Neurobiology of delayed reinforcement

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British Psychological Society Annual Conference, Imperial College London 10:10–10:50 am, Saturday 17 April 2004



Delayed reinforcement: the problems.

Delayed reinforcement: the problems

- How do animals succeed in bridging delays to reinforcement?
 - ... actions are by no means always followed by their outcomes, especially at a neuronal timescale
- 'Impulsivity' refers to several, dissociable tendencies:
 - 'preparation' impulsivity failure to collect sufficient information to make a good decision
 - 'motor'/'execution' impulsivity inability to restrain actions
 - 'outcome' impulsivity impulsive choice preference for immediate, small rewards over large, delayed rewards.

• Why do some individuals exhibit abnormally impulsive choice, choosing small, immediate rewards over large, delayed rewards?

... can be considered a normal personality trait (Aristotle, 350 BC)

... but impulsive choice contributes to attention-deficit/hyperactivity disorder (ADHD), drug addiction, mania, and personality disorders

Learning with delayed reinforcement



Discrimination learning with delayed reinforcement

FIG. 3. Rate of learning as a function of delay of reward. The reciprocal \times 1000 of the number of trials to reach the level of 75 percent correct choices is plotted against the time of delay. Experimental values are represented by black dots and the smooth curve is fitted to these data.

Grice (1948)



Free-operant learning with delayed reinforcement

Dickinson, Watt & Griffiths (1992)

Signalled and unsignalled delayed reinforcement



Cues present during the delay speed up learning



FIG. 5. Learning curves for the three different groups with five sec. delay

Grice (1948)

Choice with delayed reinforcement

Temporal discounting: devaluing the future



Smaller-sooner and larger-later rewards

Would you rather have £20 now, or £40 next year? We can call it *impulsive* to choose the smaller-sooner reward, and *self-controlled* to choose the larger-later reward. Three guesses about why people are impulsive (Ainslie, 1975):

• They lack insight into the consequences of their actions

• They are aware of the consequences of their actions, but are unable to suppress some lower principle ("the devil, repetition compulsion, classical conditioning")

• They are aware of the consequences of their actions, and choose rationally according to their value system, but their values are distorted so that imminent consequences have a greater weight than remote ones — reduced value of delayed reinforcement.

Ainslie (1975)

Impulsive and self-controlled individuals discount differently



Hyperbolic temporal discounting: irrational, but true



Choosing future rewards: preference reversal



Ainslie (1975)

Pre-commitment as a means of self-control



Homer (1700 BC?) Odyssey; Waterhouse (1891) Ulysses and the Sirens

Pigeons are often impulsive (Rachlin & Green, 1972) — but they too exhibit precommitment (Ainslie, 1974; Ainslie & Herrnstein, 1981).

Steeper temporal discounting in drug addicts



Bickel et al. (1999), smokers; Madden et al. (1999), heroin addicts

Neurochemistry of choice involving delayed reinforcement

Serotonin (5HT) in impulsive choice

• Low levels of 5HT metabolites in cerebrospinal fluid associated with impulsive aggression and violence in humans (Åsberg 1976; Linnoila *et al.* 1983).

• 5HT involved in inhibition of behaviour (impulse control)? (Soubrié 1986)

Lower levels of 5HT metabolites in cerebrospinal fluid of macaques making longer/'riskier' leaps through forest canopy! (Mehlman *et al.* 1994)

In studies specifically of **impulsive choice**:

Global 5HT depletion generally promotes impulsive choice (Wogar *et al.* 1993, Richards *et al.* 1995, Bizot *et al.* 1999, Mobini *et al.* 2000).
However, not clear cut: global 5HT depletion or antagonists do not *always* promote impulsive choice (Evenden & Ryan 1996; Crean *et al.* 2002; Winstanley et al. 2003) and 5HT2 agonists promote impulsive choice (Evenden & Ryan, 1996).

Dopamine (DA) in impulsive choice

• Amphetamine and methylphenidate (Ritalin), catecholamine releasers and reuptake blockers, are effective therapies for ADHD (Bradley, 1937, and on).

The spontaneously hypertensive rat, an animal model of ADHD, has abnormal DA systems (e.g. Russell et al. 1995). Hyperdopaminergic?
Hypodopaminergic? Debated... (e.g. Zhuang et al. 2001, Seeman & Madras 2002).

• Is impulsivity in ADHD due to steeper 'temporal discounting', due to abnormal DA systems? (e.g. Sagvolden & Sergeant, 1998).

• D2 receptors promote choice of delayed rewards. The D2 antagonist raclopride and the D1+D2 antagonist flupenthixol decrease preference for delayed reinforcement; the D1 antagonist SCH23390 has no effect (Wade *et al.* 2000).

• The effects of psychostimulants are complex (pharmacologically and behaviourally)... do they promote

• self-controlled choice? (Sagvolden '92; Richards '97/'99, Wade '00, de Wit '02)

• impulsive choice? (Evenden & Ryan '96; Charrier & Thiébot '96; Logue '92)

Choice involving delayed reinforcement: typical task



Signalled and unsignalled delayed reinforcement



The cue supports choice of the large, delayed reinforcer in rats trained in its presence



Cardinal et al. (2000)

Amphetamine *cue-independently* decreased preference for the delayed reward, but *cue-dependently* increased it



Resolves some contradictions.

Neuroanatomy of delayed reinforcement: (1) choice

Stereotaxic, excitotoxic lesions...



Nucleus accumbens core (AcbC) lesions severely impaired the ability of rats to choose a delayed reward



Cardinal et al. (2001)

... even in rats that exhibit very strong preference for the large reward when it is not delayed.



Cardinal et al. (2003)

Anterior cingulate cortex (ACC) lesions, which have been shown to produced 'motor impulsivity' in the 5-choice task, had no effect upon responding for delayed rewards



Cardinal et al. (2001)



... although they might affect 'response effort' choices

Walton et al. (2002, 2003)

ACC

PrL/IL

Medial prefrontal cortex (mPFC) lesions induced an insensitivity to the task contingencies

Lesioned subjects chose the large reward *less* frequently at zero delay, and *more* frequently at long delays.



Timing deficit? Dietrich & Allen (1998)

Cardinal et al. (2001)

Lesions of the basolateral amygdala (BLA) make rats more *impulsive* in this task; lesions of the orbitofrontal cortex (OFC) make rats more *self-controlled*.



Redrawn from Winstanley et al. (in press 2004). Lesions made after training; no stimulus in delay; 1 (immediate) v. 4 (delayed) pellets.



Mobini et al. (2002). Lesions made before training; stimulus in delay; 1 versus 2 pellets.

... but OFC lesions can also have the opposite effect!

Choice depends on reinforcer magnitude as well as delay...





Orbitofrontal cortex (OFC) lesions affect **both** delay and magnitude discounting (Kheramin *et al.*, 2002). i.e. low values of Q (relative indifference between the two reinforcers, compared to normal) can also induce 'impulsive' choice

After Bradshaw & Szabadi (1992); Ho et al. (1999); Kheramin et al. (2002)

Neuroanatomy of delayed reinforcement: (2) learning

Instrumental contingencies are harder to detect with a delay

Acquisition of free-operant instrumental responding on a fixed-ratio-1 schedule

a) Zero delay



We've seen that nucleus accumbens core (AcbC) lesions impair choice of delayed reward. Is this because they can't learn the contingency when reward is delayed?

Lesions of the AcbC again...



Cardinal & Cheung (unpublished)

AcbC lesions impair instrumental acquisition only when there is a delay between action and outcome (1)



Cardinal & Cheung (unpublished)

AcbC lesions impair instrumental acquisition only when there is a delay between action and outcome (2)



Cardinal & Cheung (unpublished)

Holds true even when experienced (rather than programmed) delays are examined



What about magnitude discrimination? The matching 'law'...



Fig. 4. The relative frequency of responding to one alternative in a two-choice procedure as a function of the relative frequency of reinforcement thereon. Variable-interval schedules governed reinforcements for both alternatives. The diagonal line shows matching between the relative frequencies. From Herrnstein (1961).

Herrnstein (1961, 1970)

Two alternatives (e.g. levers) A and B. Both deliver reinforcement intermittently and somewhat unpredictable (e.g. variable interval schedule).



where **R** is response rate; **r** is (experienced) reinforcement rate

This should be a way of testing animals' sensitivity to reinforcement magnitude... For example, if the two schedules deliver at the same rate but A delivers 1 pellet per reinforcement and B delivers 4 pellets per reinforcement, animals should allocate 80% of their responses to B.

AcbC-lesioned rats *better* at magnitude discrimination?



Considerable undermatching (common: Williams, 1994). But shams and lesions were influenced by reinforcer allocation (lines not flat), and AcbC-lesioned rats were **more** influenced by this than shams (AcbC line has a significantly steeper gradient).

Consistent with studies using other techniques (e.g. Balleine & Killcross 1994, Brown & Bowman 1995).

So a 'magnitude' explanation **can't** explain the effect of AcbC lesions to produce impulsive choice. Therefore, AcbC-lesioned rats must be specifically **hypersensitive to the effects of delays.**

The 'limbic' corticostriatal circuit: delayed reinforcement



How to avoid temptation...



Pre-commitment strategies

Cues that signal the availability of the delayed outcome

Having a good amygdala/ OFC/accumbens system to help you choose (and learn with) delayed rewards?



Draper

(1909)

Supervisor, collaborators and acknowledgements

- Barry Everitt
- Tim Cheung
- Caroline Parkinson
- John Parkinson
- David Pennicott
- Trevor Robbins
- Lakmali Sugathapala

- Helen Sweet-Gossage
- David Theobald
- Catharine Winstanley
- The UK Medical Research Council
- The Wellcome Trust
- The University of Cambridge School of Clinical Medicine

