

*Neuroeconomics*

# Social neuroeconomics: the neural circuitry of social preferences

Ernst Fehr<sup>1</sup> and Colin F. Camerer<sup>2</sup><sup>1</sup> University of Zurich, Institute for Empirical Research in Economics, Zurich, CH-8006, Switzerland<sup>2</sup> California Institute of Technology, Pasadena, CA, 91125, USA

**Combining the methods of neuroscience and economics generates powerful tools for studying the brain processes behind human social interaction. We argue that hedonic interpretations of theories of social preferences provide a useful framework that generates interesting predictions and helps interpret brain activations involved in altruistic, fair and trusting behaviors. These behaviors are consistently associated with activation in reward-related brain areas, such as the striatum, and with prefrontal activity implicated in cognitive control, the processing of emotions, and integration of benefits and costs, consistent with resolution of a conflict between self-interest and other-regarding motives.**

## Introduction

As in behaviorist psychology, the long-standing tradition in economic theory has been to treat preferences and beliefs as impossible or difficult to observe directly; instead, their effects were thought to be only revealed by direct choices. The emerging neuroeconomic approach [1–4] rejects the premise of unobservability, and seeks a microfoundation of social and economic activity in neural circuitry, using functional magnetic resonance imaging (fMRI), transcranial magnetic stimulation (TMS), pharmacological interventions and other techniques. The neuroeconomic approach hopes to unify mechanistic, mathematical and behavioral (choice-based) measures and constructs. Byproducts of such an ambitious program might include better understanding of individual differences and development over the human lifecycle (including disorders and expertise), insights into the effects of direct and social learning, empirical discipline of evolutionary modeling, and advice for how economic rules and institutions can be designed so that people react to rules in a socially efficient way.

In this review, we discuss the neural circuitry involved in altruistic, fair and trusting behaviors. Traditional economic analyses generally make the simplifying assumption that people are exclusively self-regarding, but there is now a large body of experimental evidence [5,6] indicating that many people exhibit social preferences, that is, their preferred choices are based on a positive or negative concern for the welfare of others, and on what other players believe about them. Social neuroeconomics tries

to understand the brain processes that govern these regular deviations from purely self-interested behavior. Social neuroeconomics combines the tools of social cognitive neuroscience [7–9] with well-structured tasks taken from economic theory (Boxes 1 and 2). These tasks come equipped with benchmark theoretical predictions about rational play and social efficiency of outcomes, which are useful for interpreting the results and cumulating regularity across studies.

Roughly speaking, there are two viewpoints in economic and biological sciences about why pro-social behaviors occur. One view is that behavior in one-shot anonymous games indicates a reflexive behavior that is highly adapted for repeated interactions in which immediate pro-social behavior earns future benefits. In this view, pro-sociality in one-shot games results from bounds on rationality in fully responding to changes in economic structure [10]. The other view is that pro-social behavior reflects robust social preferences for treating others generously or reciprocally, and those preferences are similar to preferences for other kinds of primary and secondary rewards (the ‘reward interpretation’; Box 2).

This paper focuses on recent studies that provide some tentative evidence of neural activity, which might eventually distinguish these two broad viewpoints. Some of the questions asked in social neuroeconomics include: what are the neural networks and the motivational forces behind charitable donations, rejections in ultimatum bargaining games, punishment of greedy behavior in third party punishment games, or decisions to trust and to reciprocate trust altruistically? To what extent do emotional and rewarding factors play a role here, and how do they interact with the human ability for rational deliberation? (Box 3).

## Theories of social preferences and the brain

Economic theories of social preferences [11–18] model the motivational forces driving the deviations from economic self-interest in a precise way. In theories of reciprocal fairness [12,17,18], for example, players are assumed to positively value kind intentions, and to negatively value hostile intentions, of other players. Thus, if player A reduces B’s payoff to his own benefit, a reciprocal player B will punish A, whereas if bad luck led to a redistribution of income from B to A, a reciprocal player B will not punish [19]. By contrast, if a player is motivated by inequity

Corresponding author: Fehr, E. (efehr@iew.uzh.ch).  
Available online 2 October 2007.

### Box 1. Measuring social preferences with games

Experimental games enable measurement of how much of their own economic payoff players are willing to sacrifice to increase or decrease the payoffs of others [5,6]. These games are typically played one-shot, with anonymous partners and with real monetary stakes. They provide a solid collection of empirical regularities from which the study of neural activity can proceed.

In a 'dictator' game [62,63], one player – the dictator – is given a sum of money that he can allocate between herself and another player – the recipient. Dictator allocations are a mixture of 50% offers and 0% offers (i.e. the dictator keeps everything), and a few offers in between 50 and 0%, but the allocations are sensitive to details of how the game is described [6], the dictator's knowledge of who the recipient is [64] and whether the recipient knows that she is part of a dictator game [65].

In an ultimatum game, the recipient can reject the proposed allocation [66]. If she rejects it both players receive nothing. Rejections are evidence of negative reciprocity [12], the motive to punish players who have treated you unfairly, or inequity aversion [14], a distaste for unfair outcomes. The strength of these motives can be measured by how much a recipient loses by rejecting a proposed allocation. Offers of less than 20% are rejected about half the time; proposers seem to anticipate these rejections and consequently offer around 40% on average. Cross-cultural studies, however, show that across small-scale societies ultimatum offers are more generous when cooperative activity and market trade are more common [67].

In a third party punishment game two players, the dictator A and the recipient B, participate in a dictator game [68]. A third player, the potential punisher C, observes how much A gives to B; then C can spend a proportion of his endowment on punishing A. This game measures to what extent 'impartial' and 'unaffected' third parties are willing to stick up for other players at their own expense, enforcing a sharing norm by punishing greedy dictators. Between 50 and 60% of the third parties punish selfish deviations from the equal split suggesting that giving less than 50% in the dictator game violates a fairness norm. In principle, the third party punishment option can be used to measure economic willingness to punish violation of any social norm (e.g. a violation of etiquette, breaking a taboo or making a linguistic slur).

In a trust game [69,70], two players, A and B, each have an initial endowment. First, A decides whether to keep his endowment or to send it to B. Then B observes A's action and decides whether to keep the amount she received or share some if it with A. The experimenter doubles or triples A's transfer, so that both players are better off collectively if A transfers money and B sends back a sufficient amount. This situation mimics a sequential economic exchange in the absence of contract enforcement institutions. B has a strong incentive to keep all the money and repay none to A; if A anticipates this behavior, however, there is little reason to transfer so a chance for mutual gain is lost. Empirically, A's invest about half of their endowment and B's repay about as much as player A invested [6]. Player A's invest less than they do in risky choices with chance outcomes, however, which indicates a pure aversion to social betrayal and inequality [46].

In a public goods game [5,6,71], which represents a generalization of the prisoners' dilemma (PD) game (Box 2), players have a token endowment they can simultaneously invest in any proportion to a private project or a public project. Investment into the public project maximizes the aggregate earnings of the group but each individual can gain more from investing into the private rather than the public project. Typically, players begin by investing half their tokens on average (many invest either all or none). When the game is repeated over time, with feedback at the end of each decision period, investments decline until only a small fraction (~10%) of the players invest anything. When players are enabled to also punish other players at a cost to themselves, many players who invested punish the players who did not invest, which encourages investment and leads players close to the efficient solution in which everyone invests [72].

aversion [14], that is, a dislike of unequal outcomes *per se*, then bad luck will induce player B to take action to redistribute income [20]. Likewise, some theories postulate an individual's desire to increase the economic welfare of the group they belong to [15,16], to experience a warm glow from altruistic giving to worthy causes [11] or to maintain a positive social image [21].

Theories of social preferences are based on the concept of decision utility [22]. A decision utility is a numerical measure that is thought to underlie observed behavior (e.g. the action chosen from a set of choices is inferred to have the highest numerical decision utility). Decision utility can, in principle, be distinguished from (i) experienced utility, which is the hedonic experience associated with the consumption of a good or an event; and from (ii) anticipated utility, which is the anticipation of experienced utility at the time of decision-making. One of the central questions in social neuroeconomics, which recent studies address, is how the brain constructs decision utilities when a person's behavior reflects their own rewards but is also governed by competing motives, such as warm glow altruism, reciprocity or inequity aversion. This general question implies a host of other important questions such as: is self-interest a primary motive that needs to be constrained by appropriate inhibitory machinery? If so, which brain circuitry is involved in these inhibitory processes? To what extent are these processes related to emotion regulation? Are deviations from economic self-interest partly governed by positive hedonic consequences associated with non-selfish behaviors and, if so, are these complex social rewards represented in the striatum and the orbitofrontal cortex (OFC) like primary or monetary rewards [23,24], or do they rely on different neural circuitries?

### Social preferences and reward circuitry

Theories of reciprocity and inequity aversion imply that subjects prefer the mutual cooperation outcome over the unilateral defection outcome in the canonical prisoners' dilemma (PD) game although unilateral defection leads to a higher economic payoff (Box 2, Tables I, II and III). Although these theories do not make assumptions about the hedonic processes associated with fairness-related behaviors (because they rely on inferred decision utilities), a plausible interpretation of these theories is that subjects in fact derive higher hedonic value from the mutual cooperation outcome [25]. Indeed, there is questionnaire evidence (M Kosfeld *et al.*, unpublished) supporting the view that mutual cooperation in social exchanges has special subjective value, beyond the value that is associated with monetary earnings (Box 2, Table IV). Therefore, a natural question is whether we can find neural traces of the special reward value of the mutual cooperation outcome. Two neuroimaging studies [26,27] report activation in the ventral striatum when subjects experience mutual cooperation with a human partner compared with mutual cooperation with a computer partner. Despite the fact that the monetary gain is identical in both situations, mutual cooperation with a human partner is associated with higher striatal activity, consistent with the reward hypothesis, given substantial evidence from other studies with primary and secondary rewards that the striatum is activated by anticipated reward.

## Box 2. Analysis of a prisoner's dilemma with social preferences

In the prisoners' dilemma (PD) each of two players makes one of two choices: cooperate or defect. The PD can either be played sequentially, where one player moves before the other, or simultaneously, where both players choose without knowing what the other one does.

**Table I. Representation of PD in terms of material payoff**

	Cooperate (C)	Defect (D)
Cooperate (C)	4, 4	0, 5
Defect (D)	5, 0	1, 1

The PD can be thought of as a paradigm for any kind of exchange that is not enforced by third parties. Assume for instance that both player A and B possess a good that they value with 1; however, they both value the other player's good with 4 so that exchanging the goods is beneficial for both. In this case, cooperation means sending the good to the other player whereas defection means keeping one's good. In case of mutual cooperation (exchange) both players receive an economic payoff of 4, in case of mutual defection both receive 1. If player A (the row player) defects and player B (the column player) cooperates, A receives an economic payoff of 5 (the value of his own good plus the value of the other's good) whereas B receives 0. See lower left corner of Table I; the first number in each cell is A's payoff, the second number is B's payoff. Regardless of what B does, it is always in the self-interest of A to defect. The same holds for B. Thus, the unique equilibrium outcome in the game is (defect, defect). However, if both players defect they are worse off than if they both cooperate, hence the dilemma.

**Table II. Utility representation of PD if players are inequity averse**

	Cooperate (C)	Defect (D)
Cooperate (C)	4, 4	$0 - 5\alpha, 5 - 5\beta$
Defect (D)	$5 - 5\beta, 0 - 5\alpha$	1, 1

If both players have a strong enough preference for reciprocity [12] or if they are inequity averse [14] their subjective preferences transform the game. In case of inequity aversion a player suffers from receiving less than the other with parameter  $\alpha_i$  (envy), and also from receiving more than the other with parameter  $\beta_i$  (compassion). An inequity averse player  $i$ 's subjective payoff  $U_i$  is a function of her own economic payoff  $x_i$  and of the payoff differences ( $x_j - x_i$ ) between the two players:  $U_i(x) = x_i - \alpha_i(x_j - x_i)$  if player  $i$  is worse off than player  $j$  ( $x_j - x_i > 0$ ), and  $U_i(x) = x_i - \beta_i(x_i - x_j)$  if player  $i$  is better off than player  $j$  ( $x_i - x_j > 0$ ). Inequity aversion makes unilateral defection less attractive by reducing the subjective payoff from 5 to  $(5 - 5\beta)$  whereas being the victim of the other player's unilateral defection is particularly painful because it reduces the subjective payoff from 0 to  $-5\alpha$  (See Table II).

**Table III. Utility representation of PD if players are inequity averse with parameters  $\alpha = 1$  and  $\beta = 0.5$**

	Cooperate (C)	Defect (D)
Cooperate (C)	4, 4	-5, 2.5
Defect (D)	2.5, -5	1, 1

Social preference theories also predict that subjects prefer to punish unfair behavior such as defection in public good and PD games (Box 2, Table V) because leaving an unfair act unpunished is associated with higher disutility than bearing the cost of punishing an unfair act. In this view, it is natural to hypothesize that the act of punishing defection involves higher activation of reward circuitry. A study using positron-emission topography (PET) [28] examined this hypothesis in the context of a PD game with a punishment opportunity similar to the one described in Box 2, Table V. This study showed that the dorsal striatum

Table III shows a representation of subjective payoffs for the special case of  $\alpha = 1$  and  $\beta = 0.5$ . Here, we can see that an inequity averse player A values the mutual cooperation outcome with 4 whereas the unilateral defection outcome is only valued with 2.5. Thus, if a player believes that the other player cooperates the player subjectively prefers to also cooperate, rendering mutual cooperation an equilibrium. However, mutual defection also remains an equilibrium: if an inequity averse player believes that the other player defects he or she prefers to defect too.

**Table IV. Player A's actual average ranking of outcomes in the PD**

	Cooperate (C)	Defect (D)
Cooperate (C)	3.2	1.6
Defect (D)	2.7	2.5

Subjects' ordinal ranking of the four outcomes in a PD game similar to the one in Table I were elicited by M Kosfeld *et al.* (unpublished). Subjects had to assign a rank between 1 and 4 to each cell of a PD. 4 represented the most highly valued outcome and 1 the least valued outcome. The numbers in the table represent the outcome from the perspective of player A only. On average, subjects valued the mutual cooperation outcome highest (3.2) and the other player's unilateral defection the least (1.6). The hypothesized ordinal ranking of outcomes by an inequity averse player A in Table III is identical to the actual ordinal ranking of outcomes in Table IV. In particular, the average player A (Table IV) prefers cooperation if she believes the opponent cooperates, and prefers defection if she believes the opponent defects.

**Table V. Utility representation if an inequity averse player A ( $\alpha = 1, \beta = 0.5$ ) punishes a selfish player B for defection**

	Cooperate (C)	Defect (D)
Cooperate (C)	4, 4	-2, 0
Defect (D)	2.5, 0	1, 1

Suppose now that the PD is played sequentially – A first chooses, then B chooses, knowing what A did. Then, A observes whether B cooperated or defected. After this A has the chance to punish B at a cost to himself. An inequity averse player A will never punish B in case of mutual cooperation or mutual defection because the players' payoffs are equal in these cases. However, he might be willing to punish if B defected unilaterally. Suppose, for example, that A can spend 1 money unit on punishment such that B's income is reduced by 5 money units, causing a material payoff distribution of  $(-1, 0)$  and a utility of  $U_A = -1 - \alpha(1 - 0) = -1 - \alpha$ . If A does not punish B the material payoff distribution is  $(0, 5)$  and A's utility is  $U_A = 0 - \alpha(5 - 0) = 0 - 5\alpha$ . In Table V we assumed  $\alpha = 1$ , implying that A's subjective payoff is  $-5$  if he does not punish whereas if he punishes it is  $-2$ . But for any  $\alpha \geq 1/4$  player A is subjectively better off if he punishes B. Note also that in this case only mutual cooperation is part of an equilibrium because a rational player B anticipates that A punishes and hence, it is not in the self-interest of B to defect.

(caudate nucleus) is strongly activated in the contrast between a real punishment condition (in which the assignment of punishment points hurt the defector in economic terms) and a symbolic punishment condition (in which the assignment of punishment points did not reduce the defector's economic payoff). In another study [29] subjects first played a sequential PD with (confederate) fair and unfair opponents. The focal subjects were then scanned (using fMRI) when a slight pain – an electrical shock – was administered either to themselves or to confederate partners who behaved fairly or unfairly. Both men and women

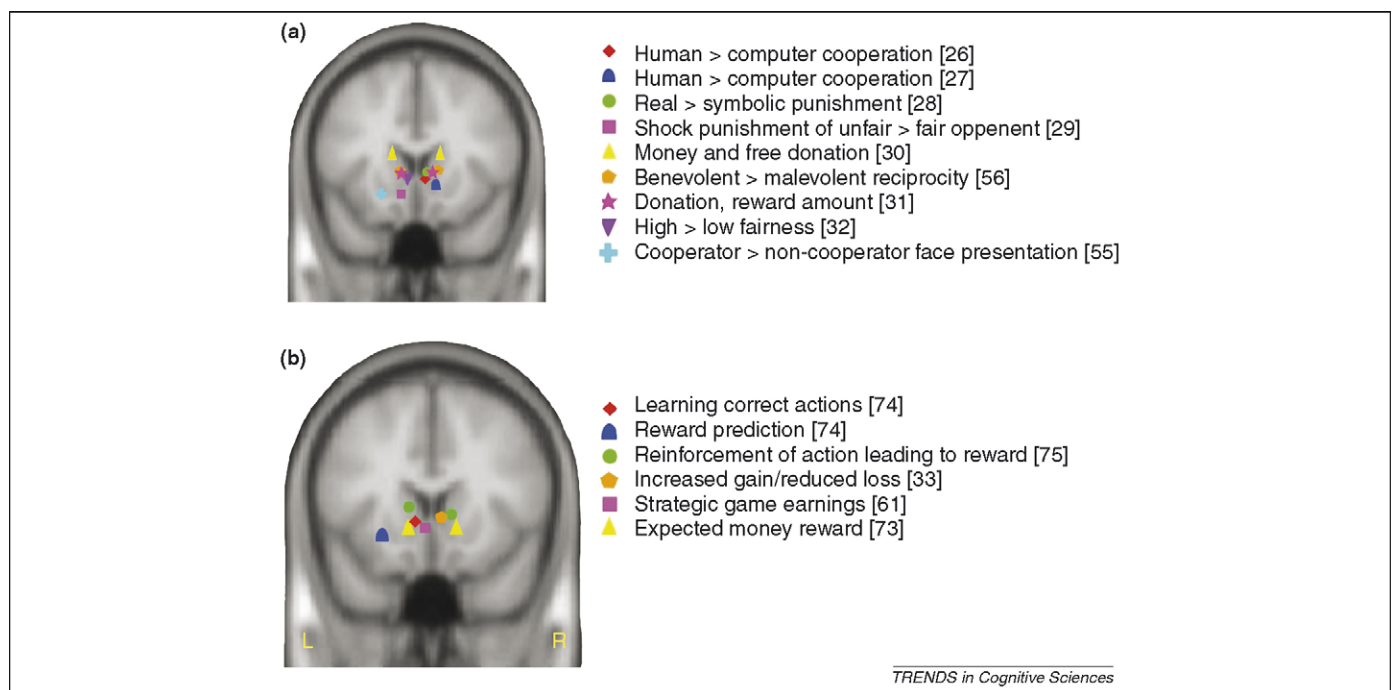
### Box 3. Questions for future research

- How does the brain relate decision utility to anticipated and experienced utility?
- What is the relationship between moral emotions, such as guilt and shame, and moral behavior? How can we measure and induce these moral emotions, and which behaviors are caused by them?
- Which behaviors, emotions and neural mechanisms of human pro-sociality are unique, and which do we share with other primates?
- What gene clusters are reliably linked to economic aspects of social behavior?
- Which computational models of brain activity predict both neural events and social behavior correctly?
- How are social disorders like autism, Asperger and Williams' syndrome, social phobias, and anti-social personality disorder linked to differences in neural activation?
- How is group membership perceived and processed neurally, and what are its implications?
- How does extensive experience (e.g. experts in negotiation or professional poker players) affect neural bases of social exchange?
- What are the neural correlates of skill in strategic interaction? How does training and experience affect neural activity?
- Many social institutions use social network connections (e.g. personal referrals) or agents (e.g. in bargaining); how does neural activity in these cases differ from personal interaction?

exhibited empathic responses in anterior cingulate and anterior insula when the fair partner received pain. However, only men report a higher desire for revenge against unfair partners, and also exhibit activation in the nucleus accumbens (NAcc) and OFC when unfair partners are shocked. Male revenge-desire ratings across subjects are also correlated with the estimate of NAcc activity,

consistent with the view that there is reward value in observing the punishment of unfair partners.

Further evidence that decisions involving social preferences are associated with activity in reward circuitry comes from fMRI studies of charitable donations [30,31] and reaction to offers in a take-it-or-leave-it ultimatum bargaining game [32]. Ventral tegmental (VTA) and striatal areas are both activated by receiving money and non-costly donations, indicating that 'giving has its own reward' [30]. Across subjects, those who made more costly donations also had more activity in the striatum. Decisions to donate, whether costly or not, activate the subgenual area, which is densely connected with mesolimbic dopaminergic and serotonergic pathways, and is implicated in social attachment mechanisms and in regulating the release of the neuromodulator oxytocin (OT) via the anterior hypothalamus. In one study, subjects are in two conditions – a forced donation and a voluntary donation condition [31]. In the forced donation condition, subjects passively observed that money is transferred to their account or to the charities account. In the voluntary condition the subjects could decide whether to accept such monetary transfers. Both in the forced and in the voluntary condition subjects reported higher satisfaction if they themselves receive more money or if the charity receives more money (controlling for the subject's cost of this transfer). Moreover, in both conditions activations in dorsal and ventral striatum are positively correlated with the money that goes to the charity and to the subjects themselves. Finally, a recent ultimatum game study [32] provides evidence suggesting that the fairness of a bargaining offer – controlling for the absolute size of the monetary gain – is associated with activations in the ventral striatum. The



**Figure 1.** Parallelism of rewards for oneself and for others: Brain areas commonly activated in (a) nine studies of social reward [26–32,55,56], and (b) a sample of six studies of learning and anticipated own monetary reward [33,61,73–75]. (All are projected onto Talairach coordinate  $y = 15$ ; voxels of peak activation in original studies range from  $y = 4$  to  $y = 24$ ). Numbers in square brackets are reference numbers.

same dollar bargaining offer of, say \$5, elicits higher striatal activation if the offer represents a fair share (say 50%) of the amount which is being bargained over, compared to when that dollar offer represents a small share (say, only 15%).

The activations observed in these studies and several others indicate that social rewards commonly activate the dorsal or ventral striatum (Figure 1a). There is substantial overlap between these areas of activation and activation observed in studies of reinforcement learning or anticipated money reward (Figure 1b). This overlap is consistent with the hypothesis that social preferences are similar to preferences for one's own rewards in terms of neural activation.

### Do activations in reward circuitry predict choices?

The above evidence is consistent with the view that costly pro-social acts of charitable donation and punishment of unfair behaviors are both rewarding. However, the hedonic interpretation of social preference theories also implies that such acts occur because they are rewarding. Evidence for causality is also important for moving from correlation to causality, and because some studies suggest monetary gains and losses are both fully processed by a unitary system, centered on the striatum [33,34]. If this unitary activity holds more generally for positively and negatively valenced goods, the mere fact that studies show higher fMRI blood oxygen-dependent level (BOLD) responses for costly altruistic acts might indicate a costly experience rather than a rewarding one. But, if it could be shown that higher activations in the striatum imply a higher willingness to act altruistically, the case for the reward interpretation would be strengthened considerably (because it is implausible to observe this relation between striatum activation and altruistic acts if striatum activation represents the cost of the act rather than its reward value).

Neuroimaging data do not enable causal inferences, but it is possible to move towards causality by predicting from neural activity in one treatment to choice behavior in another treatment ('out of treatment' forecasting). For example, individual differences in caudate nucleus activation when punishment is costless for the punisher can be used to predict how much individuals actually pay for punishment when it is not costless [28]. Likewise, individual differences in striatal activity in the condition in which donations are forced can be used to predict subjects' willingness to donate money to charities in the condition in which donations are voluntary [31] (Figure 2). These results further support the reward interpretation of social preferences, which, in turn, provides support for the hypothesis of a common neural currency of socially preferred and other primary and secondary rewards [35].

### The role of the prefrontal cortex (PFC) in decisions involving social preferences

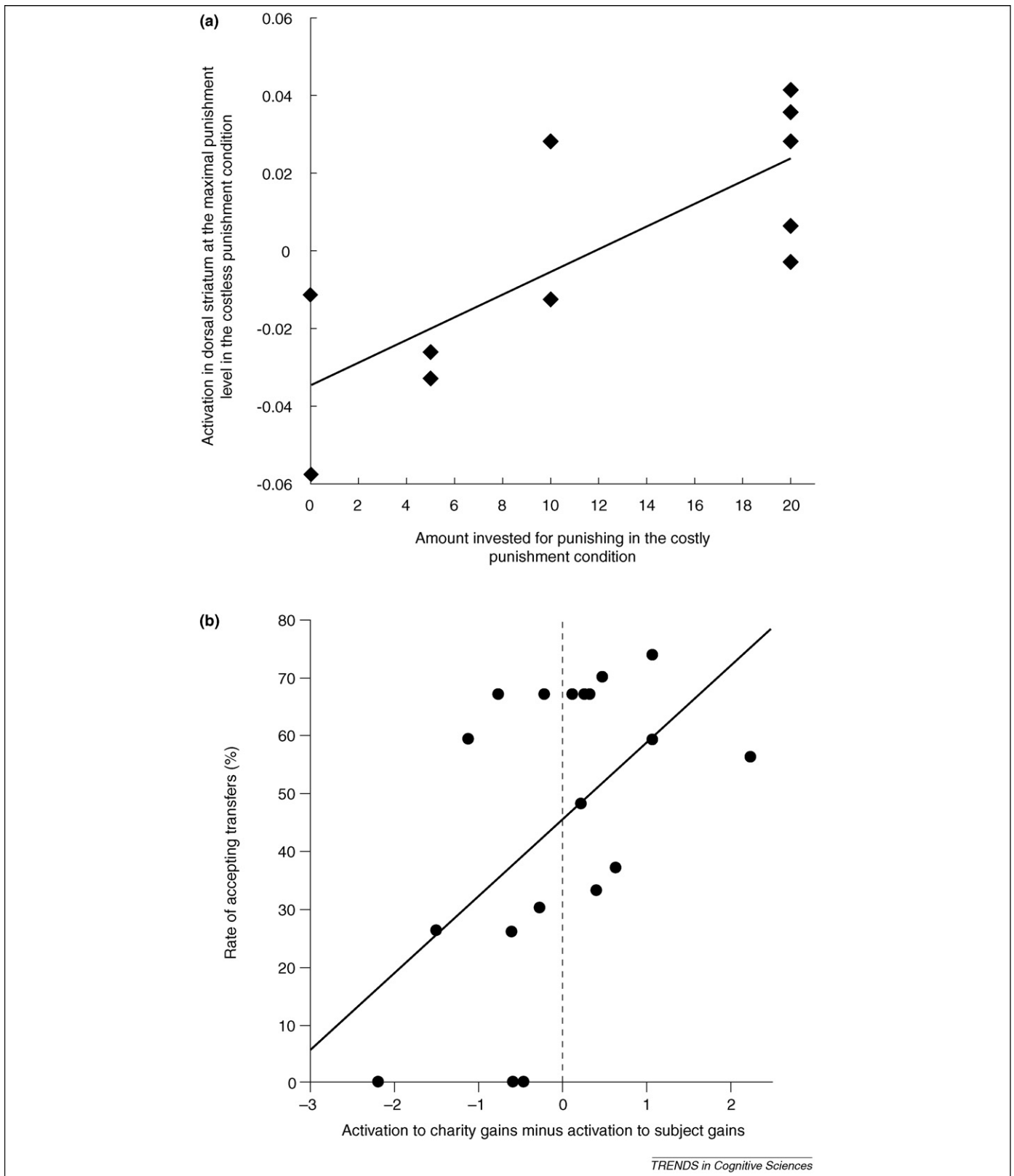
If people have social preferences the brain must compare social motives and economic self-interest and resolve conflict between them. Several studies indicate that the PFC plays a decisive role in such conflict resolution. For example, in the contrast between costly punishment condition and

costless punishment of players who behaved unfairly, the ventromedial PFC (VMPFC) Brodmann areas 10 and 11 (BA 10, 11) has been implicated [28], consistent with the hypothesis that this area is involved in the integration of separate benefits and costs in the pursuit of behavioral goals [36]. The crucial role of VMPFC in decisions involving social preferences is also supported by evidence [37] that subjects with brain lesions in VMPFC reject ultimatum game offers more frequently, suggesting that the cost of rejecting positive offers has less weight in the decision process if VMPFC is impaired. Finally, in charitable donations [30] the contrast between altruistic decisions involving costs and no costs also activated the VMPFC (BA 10, 11, 32) and the dorsal anterior cingulate cortex (ACC). Because the ACC is thought to play a key role in conflict monitoring [38], activity in this region is consistent with the existence of a tradeoff between self-interest and pro-social motives.

The role of the VMPFC in decisions involving costly altruism is also interesting because of related activation in this region in other studies. The VMPFC is involved in emotional processing and moral judgment [39,40], and in integrating the value of consumer products and their prices [41]. Lesions to VMPFC are also associated with poor choices in various situations that require integrating costs and benefits [42,43]. These studies and those on the VMPFC's role in expression of social preference suggest a general role in integrating emotional feelings about costs and benefits, regardless of whether these choices involve economic consumption goods or 'non-economic' goods, such as the subjective value of acting altruistically.

Two neuroimaging studies [32,44] suggest that the dorsolateral (DLPFC) and ventrolateral (VLPFC) prefrontal cortex also play an important role in the processing of decisions involving social preferences. These studies examined the neural circuitry involved in the recipient's behavior in an ultimatum game where the rejection of low positive offers involves a motivational conflict between fairness and economic self-interest. The first study reports differential activation of bilateral DLPFC, bilateral anterior insula (AI), as well as ACC, in the contrast between unfair and fair offers [44]. In addition, the higher the activation of right AI the more probable a subject rejects an unfair offer suggesting that AI activation might be related to the degree of emotional resentment of unfair offers. Owing to the role of ACC in conflict monitoring [38], the activation of ACC in this task might reflect the motivational conflict between fairness and self-interest when facing unfair offers. Finally, the DLPFC activation might represent the cognitive control of the emotional impulse to reject unfair offers. The second study also finds that AI is more active during rejected trials. In addition, the right VLPFC is more activated (relative to a resting baseline) when unfair offers are accepted, which might indicate that this region downregulates the resentment associated with unfair offers [32].

The interpretation that DLPFC activity represents the cognitive control of the impulse to reject implies that interfering or disrupting DLPFC activity reduces the control of the impulse and should, thus, increase the rejection rate. Knoch *et al.* [45] examined this hypothesis by reducing the activation in right and left DLPFC with



**Figure 2.** Predicting the frequency of subjects' altruistic choices with brain activations 'out of treatment'. **(a)** Subjects punished maximally when punishment of defection was costless but they still exhibit different activations in the dorsal striatum (y-axis). These activations predict how much subjects spend on punishment when it is costly (x-axis) [28]. Reproduced with permission from [28] **(b)** The individual differences in striatal activations between the condition where subjects just observe the charity receiving money (i.e. they cannot decide) and the condition where they themselves receive money can be interpreted as a measure of hedonic altruism (x-axis). Consistent with this interpretation, subjects with a higher difference in these activations accept transfers to charities more often when they have the freedom to do so (y-axis) [31]. Figures reproduced with permission from AAAS [28,31].

low-frequency TMS. Surprisingly, the study found that TMS of right DLPFC increases the acceptance rate of unfair offers relative to a placebo stimulation (from 9% to 44%), whereas TMS of left DLPFC did not affect behavior significantly. This finding suggests that right DLPFC is not involved in controlling the impulse to reject unfair offers but in controlling the impulse that pushes subjects towards accepting unfair offers, that is, with controlling economic self-interest. Interestingly, the disruption of right DLPFC only affects subjects' fairness-related behaviors but not their fairness judgments, that is, they still judge low offers as unfair, but they nevertheless accept them more frequently and more quickly.

### Trust, reputation and social preferences

Social preference models predict that trusting other individuals, by making investments that might not be repaid, is not just a decision involving monetary risk. Reciprocal and inequity averse subjects derive a special disutility from betrayal of trust, along with the associated economic loss, which is consistent with behavioral studies [46] indicating a pure aversion to social betrayal. The first evidence that the brain distinguishes between social trust and monetary risk-taking comes from [47] who infused the synthetic neuropeptide oxytocin (OT) intranasally to players in a trust game. OT-infused players were more trusting than a placebo control group, although OT-players' beliefs about the chances of being repaid were not higher and OT did not affect risk-taking in a pure risk condition. Thus, OT seems to limit the fear of betrayal in social interactions, consistent with animal evidence that it inhibits defensive behavior and facilitates maternal behavior and pair-bonding [48]. The hypothesis that the fear-of-betrayal-reducing effect of oxytocin might be a result of a reduced activation of the amygdala is consistent with a study [49] showing that OT dampens amygdala activity and its connections to the brainstem if subjects view emotionally arousing pictures. Amygdala involvement has been shown to occur in assessing the trustworthiness of faces [50,51] and the processing of ambiguous events [52], which both have social implications.

Because trust decisions are also likely to involve perspective taking, they should also activate areas implicated in theory of mind tasks, such as the paracingulate cortex and the posterior superior temporal sulcus (pSTS) [53]. One of the earliest neuroeconomic studies [54] reports activation of the paracingulate in a trust game when subjects play against another person compared to a computerized opponent. Another study found that pSTS is activated simply by showing the faces of intentional cooperators compared to nonintentional agents [29].

In repeated trust games a player learns about his opponent's choices so that the opponent acquires a reputation. In game theory, this reputation is defined as the subjective probability that the opponent is the type of player who prefers to reciprocate trust. In this approach, players' preferences and their subjective beliefs are distinct concepts and a rational player's beliefs are not colored by his preferences or his emotions towards the opponent. It is interesting to examine whether the brain also makes this distinction, that is, whether the neural networks

involved in hedonic preferences and emotional processing are distinct from the networks involved in assessing the opponent's reputation or whether there is substantial overlap in these neural networks.

Preliminary evidence suggests that the latter is likely to be true. In one study [55] players faced a series of cooperative and noncooperative opponents in a sequential PD game. The authors found that simply displaying the faces of cooperative partners (relative to neutral faces) in a subsequent gender-assessment task activated striatal and emotion-related areas, such as the amygdala, the insula and the putamen. This suggests that a trustworthy person's face automatically triggers emotions and reward expectations, as if simply seeing another person's face activates its representation as a future exchange value.

The importance of the striatum in learning the opponent's trustworthiness has been demonstrated by two other studies. In one study, the activity in caudate nucleus signals whether the other player reciprocates an earlier move [56]. A further study using the same trust-game paradigm showed specializations in the cingulate for encoding decisions of others and oneself [57]. In a third study, trustors repeatedly face three partners whose (fictional) profiles make them seem morally good, bad or neutral (instilling a prior belief about trustworthiness) [58]. By design, all three fictional partners repay in the trust game with the same frequency. During the outcome phase the caudate nucleus activates more strongly for repayment outcomes from the neutral partner, but not from the other partners, presumably because the neutral partner represents unpredictable outcomes and there is more to learn.

### Conclusions and research directions

In this review, we showed how theories of social preferences guided the conduct of neuroeconomic experiments and the interpretation of the resulting brain data. One emerging theme of the studies reviewed above is that social reward activates circuitry that overlaps, to a surprising degree, with circuitry that anticipates and represents other types of rewards. These studies reinforce the idea that social preferences for donating money, rejecting unfair offers, trusting others and punishing those who violate norms, are genuine expressions of preference. The social rewards are traded off with subjects' economic self-interest and the DLPFC and the VMPFC are likely to be crucially involved in the balancing of competing rewards. These processes can also be altered by treatments like oxytocin infusion and TMS disruption, actually changing behavior in ways that are consistent with hypotheses derived from fMRI.

Economics and other social sciences might benefit from social neuroeconomics because of the potentially unifying force of neural data for choice-based approaches. Most of economics assumes, for example, that beliefs about other people's behavior are based on a rational assessment of the available information; they are neither directly affected by preferences nor are they disturbed by emotions. However, beliefs about other people's trustworthiness are strongly affected by reward and emotion circuitry [55]. This study

also creates a paradigm for exploring how rapidly reputations increase and decrease, and whether good reputations built up by particular players in racial, gender or class-based social groups might generalize to new players in those same groups. This kind of group-based neural generalization of expected reward could be important in understanding the powerful role of social networks and physical cues in labor market discrimination, for example.

Another example illustrating the potential of neuroeconomics comes from recalling the economist's concepts of risk preferences, time preferences (the willingness to postpone consumption) and social preferences. Most economic analyses treat these as separate types of preference. However, suppose all three types of preference share some common neural circuitry for controlling automatic emotional impulses (based on fear, temptation and selfishness, respectively), by integrating all the costs and benefits of choices. If there is such a shared basis of preference, it is important for economists and other social scientists to understand, and this can be best established by data from imaging, lesion patient studies and other neuroscientific measures. For example, studies by Knoch *et al.* [45,59,60] show that disruption of right, but not left, DLPFC with TMS increases both risk-taking in choice tasks and self-interested choices in ultimatum games.

Future studies should exploit the wide range of tools available to neuroscientists, using multiple measures at the same time (e.g. hormone measurement or TMS and fMRI), combined with the parametric value and predictions about more complex games. Focusing on neural bases also opens up research directions which are, perhaps surprisingly, unexplored in economics. For example, the standard analysis in game theory assumes that players are 'in equilibrium' (i.e. they correctly anticipate, from introspection or learning, what others are planning to do). In equilibrium analysis there is no room for differences in strategic skill. Yet in one study skill varied systematically across players (as measured by belief accuracy and differences in earnings) [61]. When making choices, more skilled players showed more activity in the ventral striatum and precuneus (as if they anticipated higher money rewards) and less skilled players showed more activity in the insula (as if they were feeling discomfort from strategic uncertainty).

These directions might eventually provide a biological basis for a mathematical characterization of social exchange that is rooted in neural details but can also make predictions about activity in strategic interaction and market trading, and about how behavior can change when causally manipulated by pharmacology, TMS and other tools.

## References

- Glimcher, P.W. and Rustichini, A. (2004) Neuroeconomics: the consilience of brain and decision. *Science* 306, 447–452
- Camerer, C. *et al.* (2005) Neuroeconomics: how neuroscience can inform economics. *J. Econ. Lit.* 43, 9–64
- Fehr, E. *et al.* (2005) Neuroeconomic foundations of trust and social preferences: initial evidence. *Am. Econ. Rev.* 95, 346–351
- Sanfey, A.G. *et al.* (2006) Neuroeconomics: cross-currents in research on decision-making. *Trends Cogn. Sci.* 10, 108–116
- Fehr, E. and Fischbacher, U. (2003) The nature of human altruism. *Nature* 425, 785–791
- Camerer, C.F. (2003) *Behavioral Game Theory – Experiments in Strategic Interaction*, Princeton University Press
- Adolphs, R. (2003) Cognitive neuroscience of human social behaviour. *Nat. Rev. Neurosci.* 4, 165–178
- Blakemore, S.J. *et al.* (2004) Social cognitive neuroscience: where are we heading? *Trends Cogn. Sci.* 8, 216–222
- Lieberman, M.D. (2007) Social cognitive neuroscience: a review of core processes. *Annu. Rev. Psychol.* 58, 259–289
- Samuelson, L. (2005) Foundations of human sociality: a review essay. *J. Econ. Lit.* 43, 488–497
- Andreoni, J. (1990) Impure altruism and donations to public goods: a theory of warm glow giving. *Econ. J.* 100, 464–477
- Rabin, M. (1993) Incorporating fairness into game theory and economics. *Am. Econ. Rev.* 83, 1281–1302
- Levine, D.K. (1998) Modeling altruism and spitefulness in experiments. *Rev. Econ. Dynam.* 1, 593–622
- Fehr, E. and Schmidt, K.M. (1999) A theory of fairness, competition, and cooperation. *Q. J. Econ.* 114, 817–868
- van Lange, P.A.M. (1999) The pursuit of joint outcomes and equality in outcomes: an integrative model of social value orientation. *J. Pers. Soc. Psychol.* 77, 337–349
- Charness, G. and Rabin, M. (2002) Understanding social preferences with simple tests. *Q. J. Econ.* 117, 817–869
- Dufwenberg, M. and Kirchsteiger, G. (2004) A theory of sequential reciprocity. *Games Econ. Behav.* 47, 268–298
- Falk, A. and Fischbacher, U. (2006) A theory of reciprocity. *Games Econ. Behav.* 54, 293–315
- Blount, S. (1995) When social outcomes aren't fair – the effect of causal attributions on preferences. *Organ. Behav. Hum. Decis. Process.* 63, 131–144
- Dawes, C.T. *et al.* (2007) Egalitarian motives in humans. *Nature* 446, 794–796
- Benabou, R. and Tirole, J. (2006) Incentives and prosocial behavior. *Am. Econ. Rev.* 96, 1652–1678
- Kahneman, D. (1994) New challenges to the rationality assumption. *J. Inst. Theor. Econ.* 150, 18–36
- O'Doherty, J.P. (2004) Reward representations and reward-related learning in the human brain: insights from neuroimaging. *Curr. Opin. Neurobiol.* 14, 769–776
- Knutson, B. and Cooper, J.C. (2005) Functional magnetic resonance imaging of reward prediction. *Curr. Opin. Neurol.* 18, 411–417
- Thibaut, J.W. and Kelley, H.H. (1959) *The Social Psychology of Groups*, Wiley
- Rilling, J. *et al.* (2002) A neural basis for social cooperation. *Neuron* 35, 395–405
- Rilling, J.K. *et al.* (2004) Opposing BOLD responses to reciprocated and unreciprocated altruism in putative reward pathways. *Neuroreport* 15, 2539–2543
- de Quervain, D.J. *et al.* (2004) The neural basis of altruistic punishment. *Science* 305, 1254–1258
- Singer, T. *et al.* (2006) Empathic neural responses are modulated by the perceived fairness of others. *Nature* 439, 466–469
- Moll, J. *et al.* (2006) Human fronto-mesolimbic networks guide decisions about charitable donation. *Proc. Natl. Acad. Sci. U. S. A.* 103, 15623–15628
- Harbaugh, W.T. *et al.* (2007) Neural responses to taxation and voluntary giving reveal motives for charitable donations. *Science* 316, 1622–1625
- Tabibnia, G. *et al.* The sunny side of fairness – preference for fairness activates reward circuitry. *Psychol. Sci.* (in press)
- Tom, S.M. *et al.* (2007) The neural basis of loss aversion in decision-making under risk. *Science* 315, 515–518
- Seymour, B. *et al.* (2007) Differential encoding of losses and gains in the human striatum. *J. Neurosci.* 27, 4826–4831
- Montague, P.R. and Berns, G.S. (2002) Neural economics and the biological substrates of valuation. *Neuron* 36, 265–284
- Ramani, N. and Owen, A.M. (2004) Anterior prefrontal cortex: insights into function from anatomy and neuroimaging. *Nat. Rev. Neurosci.* 5, 184–194
- Koenigs, M. and Tranel, D. (2007) Irrational economic decision-making after ventromedial prefrontal damage: evidence from the ultimatum game. *J. Neurosci.* 27, 951–956

- 38 Botvinick, M.M. *et al.* (2001) Conflict monitoring and cognitive control. *Psychol. Rev.* 108, 624–652
- 39 Moll, J. *et al.* (2005) Opinion: the neural basis of human moral cognition. *Nat. Rev. Neurosci.* 6, 799–809
- 40 Koenigs, M. *et al.* (2007) Damage to the prefrontal cortex increases utilitarian moral judgements. *Nature* 446, 908–911
- 41 Knutson, B. *et al.* (2007) Neural predictors of purchases. *Neuron* 53, 147–156
- 42 Damasio, A.R. (1995) *Descartes' Error: Emotion, Reason and the Human Brain*, Harper
- 43 Bechara, A. *et al.* (1997) Deciding advantageously before knowing the advantageous strategy. *Science* 275, 1293–1295
- 44 Sanfey, A.G. *et al.* (2003) The neural basis of economic decision-making in the ultimatum game. *Science* 300, 1755–1758
- 45 Knoch, D. *et al.* (2006) Diminishing reciprocal fairness by disrupting the right prefrontal cortex. *Science* 314, 829–832
- 46 Bohnet, I. and Zeckhauser, R. (2004) Trust, risk and betrayal. *J. Econ. Behav. Organ.* 55, 467–484
- 47 Kosfeld, M. *et al.* (2005) Oxytocin increases trust in humans. *Nature* 435, 673–676
- 48 Insel, T.R. and Young, L.J. (2001) The neurobiology of attachment. *Nat. Rev. Neurosci.* 2, 129–136
- 49 Kirsch, P. *et al.* (2005) Oxytocin modulates neural circuitry for social cognition and fear in humans. *J. Neurosci.* 25, 11489–11493
- 50 Winston, J.S. *et al.* (2002) Automatic and intentional brain responses during evaluation of trustworthiness of faces. *Nat. Neurosci.* 5, 277–283
- 51 Adolphs, R. *et al.* (2005) A mechanism for impaired fear recognition after amygdala damage. *Nature* 433, 68–72
- 52 Hsu, M. *et al.* (2005) Neural systems responding to degrees of uncertainty in human decision-making. *Science* 310, 1680–1683
- 53 Frith, U. and Frith, C.D. (2003) Development and neurophysiology of mentalizing. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 358, 459–473
- 54 McCabe, K. *et al.* (2001) A functional imaging study of cooperation in two-person reciprocal exchange. *Proc. Natl. Acad. Sci. U. S. A.* 98, 11832–11835
- 55 Singer, T. *et al.* (2004) Brain responses to the acquired moral status of faces. *Neuron* 41, 653–662
- 56 King-Casas, B. *et al.* (2005) Getting to know you: reputation and trust in a two-person economic exchange. *Science* 308, 78–83
- 57 Tomlin, D. *et al.* (2006) Agent-specific responses in the cingulate cortex during economic exchanges. *Science* 312, 1047–1050
- 58 Delgado, M.R. *et al.* (2005) Perceptions of moral character modulate the neural systems of reward during the trust game. *Nat. Neurosci.* 8, 1611–1618
- 59 Knoch, D. *et al.* (2006) Disruption of right prefrontal cortex by low-frequency repetitive transcranial magnetic stimulation induces risk-taking behavior. *J. Neurosci.* 26, 6469–6472
- 60 Knoch, D. and Fehr, E. (2007) Resisting the power of temptations: the right prefrontal cortex and self-control. *Ann. NY. Acad. Sci.* 1104, 123–134
- 61 Bhatt, M. and Camerer, C.F. (2005) Self-referential thinking and equilibrium as states of mind in games: fMRI evidence. *Games Econ. Behav.* 52, 424–459
- 62 Mikula, G. (1972) Reward allocation in dyads regarding varied performance ratio. *Z. Sozial Psychol.* 3, 126–133
- 63 Kahneman, D. *et al.* (1986) Fairness as a constraint on profit seeking: entitlements in the market. *Am. Econ. Rev.* 76, 728–741
- 64 Eckel, C. and Grossman, P. (1996) Altruism in anonymous dictator games. *Games Econ. Behav.* 16, 181–191
- 65 Dana, J. *et al.* (2006) What you don't know won't hurt me: costly (but quiet) exit in dictator games. *Organ. Behav. Hum. Decis. Process.* 100, 193–201
- 66 Güth, W. *et al.* (1982) An experimental analysis of ultimatum bargaining. *J. Econ. Behav. Organ.* 3, 367–388
- 67 Henrich, J. *et al.* (2001) In search of homo economicus: behavioral experiments in 15 small-scale societies. *Am. Econ. Rev.* 91, 73–78
- 68 Fehr, E. and Fischbacher, U. (2004) Third-party punishment and social norms. *Evol. Hum. Behav.* 25, 63–87
- 69 Camerer, C. and Weigelt, K. (1988) Experimental tests of a sequential equilibrium reputation model. *Econometrica* 56, 1–36
- 70 Berg, J. *et al.* (1995) Trust, reciprocity and social history. *Games Econ. Behav.* 10, 122–142
- 71 Ledyard, J. (1995) Public goods: a survey of experimental research. In *Handbook of Experimental Economics* (Kagel, J. and Roth, A., eds), pp. 111–194, Princeton University Press
- 72 Fehr, E. and Gächter, S. (2002) Altruistic punishment in humans. *Nature* 415, 137–140
- 73 Preuschhoff, P.K. *et al.* (2006) Neural differentiation of expected reward and risk in human subcortical structures. *Neuron* 51, 381–390
- 74 O'Doherty, J. *et al.* (2004) Dissociable roles of ventral and dorsal striatum in instrumental conditioning. *Science* 304, 452–454
- 75 Tricomi, E.M. *et al.* (2004) Modulation of caudate activity by action contingency. *Neuron* 41, 281–292

## Elsevier.com – linking scientists to new research and thinking

Designed for scientists' information needs, Elsevier.com is powered by the latest technology with customer-focused navigation and an intuitive architecture for an improved user experience and greater productivity.

The easy-to-use navigational tools and structure connect scientists with vital information – all from one entry point. Users can perform rapid and precise searches with our advanced search functionality, using the FAST technology of Scirus.com, the free science search engine. Users can define their searches by any number of criteria to pinpoint information and resources. Search by a specific author or editor, book publication date, subject area – life sciences, health sciences, physical sciences and social sciences – or by product type. Elsevier's portfolio includes more than 1800 Elsevier journals, 2200 new books every year and a range of innovative electronic products. In addition, tailored content for authors, editors and librarians provides timely news and updates on new products and services.

Elsevier is proud to be a partner with the scientific and medical community. Find out more about our mission and values at Elsevier.com. Discover how we support the scientific, technical and medical communities worldwide through partnerships with libraries and other publishers, and grant awards from The Elsevier Foundation.

As a world-leading publisher of scientific, technical and health information, Elsevier is dedicated to linking researchers and professionals to the best thinking in their fields. We offer the widest and deepest coverage in a range of media types to enhance cross-pollination of information, breakthroughs in research and discovery, and the sharing and preservation of knowledge.

**Elsevier. Building insights. Breaking boundaries.**  
**www.elsevier.com**