultimately making farming an unproductive activity due to mutual adherence to more traditional norms of sharing.

3.5 Conclusion

We have proposed that in addition to individual reciprocal arrangements, humans appear to be able to take advantage of gains from cooperation in ways that are unexpected by pair-wise game models. We suggested that people engage in multi-individual processes of norm negotiation (both verbal and nonverbal) that allow gains from cooper-

ation and minimize risks of free-riding. The framework we proposed, however, is qualitative and far from fully specified. It clearly requires formal models to evaluate its plausibility.

We suspect that given the absence of state controls, the systems of exchange and cooperation found in traditional societies would not be stable without the complex web of kinship connections characterizing their residential groups. Those connections have two effects. First, as discussed earlier in this chapter, they reduce conflicts of interest between individuals and families. In fact, the marriage alliances between families (observed and commented on since the earliest days of anthropology) may be a way to minimize such conflicts of interest through the production of descendents sharing genes from both sets of families. Second, kinship connections lower the variance in the payoffs associated with norms of sharing and cooperation. For example, norms that allocate larger shares to families with more children to feed may be disadvantageous for individuals in small families, but because, members of small families are likely to have close kin (nieces, nephews, brothers, sisters, and grandchildren) in large families, the total net results of the norm for their genetic lineage may be positive. Since most other species that have elaborate systems of resource sharing and cooperation—such as social insects and group-hunting predators—organize cooperation along kinship lines, it is likely that kinship played an important role in the evolution of cooperation in humans. Models of multi-individual norm negotiation with and without kinship will be particularly useful in evaluating this intuition.

In part III of this chapter, we suggested that norms of sharing and cooperation would reflect the ecology of subsistence, as well as the associated variability in the gains from cooperation and possibilities for free-riding. However, it is possible that similar ecologies may result in very different equilibria, depending upon historical conditions and perhaps even essentially random perturbations. Formal models would also be useful for evaluating this possibility. If multiple equilibria are possible, then cultural or trait group selection may determine which equilibria come to dominate over time. Given the kinship relations organizing the formation of groups in traditional societies, cultural and genetic selection among groups and lineages may occur simultaneously.

Finally, informal observation (and the results of behavioral genetics studies) suggest that there may be significant individual differences within groups in terms of free-riding and obedience to group norms.
The existence of varying degrees of free-riding by individual members of social groups may be an inevitable outcome of cooperative norms that can only be partially enforced. The optimal amount of effort allocated to police free-riding may itself be subject to negotiation, as are allocations to law enforcement in state societies.

This chapter represents a first step in developing a multiindividual approach to cooperation among traditional human societies and to the psychology that underlies it. Our hope is that this paper will help stimulate the development of formal analyses of those processes.

Notes

1. Chimpanzee mothers do share some difficult-to-acquire solid foods with weaned offspring (Silk 1979), but chimpanzee young are largely self-sufficient after they are weaned.

2. While computer simulations reveal that significant correlations between individuals in amounts given and received are possible when tolerated theft is the sole cause of food sharing, correlations greater than 0.2 were only found in groups of fewer individuals than was common in the above groups.

3. The consumption and production of women is not included in this calculation since, on average, women produce just enough to support their own consumption or a bit less.

4. Among the Machiguenga of Yemiuwato, the best hunter produced more than half of the meat for the whole village over a year period.

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