

# NIH Public Access

**Author Manuscript** 

Prog Neurobiol. Author manuscript; available in PMC 2013 June 09.

Published in final edited form as:

Prog Neurobiol. 2010 November ; 92(3): 345–369. doi:10.1016/j.pneurobio.2010.06.007.

# Toward a Neurobiology of Delusions

P.R. Corlett<sup>1,\*</sup>, J.R. Taylor<sup>1</sup>, X.-J. Wang<sup>2</sup>, P.C. Fletcher<sup>3</sup>, and J.H. Krystal<sup>1</sup>

<sup>1</sup>Department of Psychiatry, Yale University, School of Medicine, New Haven, Connecticut, USA

<sup>2</sup>Department of Neurobiology and Kavli Institute of Neuroscience, Yale University School of Medicine, New Haven, Connecticut, USA

<sup>3</sup>Department of Psychiatry, Brain Mapping Unit and Behavioural and Clinical Neurosciences Institute, University of Cambridge, School of Clinical Medicine, Cambridge

# Abstract

Delusions are the false and often incorrigible beliefs that can cause severe suffering in mental illness. We cannot yet explain them in terms of underlying neurobiological abnormalities. However, by drawing on recent advances in the biological, computational and psychological processes of reinforcement learning, memory, and perception it may be feasible to account for delusions in terms of cognition and brain function. The account focuses on a particular parameter, prediction error – the mismatch between expectation and experience – that provides a computational mechanism common to cortical hierarchies, frontostriatal circuits and the amygdala as well as parietal cortices. We suggest that delusions result from aberrations in how brain circuits specify hierarchical predictions, and how they compute and respond to prediction errors. Defects in these fundamental brain mechanisms can vitiate perception, memory, bodily agency and social learning such that individuals with delusions experience an internal and external world that healthy individuals would find difficult to comprehend. The present model attempts to provide a framework through which we can build a mechanistic and translational understanding of these puzzling symptoms.

## Keywords

Delusions; Prediction; Error; Learning; Memory; Reconsolidation; Habit

# 1. Introduction

Delusions are the extraordinary and tenacious false beliefs suffered by patients with various ailments ranging from schizophrenia (Schneider, 1959), to traumatic brain injury (Coltheart *et al.*, 2007), Alzheimer's (Flint, 1991) and Parkinson's disease (Ravina *et al.*, 2007), the ingestion of psychotogenic drugs (Corlett *et al.*, 2009a) and, less frequently, autoimmune disorders such as Morvan's syndrome (Hudson *et al.*, 2008) or potassium channel encephalopathy (Parthasarathi *et al.*, 2006). Given this range of potential diagnoses, each with its own candidate neuropathology, it is perhaps unsurprising that we have not converged upon an agreed neurobiology of delusions. Delusions are particularly hard to study because of their insidious onset and tonic nature, their conceptual rather than behavioral basis (making them difficult to study using animal models), and the absence of a coherent theoretical model. We aim to address these issues in the current review by developing a translational model of delusion formation which we believe makes delusions

<sup>\*</sup>Corresponding Author: Dr. Philip R. Corlett (BA, PhD), philip.corlett@yale.edu, Yale University School of Medicine, Department of Psychiatry, Connecticut Mental Health Centre, Abraham Ribicoff Research Facility, 34 Park Street, New Haven, CT 06519.

tractable for animal modeling, amenable to investigation with functional neuroimaging and grounded within a theoretical framework that makes testable predictions.

Our task is made more difficult when one considers the range of odd beliefs from which people suffer; fears of persecution by clandestine forces (Melo *et al.*, 2006); beliefs that televisions or newspapers are communicating a specific and personal message (Conrad, 1958b; Startup and Startup, 2005), the conviction that one's thoughts and movements are under the control of an external agent or are broadcast out loud (Schneider, 1959); an unrealistic belief in one's own fame or power (Karson, 1980; Kraeplin, 1902), that one is infested with parasites (Thiebierge, 1894) or deceased (Cotard, 1880), or the subject of a stranger's love (De Clerambault, 1942), or that family members have been replaced by imposters or even robots (Capgras, 1923).

We take a cognitive neuropsychiatric approach to delusions. That is, the starting point is to review what we understand about the healthy functioning of a particular process, e.g. familiar face recognition, before extrapolating to the disease case, when face recognition fails and delusions of misidentification form (Halligan and David, 2001). This approach has proven successful for explaining certain delusions (Ellis and Young, 1990) but not yet for delusions in general. Perhaps this is because there are difficulties defining delusions as well as deciding what they have in common (if anything) with normal, healthy beliefs (Berrios, 1991; Delespaul and van Os, 2003; Jones, 2004; Owen *et al.*, 2004). Beliefs are not easily accessible to the techniques of neuroscience which are more suited to representing states with clear experiential boundaries (Damasio, 2000). (Knobel *et al.*, 2008).

Furthermore, delusions are difficult to model in animals, given that they involve dysfunctions of what many consider uniquely human faculties like consciousness, language, reality monitoring and meta-cognition (Angrilli *et al.*, 2009; Berrios, 1991; Moritz *et al.*, 2006). Computational models of core cognitive functions (such as working memory) are being applied to gain insights into neural dysfunction in schizophrenia (Seamans and Yang, 2004; Winterer, 2006) and some are beginning to address the phenomenology of specific psychotic symptoms (Loh *et al.*, 2007), however, these models have focused on circuit mechanisms within a local area (like prefrontal cortex), they are unable to capture the content of particular symptoms which involve information processing across large networks of interacting brain regions (Fuster, 2001).

There is a need for a testable conceptual model of delusions, one that is rooted in translational cognitive neuroscience. We, and others, propose that beliefs (both normal and abnormal) arise through a combination of innate or endowed processes, learning, experience and interaction with the world (Friston, 2010). Like other forms of information, beliefs are represented in the brain through the formation and strengthening of synaptic connections between neurons, for example causal beliefs may be mediated by a strengthening of the synaptic associations between pools of neurons representing a particular cause and their counterparts representing an associated effect (Dickinson, 2001; McLaren and Dickinson, 1990; Shanks, 2010). There are neural (and hence cognitive) limits set on the range of possible connections that can be made (Kandel, 1998). The strength of those connections is modifiable such that those conveying an adaptive advantage are strengthened and those that are disadvantageous are weakened (Hebb, 1949b; Thorndike, 1911).

This set of sculpted connections is used to predict subsequent states of the internal and external world and respond adaptively (Friston, 2005b); however, should that next state be surprising, novel or uncertain new learning is required (Schultz, 2000). Our premise is based upon the idea that the brain is an inference machine (Helmholtz, 1878/1971) and that delusions correspond to false inference. This inference is necessarily probabilistic and rests

upon some representation of predictions (prediction error) and uncertainty (i.e., precision) about those predictions. Within this framework, we see delusions as maladaptive beliefs that mis-represent the world. They might arise through any number of perturbations within this scheme, from an unconstrained specification of the possible or lawful set of neural connections (Hoffman and Dobscha, 1989); providing the potential for bizarre beliefs to form (Hemsley and Garety, 1986a), to an adventitious and inappropriate reinforcement of particular neural connections (King et al., 1984; Shaner, 1999); engendering unexpected percepts, attentional capture and beliefs that deviate grossly from reality (Corlett et al., 2009a; Corlett et al., 2007a; Fletcher and Frith, 2009). Impaired predictive mechanisms have been previously implicated in delusions of alien control; whereby the sufferer believes their movements are under the control of an external agent because of an inability to appropriately predict the sensory consequences of their actions (Frith et al., 2000b). We propose that this account generalizes from actions to numerous cognitive processes, that predictive learning and prediction errors are general mechanisms of brain function (Friston, 2005b; Schultz and Dickinson, 2000) and that aberrant predictions and prediction errors provide a unifying explanation for delusions with disparate contents.

A crucial distinction, which we will appeal to repeatedly, is between prediction errors per se and the precision or uncertainty about those errors. We will develop the argument that delusions (and their neurotransmitter basis) represent a failure to properly encode the precision of predictions and prediction errors; in other words, a failure to optimise uncertainty about sensory information. Here, prediction errors encode information that remains to be explained by top-down predictions (Rao and Ballard, 1999). This distinction is important because it is easy to confuse the role of phasic dopaminergic discharges as encoding reward prediction error (Montague et al., 1996; Schultz et al., 1997), and the role of dopamine in modulating or optimising the precision of prediction errors that may or may not be reward-related (Friston et al., 2009), for example by modulating the signal to noise response properties of neural units encoding prediction error. In what follows, we will assume that the pathophysiology of delusions involves a misrepresentation of salience, uncertainty, novelty or precision (mathematically precision is the inverse of uncertainty). Biologically, this corresponds to aberrant modulation of post synaptic gain that, presumably, involves NMDA receptor function (Friston, 2010). This fits comfortably with the role of dopamine in controlling signal to noise and the numerous proposals that dopamine (at least in terms of its tonic discharge rates) encodes uncertainty or violation of expectations (Fiorillo et al., 2003; Preuschoff et al., 2006).

The challenge is to provide empirical data that test the hypothesis. Numerous investigators have accepted this challenge and, by sharing a set of common simplifying assumptions, we are beginning to develop an understanding of delusions in the brain. Here, we review this growing understanding, beginning with a set of principles which, we believe, are important in developing our understanding of the neurobiology of delusions.

# 2. Reductionist principles for a neuroscience of delusion

The four principles are as follows: Beliefs and memories share cognitive and neural mechanisms (1); learning memory and belief influence perception (2); affect impacts upon learning and memory and hence belief (3); our sense of self, agency, free will and beliefs about others are governed by the same simple neural learning mechanisms (4). By taking a reductionist approach, grounded in formal animal learning theory, computational and cognitive neuroscience we can begin to tackle the hard problems of belief, delusion, and the brain; problems often considered beyond the scope of neuroscience. Below, we consider the principles in more detail before discussing their implications for understanding the cognitive neuroscience of delusions.

#### 2.1 Beliefs and memories share cognitive and neural underpinnings

Beliefs are notoriously difficult to define (Dennett, 1995), but generally refer to the attitude we have with regard to propositions about the world. Perhaps a pragmatic analysis might help. What functions do beliefs serve? Like memories, beliefs help us to organize incumbent information and coordinate adaptive responses (Dennett, 1995). In other words, though beliefs and memories are based on past experiences they are utilized to predict the future and respond accordingly (Corlett, 2009). The most rigorous and formal definition of beliefs appeals to probability theory, and in particular Bayesian formulations (Bayes, 1763). This framework, which we use later, associates beliefs with probability distributions that are represented by the brain (Fiser *et al.*, 2010). These comprise posterior beliefs that are conditioned upon sensory information and are constrained by prior beliefs. In the context of hierarchical Bayesian inference, the posterior belief (having seen the evidence) rests on empirical priors. Empirical priors are prior beliefs that are themselves optimised during hierarchical inference (Friston, 2005b). Assuming that the brain uses hierarchical inference to make predictions about the world, most of the beliefs it entertains can be regarded as empirical prior beliefs. From now on, we will refer to these as prior beliefs or priors and associate these with the source of top-down predictions that are used to form prediction errors. Some have equated beliefs with stimulus-response habits in experimental animals: the behaviors that track previously experienced contingencies but are insensitive to alterations in those contingencies (Eichenbaum, 2000). Indeed, in view of their tenacity and tendency to misrepresent true contingency, some have pointed out the similarities of beliefs to superstitious behaviors (Beck et al., 2007). Thus, beliefs, and therefore delusions, are regarded as representing adventitiously reinforced superstitions; predictions about the future that were formed accidentally and inappropriately but that nevertheless persist (Freeman et al., 2009; Shaner, 1999). Despite capturing aspects of belief phenomenology, these theories offer neither a mechanistic nor a neurobiological explanation of belief or delusion formation. This is what we seek here.

One compelling approach equates the process of human belief formation with Pavlovian conditioning. The same processes that drive animals to learn predictive associations between sensory stimuli and salient events (rewards or punishments) also contribute to the acquisition of beliefs in humans (Dickinson, 2001). Expectancy and experience, or, more specifically, mismatches between the two, are crucial for learning (Alloy and Tabachnik, 1984; Courville et al., 2006; Waldmann, 1998). This mismatch, or prediction error, is central to formal associative learning theories, driving learning directly (Rescorla, 1972) and indirectly, via the allocation of attention toward potentially explanatory cues (Pearce and Hall, 1980). However, there is also a tendency to focus on, and learn about, highly salient stimuli that consistently predict important consequences (Mackintosh, 1975). Under one account (Grossberg, 1982), the occurrence of an expected event that matches an active expectancy will amplify its representation in short-term memory, increasing the likelihood that it will be consolidated within long-term memory as well as the strength of this consolidation. By contrast, when an unexpected event violates the active expectancy, an orienting system is activated which resets short-term memory (dropping active expectancies) and engages an orienting response, permitting the acquisition of new explanatory associations. In essence, organisms learn associations between stimuli, events, thoughts and percepts to build an internal model of their environment. (Sokolov, 1960; Tolman, 1932). This model is itself predictive and, whenever significant novelty is detected due to a mismatch between its predictions and actual experience it must be updated (Grossberg, 1982). In short, the allocation of attention toward appropriately salient events depends upon the optimization of the precision of top-down priors, relative to bottom-up evidence; both in sensory cortices [involving acetylcholine (Yu and Dayan, 2005)] and in frontrostriatal circuits [involving dopamine (Friston et al., 2009)].

This presents the organism with a challenge: to navigate the world successfully, we must sustain a set of prior beliefs (our internal model), sufficiently robust that we do not react reflexively and chaotically to any incoming sensory stimulus. At the same time, these beliefs (priors) must not be so immutable that our responses become fixed, stereotypical and insensitive to change(Corlett *et al.*, 2009b). According to learning models of delusions, during the earliest phases of delusion formation aberrant novelty, salience or prediction error signals drive attention toward redundant or irrelevant environmental cues, the world seems to have changed, it feels strange and sinister, such signals and experiences provide an impetus for new learning which updates the world model inappropriately, manifest as a delusion (Corlett *et al.*, 2009a; Corlett *et al.*, 2007a; Gray, 2004, 1991; Hemsley, 1994; Kapur, 2003). The insight relief that delusions bring engages strong memory consolidation, furthermore, they are deployed reflexively in response to similar aberrant experiences (Mishara, 2009) and as such, they are rapidly rendered impervious to contradiction (Corlett *et al.*, 2009a, see below).

2.1.1 Neural instantiation of predictive learning and belief—Midbrain dopamine neurons in substantia nigra (SN) and ventral tegmental area (VTA) code a reward prediction error (Montague et al., 1996; Schultz et al., 1997). When primates (Schultz et al., 1993; Waelti et al., 2001) and rodents (Takahashi et al., 2009) learn, activity in these neurons reflects a mismatch between expected and experienced reward that is redolent of the prediction error signal from formal learning theories (Waelti et al., 2001) and machine learning models (Montague et al., 1996; Sutton, 1998). However, recent studies have identified punishment prediction error signals (Matsumoto and Hikosaka, 2009) and mismatches between expected and experienced information (Bromberg-Martin and Hikosaka, 2009) in distinct anatomical populations of midbrain dopamine neurons, suggesting that these neurons and the circuits in which they are embedded are involved in the processing of salient events that will guide future adaptive behavior, for both positively and negatively valenced events (Hikosaka et al., 2008a). In human subjects, a circuit involving the midbrain and its projection sites in the striatum and prefrontal cortex signal prediction errors that guide causal learning (Corlett et al., 2004; Fletcher et al., 2001; Turner et al., 2004).

Prediction error-driven learning and memory may represent a basic mode of brain function, referred to as predictive coding (Friston, 2005b, 2009; Schultz and Dickinson, 2000), that is, brains, component brain systems and even single neurons minimize uncertainty about incident information (either external or internal) by structurally or functionally embodying a prediction and responding to errors in the accuracy of the prediction (Fiorillo, 2008). Rapid excitatory and inhibitory neurotransmitters (glutamate and GABA) interact with slower neuromodulatory transmitters to instantiate this predictive coding scheme (Friston, 2005b, 2009), but the precise mechanism for computing prediction error signals remain poorly understood. Across successive levels of cortical hierarchies, top-down signaling from neurons in layers higher up the hierarchy confer expectancies, possibly through glutamatergic NMDA receptors but this is still not established empirically. Bottom-up inputs to a layer are signaled from the layer below through fast glutamatergic and GABAergic mechanisms. At a given level, any mismatch between expectancy and experience is transmitted up the cortical hierarchy to the level above via AMPA receptor signaling (Angelucci et al., 2002a; Angelucci et al., 2002b; Friston, 2005b, 2009; Sherman and Guillery, 1998). Slower neuromodulatory transmitters, like dopamine, acetylcholine, serotonin and cannabinoids are engaged (Corlett et al., 2009a), mediating the post prediction error response by encoding the precision of or uncertainty associated with a particular prediction error(Friston, 2005c). Such uncertainty signals engage subsequent processing such as enhancing neural maintenance of working memory (Lavin et al., 2005) and modulating synaptic plasticity down the hierarchy thus tuning subsequent responses (Grace,

1991; Herrero *et al.*, 2008). We shall refer this perspective on cortical processing, through feedforward signaling of sensory stimuli and feedback signaling of expectation and priors, as the Bayesian model.

According to this model, a prior belief is updated by prediction errors to provide a probabilistic prediction of expected inputs. Input probabilities are learnt at synapses by virtue of experience-dependent learning (Soltani and Wang, 2010), and read out at the level of neural activity populations (Ma et al., 2006). However, beliefs and priors are more than expectancies; strong prior beliefs can enhance, attenuate or vitiate sensed inputs sculpting them to conform to expectations (Jaspers, 1963). The power of prior expectancies can be observed in visual illusions, for example the hollow mask illusion in which a hollow mask is perceived as a convex face as a result of extended lifetime experience that faces are not concave but convex. Likewise strong neural priors can sculpt input signals so that they conform to expectancies (Rao and Ballard, 1999). Beliefs then, not only provide a mechanism through which current information is interpreted in light of the past; they involve an inductive inference that ensures experiences conform with expectancies (Clifford, 1877). In associative learning, such behavioral inflexibility involves training in which expectancies are continuously confirmed (Adams, 1981). The representations and neural circuits controlling behavior gradually shift from more plastic goal-directed, knowledge-based frontal, temporal and ventral striatal regions of the brain toward more inflexible habitual behavior, decreased involvement of frontal cortices and a shift toward dorsal striatal circuits (Belin et al., 2009; Daw et al., 2005; Eichenbaum, 2000). This shift is marked by an increasing strength of the behavior even when the contingency no longer pertains or when the consequences of that behavior are no longer desired.

Whilst Bayesian models are often considered rational and optimal (Shanks, 2008), they have nevertheless been deployed to explain irrational processes such as the spread of panic and rumor within a crowd (which occurs rapidly in salient situations with few explanatory priors; Butts, 1998) and, more recently, a biophysically plausible model offers an explanation for base rate neglect in probabilistic decision making (Soltani & Wang, 2010). Essentially we advocate an explanation of delusions as a disruption to the normal Bayesian predictive mechanisms of the brain such that predictable and irrelevant events mismatch with expectancies and their salience demands new learning and explanation; a delusion represents an explanatory mechanism, an attempt to impose order on a disordered perceptual and cognitive world (McReynolds, 1960; Maher, 1975; Gray et al, 1991; Kapur, 2003; Corlett et al, 2007; Fletcher & Frith, 2009; Corlett et al, 2009a).

2.1.2 Oscillation signatures of match and mismatch events—In our introduction we alluded to the importance of dysfunctional neural circuits (rather than isolated regions) when considering the pathophysiological mechanisms underpinning delusions. That is, psychoses could be conceived as 'disconnection syndromes' (Friston and Frith, 1995). Interand intra-regional neural connections and disconnections are still poorly understood at the present time. One of the active research areas is the examination of the role of neural oscillations in inter-areal communication (Uhlhaas et al., 2008; Uhlhaas et al., 2006a; Uhlhaas and Singer). For example, oscillatory activity in the gamma frequency band (30-50hz) contributes to synchronizing populations of neurons in different brain regions, mediating the temporal structuring of neural activity necessary for sharing, transfer and storage of information (or learning) between these groups of coordinated cells or cell assemblies (Buzsaki, 2007). Such oscillations are thought to reflect the engagement of high level cognitive processes such as attention (Joliot et al., 1994). A recent computational model of selective attention, consisting of a reciprocally connected loop between a sensory circuit and a high-level cognitive circuit, found that top-down signaling enhances gammaband oscillatory coherence only when there is a match between the attended stimulus feature

(expectation) and the actual stimulus feature (experience), and that this occurs exclusively in sensory neurons selective for the actual feature and in memory neurons (that are the source of top-down signaling) selective for the attended feature (Ardid *et al.*, 2010).

Learning from the violation and confirmation of our expectancies can both be traced in oscillatory activity of recurrent neural circuits (Grossberg, 2009). Match based learning captures the Hebbian notion of cell assemblies; collections of synaptically interconnected cells whose pre- and post- synaptic firing correlates and becomes mutually excitatory such that when a fraction of an input pattern is incident upon the assembly, the whole output is realized (Hebb, 1949a). In human learners, gamma oscillations (measured using EEG) increase during acquisition of new associations, as does the coherence of oscillations in cortical regions representing the stimuli being associated (Miltner *et al.*, 1999). Neural synchrony impacts on learning because synaptic plasticity depends on the timing of pre- and post- synaptic neural spikes (Bi and Poo 2001).

But as we have observed, learning does not proceed by contiguity alone (Konorski, 1948). Cell assemblies also represent events that do not match our expectancies (O'Donnell, 2003). In terms of synaptic machinery, one type of mismatch based learning, which is based on expected rewards, appears to be implemented in the mesocorticolimbic system through a trisynaptic arrangement between pre and post-synaptic glutamatergic signaling with a modulatory role for the dopaminergic prediction error input from VTA (Pennartz et al., 2000; Schultz, 1998). Ensembles of neurons are defined by their membrane potential states; periods of very negative resting membrane potential or down states are periodically interrupted by a plateau depolarization or Up state (Haider et al., 2006; Ros et al., 2009; Sanchez-Vives and McCormick, 2000). Striatal up states are synchronized with those in frontal cortex (Goto and O'Donnell, 2001). Dopamine D<sub>2</sub> receptor signaling is associated with an instability of prefrontal representations (Seamans and Yang, 2004), providing an ensemble-level mechanism for surprise driven resetting of representations, search and new learning (Braver and Cohen, 1999; Grossberg, 1982). On the other hand, dopamine, acting through D<sub>1</sub> receptors and their interaction with NMDA channels facilitates the maintenance of Up-states in target neurons (Cepeda and Levine, 1998; Wang and O'Donnell, 2001) and reinforces cell assemblies representing expected salient events (O'Donnell, 2003). In this scheme, the excessive  $D_2$  signaling, impaired  $D_1$  and impoverished NMDA signaling that comprise psychotic states would lead to a poor specification of prior expectancies and frontostriatal cell assemblies comprised of cells representing merely coincident events and spurious associations.

But, how are predictions and prediction errors reflected more generally in the oscillatory signals of cortical hierarchies? While gamma oscillations are commonly enhanced under conditions that involve cognitive control, the top-down specification of priors may be reflected in beta-band (15–30 Hz) oscillations (Wang, In Press). For instance, when recordings are made from the lateral intra-parietal cortex and prefrontal cortex of behaving monkeys during a visual search task, the inter-areal cohence is enhanced in the beta frequency band when the target and distractors are similar and visual search depends on top-down signalling, relative to when the target and distractors are dissimilar and target detection is based by feedforward perceptual 'pop-out' (Buschman and Miller, 2007).

Cortical areas have a well defined laminar structure and, in the neocortex, gamma band oscillations are prominent in superficial layers (2/3) endowed with abundant horizontal connections (Binzegger *et al.*, 2004; Buhl *et al.*, 1998). In contrast, deeper layers (5/6) tend to display lower frequency beta-band oscillations (Wang, In Press). Between reciprocally connected cortical areas, feedforward projections from a lower to a higher area originate in superficial layers of the lower area. Feedback connections begin in deep layers of the higher

area and project to superficial layers of the lower area as well as subcortical structures. Thus beta oscillations, produced in the deep layers, may be especially involved in long distance signaling along feedback pathways. Top-down beta oscillations may encode the expectations that guide match-based learning and perception (Berke *et al.*, 2008). Moreover, prior specifying, beta-frequency oscillatory feedback signals emanating from a 'cognitive area' project to superficial layers 2/3 in a 'sensory area', hence are well suited to modulating gamma oscillations that are locally generated in the superficial layers, in a context dependent manner (Wang, 2010).

There are competing theories regarding the roles of different oscillatory bands in conveying neuronal predictions and prediction errors (Grossberg, 2009). For example, the relationship between high frequency gamma and lower frequency theta band oscillations in hippocampal neurons appears important for the recall of temporal sequences of events (Lisman and Buzsaki, 2008), this form of coding may be especially important in specifying predictions about the future (Lisman and Redish, 2009) and, if it is disrupted, prediction errors may result (Lisman and Grace, 2005); these aberrant errors may be propagated to target structures though inappropriate entrainment of oscillations between structures (Sirota et al., 2008). Furthermore, there are magnetoencephalography data suggesting that, during a face perception task in human subjects, higher-frequency gamma oscillations at lower levels of a neural hierarchy can entrain lower frequency (alpha-band) oscillations in regions higher up the hierarchy, which may represent accumulating prediction error for perceptual synthesis (Chen et al, 2010). Through nonlinear coupling, gamma oscillations in the higher region increase, providing a mechanism through which ascending prediction errors are damped down or explained away (manifest as a decrease in alpha-band power; Chen et al, 2010). More data are clearly required. However, we can predict that in delusion-prone individuals, if predictions are poorly specified and errors signalled inappropriately, then low frequency oscillations, gamma oscillations and their interaction should be perturbed. Consistent with this prediction, highly schizotypal subjects have electrocortical responses to sensory stimulation in the gamma and beta frequency ranges that were slower to habituate following repeated presentation of the stimuli, indicative of maladaptive prior expectancies as well as aberrant prediction error responses (Vernon et al., 2005). Furthermore, patients with schizophrenia have reduced long-range phase synchrony in the beta band during gestalt stimulus perception, perhaps indicative of aberrant prediction error. This aberrant signalling correlated with delusion severity across subjects (Uhlhaas et al., 2006a).

2.1.3 Delusions as aberrant neural learning—Excessive and inappropriate dopamine signaling is thought to render merely coincident events highly salient (Gray, 1991; Hemsley, 1994; Kapur, 2003), this may result from a dysfunction in glutamatergic and GABAergic signaling and thence, the regulation of dopamine signaling (Carlsson et al., 2001; Grace, 1991; Laruelle et al., 2003). Either directly or indirectly, this dysregulation leads to the inappropriate signaling of prediction error (Corlett et al., 2007a; Grace, 1991; Gray, 1991). Since prediction error may guide attention toward events that may explain the feeling of surprise or uncertainty (Pearce and Hall, 1980) and engage learning mechanisms (Rescorla, 1972), we can see that such a disruption has could lead to altered attention, learning, and ultimately belief formation. To consider the nature of this disruption in a little more detail, inappropriate prediction error signals could be conceived of as resulting from a change in the signal to noise properties of dopamine signaling (Grace, 1991; Miller, 1976; Spitzer, 1995); due to deficits in glutamatergic regulation of VTA dopamine neurons. Physiological noise is perceived by the system as real signal that engenders the cascade of events that a true prediction error would engage, namely a search for explanation and new learning. Ultimately, both of these possibilities; inappropriate prediction error and an altered signal to noise ratio of the dopamine system; are reflective of poor precision in the estimation of prediction error (Friston et al., 2009; Preuschoff et al., 2006), which will vitiate inference,

The relationship between conditioning and delusions has also been confirmed in the context of a reward learning task (Schlagenhauf *et al.*, 2009) and an aversive conditioning task (Holt *et al.*, 2008); in both cases, aberrant learning was related to the severity of delusional beliefs. It appears that the brain systems that govern normal causal belief formation are internally and inappropriately engaged when delusions form.

2.1.4 Multiple neural origins for prediction error and its dysfunction?—The computation of VTA prediction error signals involves the interplay between the basal ganglia and the prefrontal cortex (Schultz, 2007; Soltani and Wang, 2008), especially the anterior cingulate cortex (Matsumoto et al., 2007; Rushworth and Behrens 2008) and the orbitofrontal cortex (Schoenbaum et al., 2010). Other studies point to hippocampus, specifically for signaling novelty in the form of mismatches between actual and expected information (i.e. prediction errors) which may then be transmitted to the VTA via the striatum (Lisman and Grace, 2005). This signaling of unexpected and salient events causes the organism to stop its ongoing behavior and search for explanatory cues (Gray, 1991). Patients with psychosis have increased regional cerebral blood flow (an indirect measure of neural activity) in CA1 and, those in whom this effect is most pronounced have the most severe delusions (Schobel et al., 2009). Likewise, individuals in the prodrome (the very earliest phases of psychosis) release more striatal dopamine than controls and again, the magnitude of that dopamine release correlates with the severity of delusion like ideas (Howes et al., 2009). Contrary to the predictions of Gray and Kapur, this dopamine dysfunction has been observed not in the limbic striatum but in the associative striatum, a sub-region that is reciprocally connected with the dorsolateral prefrontal cortex (Haber, 2003; Haber et al., 2006). The latter is a part of the circuit engaged by prediction error driven learning and, moreover, shows aberrant responses in subjects experiencing disturbed percepts and odd beliefs (Corlett et al., 2004; Corlett et al., 2006; Corlett et al., 2007b). We dicuss these observation in more detail below.

The rapidity with which reward prediction error signals are registered in VTA (of the order of milliseconds) may be incommensurate with the calculation of a reward prediction error (Redgrave and Gurney, 2006). Instead these signals could represent unexpected sensory events through cholinergic inputs from the pedunculopontine tegmentum (Dommett *et al.*, 2005), or PPT, inputs which are combined with context representations from the prefrontal cortex and hippocampus as well as motor representations from the putamen in order to ascertain whether the organism or the environment was responsible for the unpredicted event. This agency account suggests that dysfunctions in dopamine signaling could explain both the sense of excessive agency for events in the world associated with paranoia (Kaney and Bentall, 1992) as well as the externalization of agency associated with delusions of passivity (Blakemore *et al.*, 2002; Frith *et al.*, 2000a). See below.

A further candidate site for prediction error dysfunction in psychosis is the habenula (Shepard *et al.*, 2006). The habenula, in concert with the prefrontal cortex, is responsible for instantiating negative prediction error signals in the VTA; the dips below baseline firing that engage extinction learning; abandoning what we have previously learned in favor of a new prediction (Pan *et al.*, 2008). A deficit in this signaling would raise baseline

mesocorticolimbic dopamine levels (Lecourtier *et al.*, 2008) and impair extinction learning (Holt *et al.*, 2008; Waltz *et al.*, 2007), perhaps explaining why deluded individuals stick with maladaptive and erroneous ideas (or corticostriatal cell assemblies) despite their demonstrable falsehood(Corlett *et al.*, 2009c).

Bringing these observations together, it appears that the mesocorticolimbic dopamine system codes numerous types of expectation, their violation and the new learning that expectancy violation engenders; permitting adaptation to prevailing environmental contingencies (Schultz and Dickinson, 2000). When events that violate perceptual expectations are experienced, the hippocampal projection to the striatum engages a broader population of dopamine neurons in the VTA (Lodge and Grace, 2006a). Furthermore, the prefrontal cortex maintains higher level expectancies representing goals and the actions required to achieve those goals (Grace et al., 2007; Sesack and Grace, 2010) as well as reward values for sensory stimuli (Schoenbaum et al., 2010) and actions (Rushworth and Behrens 2008). When events occur that violate those expectancies, PFC modulates the responses of active VTA dopamine neurons: engaging burst firing through its influence over the PPT (Lodge and Grace, 2006a, b), allowing updating of expectancies through new learning. Furthermore, PFC enables the quiescence of those same VTA neurons (through its influence on the habenula) when contingencies change and learning is extinguished (Hikosaka et al., 2008b; Pan et al., 2008). Reciprocal connections between VTA, PFC, striatum and hippocampus are involved in this updating process so that future expectancies conform to the prevailing environmental contingencies.

We predict that delusions are associated with a threefold disturbance in this circuitry: (i) Excessive hippocampal drive to VTA (via striatum) engaging a broader population of VTA dopamine neurons; (ii) Inappropriate engagement of PPT due to PFC dysfunction, instigating burst firing in that expanded pool of recruited neurons and (iii) Impaired habenula mediated inhibition of VTA dopamine neurons (which would normally instantiate extinction learning when an expected event fails to occur).

These three deficits would confer the cardinal characteristics of delusions, their bizarreness and tenacity: Bizarreness; due to the aberrant recruitment of VTA cells and their incorporation into cell-assemblies which sculpt future expectancies; and tenacity; due to the failure of PFC to control the habenula, and hence co-ordinate the dips in VTA neuron firing below baseline that engage extinction learning when the predictions of the delusion are not borne out.

While this model begins to implicate aberrant learning processes in delusion formation, it does not address the range of different themes that form the content of delusions, nor does it fully explain the behaviors in which deluded individuals engage when confronted with evidence that challenges their belief (see below). In order to extend out explanation to encompass these characteristics, we discuss below what we consider to be key factors: the role of beliefs in instrumental conditioning (learning the relationships between our actions and their effects in the world) and the impact of repeated recall and rehearsal of that information on subsequent processing.

#### 2.2 Learning, memory and belief alter perception

Perception is substantially constructive. That is, our expectancies (based on previous experience) contribute to what we currently perceive by sculpting sensory inputs (Bruner *et al.*, 1949; Helmholtz, 1878/1971). The concepts and categories we have learned through experience can influence what we perceive, for example, if subjects are shown simple objects and asked to reproduce their colors, their responses are heavily influenced by the shape of the object (Goldstone, 1995). Motivational state, itself interacting with learning and

Corlett et al.

memory (Berridge, 2007), can impact upon perceptual judgment; poorer children judge coins to be larger and heavier than do richer children (McCurdy, 1956). When presented with noisy, unstructured visual inputs, hungry subjects claim to see food objects (Atkinson, 1948). The impact of motivation on bottom-up perceptual inputs may be mirrored in the mechanisms we use to imagine given percepts, a mechanism which, when inappropriately engaged may elicit hallucinations (Grossberg, 2000) and the impaired reality monitoring associated with delusional ideation (Simons *et al.*, 2008)). For example, the spontaneous confabulations of patients with orbitofrontal lesions represent an excessive influence of past experience on current perceptual expectations such that known people or places lack a sense of familiarity (Fleminger, 1992).

2.2.1 Neural mechanisms of the memory-perception cycle—Predictive coding and prediction error may be a basic mode of brain function (Friston, 2005b, 2009; Mesulam, 2008). This theory is best encapsulated by sensory cortices, in particular the visual cortex; whose anatomy recapitulates the idea of a hierarchically organized predictive coding system. Further up the neural hierarchy, more distal to the site of sensory input, approaching association cortices, the representations of sensory stimulation become more abstract (Mesulam, 2008). But the percept does not emerge as a consequence of a simple unidirectional progression up this hierarchy (Sperry, 1990). Rather the hierarchy is nested (Feinberg, 2000; Feinberg and Keenan, 2005) or enriched by interactions (feedback as well as feedforward) between its layers (Friston, 2005b, 2009). These interactions are instantiated by sparse and rapid feedforward AMPA and GABAergic signaling meeting feedback (possibly NMDA-mediated) signaling representing predicted inputs embodied in the layers above (Friston, 2005b). Any mismatch between expectancy and experience (signaled via AMPA receptors) can serve to update future priors. Dysinteractions within this Bayesian hierarchical arrangement may be responsible for the symptoms of psychosis (Corlett et al., 2009a; Fletcher and Frith, 2009). For example, in the absence of stable prior expectancies, certain perceptual illusions may not be perceived by patients with schizophrenia (Dakin et al., 2005; Dima et al., 2009; Emrich, 1989) nor individuals administered NMDA receptor antagonists (Phillips and Silverstein, 2003) perhaps indicative of a common underlying mechanism [although see (Passie et al., 2003) for a dissociation between the effects of ketamine on a perceptual illusion and its psychotomimetic effects]. See Figure 2.

The thalamus has also been strongly implicated in conscious perception. Thalamocortical circuits have intrinsic resonance in the gamma frequency range which is critical for conscious perception, prediction and learning (Steriade *et al.*, 1991). GABAergic neurons in the basal ganglia projecting to the thalamus exert an inhibitory influence on thalamocortical neurons thus protecting the cortex from sensory overload (Sharp *et al.*, 2001). Hyperactivity of dopamine or hypo-activity of glutamate in the striatum would compromise these protective mechanisms leading to excessive cortical stimulation and psychosis (Carlsson *et al.*, 2001; Carlsson and Carlsson, 1990; Geyer and Vollenweider, 2008). Such a deficit could conceivably alter the sense of background and foreground that permeates normal perception (Conrad, 1958a). This could explain why other Gestalt principles, which involve grouping the perceptual field on the basis of learned environmental regularities (Fiser, 2009; Vickery and Jiang, 2009), are impaired by psychotomimetic drugs that alter dopaminergic and glutamatergic function (Kurylo and Gazes, 2008). Gestalt organizing principles are similarly disrupted in patients with schizophrenia (Silverstein *et al.*, 2006; Uhlhaas and Mishara, 2007; Uhlhaas *et al.*, 2006b).

Like other systems, thalamocortical circuits and their interaction with cortical information processing have been subject to a Bayesian analysis (Koechlin *et al.*, 1996). According to this scheme, thalamocortical information represents the feedforward aspect (the information

being represented) and cortico-cortical processing represents the prior expectancies, the operations to be performed on that information. Similar models have been developed to account for perception of coherent visual motion and mental rotation, as well as the predictive functions involved in enacting adaptive movements (Koechlin *et al.*, 1996; Llinas and Roy, 2009).

Inherent in all of these related schemes is the notion of a balance, between bottom-up and top-down (or feedforward and feedback) signaling. This balance is necessary in order to meet the afore-mentioned challenge of a system that is robust to noisy inputs (through reliance on empirically derived prior expectations) but is also flexibly responsive to new contexts and situations (through the capacity to alter priors on the basis of bottom-up signal). With this in mind, it is clear that, in addition to poorly specified predictions, excessively strong priors may be profoundly disruptive and psychotogenic (Corlett *et al.*, 2009a). Perceptual associations between sensory modalities appear to be learned using mesocorticolimbic prediction error signals (den Ouden et al., 2010; den Ouden et al., 2009), which may explain the phenomenon of sensory conditioned hallucinations (Ellison, 1941; Seashore, 1895) whereby, learned associations between sensory stimuli (a tone predicts light stimulation for example) alter perception, such that presentation of one stimulus (tone) induces experience of the other (light) even though the latter is not present. Learned associations can alter perception; Hallucination-prone individuals are more susceptible to experiencing sensory conditioned hallucinations (Kot and Serper, 2002). Likewise delusional beliefs can alter percepts such that they conform to the delusion (Jaspers, 1963). Excessively strong top-down predictions may explain the psychotogenic effects of LSD and sensory deprivation (Corlett et al., 2009a). Furthermore, individuals prone to abnormal experiences and beliefs are more susceptible to the Deese-Roediger-McDermott memory illusion whereby they claim to have experienced an event that was strongly expected but nevertheless did not occur (Corlett et al., 2009d). We predict that such expectation-based psychotic phenomena would be associated with inappropriate gamma and beta oscillations, reflective of inappropriate reverbatory activity in recurrent neural circuits and of pattern completion within Hebbian cell assemblies that are not relevant to the situation at hand.

#### 2.3 Affect impacts upon learning, memory, perception and hence belief

The aberrant percepts that drive delusion formation often occur during periods of stress and are themselves anxiogenic (Keinan and Keinan, 1994). Furthermore, individuals with a low tolerance for ambiguity are more prone to paranormal beliefs and odd experiences (Houran and Houran, 1998). Some models posit a vicious circle in which fear and aberrant perception are mutually reinforcing and demand explanation, culminating in a delusion which then subtends future aberrant percepts and inappropriate fear (Lange *et al.*, 1998; Pally, 2007). These models are descriptively compelling but are expressed largely at the higher cognitive level. We seek a more fundamental neural and cognitive explanation. Simply put, we argue that affectively charged uncertainty drives delusion formation, through establishment of predictive associations that, whilst maladaptive, represent attempts to render the world more predictable.

**2.3.1 Neural mechanisms of affective modulation**—The uncertainty engendered by aberrations of experience is affectively charged (Vinogradov *et al.*, 1992). Affective learning is also prediction error driven, involving a circuit incorporating the VTA, amygdala and hippocampus as well as the striatum and prefrontal cortex (Delgado *et al.*, 2008b; Laviolette and Grace, 2006; Milad *et al.*, 2007; Milad *et al.*, 2004; Schiller *et al.*, 2008). Dysfunctions within these nodes could engender fear in the wrong context, leading to maladaptive learning about the danger of adverse consequences. The top-down instantiation of extinction learning is particularly interesting in this respect; the dopaminergic and GABAergic

mechanisms that override old fear learning with new extinction learning (Bissiere *et al.*, 2003) may be impaired in schizophrenia (Holt *et al.*, 2008). It is clear that paranoia could be accounted for parsimoniously by appealing to an inappropriate engagement of the brain's fear system and its persistence by an impairment of the brain's mechanisms of extinction.

The amygdala is crucial for fear learning in rodents and humans (Critchley *et al.*, 2002; Morris *et al.*, 2002). However, its role may not be limited to fear; the amygdala is involved in coding, processing and learning about salient stimuli (Balleine and Killcross, 2006; Paton *et al.*, 2006). The link between fear and uncertainty is emphasized by theorists who posit that the amygdala is also engaged during conditions of uncertainty about biologically relevant stimuli that warrant vigilance (Sander *et al.*, 2003; Whalen *et al.*, 1998). For example, fearful faces represent ambiguous stimuli, since they signal the presence but not the source of threat (Whalen *et al.*, 2001). Amygdala responses to appetitive and aversive events are modulated by predictability, being more marked when salient events are uncertain (Belova *et al.*, 2007). In this respect, it is noteworthy that animals with lesions of the central nucleus of the amygdala do not allocate more attention to surprising events (Holland and Gallagher, 1993b).

Cholinergic interneurons in the substantia innominata/nucleus basalis and their projections to posterior parietal cortices are important for the surprise-induced enhancement of attention (Chiba et al, 1995; Bucci et al, 1998; Han et al, 1999). In humans, cues that predict aversive events engage both striatum (Delgado et al., 2008a) and amygdala (Schiller et al., 2008) but only the striatum codes aversive prediction error (Schiller et al., 2008), suggesting that the amygdala is involved in representing the salience of events learned as a consequence of prediction error signals transmitted from other regions. Aberrant prediction error responses in the midbrain or striatum could therefore encourage inappropriate assignment of significance to stimuli, thoughts and percepts (Kapur, 2003) which are then allocated attention in the amygdala (Laviolette and Grace, 2006) through changes in fronto-parietal spatial representations (Mohanty et al., 2009). These environmental contingencies are also subjected to strong consolidation through changes in synaptic strength in the rhinal and entorhinal cortices (Hikosaka et al., 2008a), hence, future encounters with similar cues will engender rapid and powerful predictions of aversive stimulation which would engage avoidance behaviors. Impairments in this system could then contribute to the maintenance of paranoia (Freeman et al., 2007; Moutoussis et al., 2007).

Uncertainty is a powerful and uncomfortable experience. A consequence of such perceived and unsettling lacking of control is that subjects strive to find consistent relationships. They consequently become prone to finding illusory patterns, seeing figures in noise, recognizing correlations between unrelated events, creating superstitious rituals and endorsing conspiracy beliefs (Whitson and Galinsky, 2008). We contend that these healthy coping mechanisms are magnified in individuals with psychosis, culminating in the formation of delusions. These 'filling in' processes may result from top-down influences of orbitofrontal cortex, which receives information from the each modality-specific cortical pathway specifying what a particular sensory object is (Rolls et al., 2008), for example; the inferior temporal cortex where object and face identity are encoded (Rolls, 2007) and the superior temporal sulcus where face expression and gesture are represented (Hasselmo et al., 1989a; Hasselmo et al., 1989b). Furthermore, the orbitofrontal cortex has inputs from the amygdala and the ventral tegmental area (Takahashi et al., 2009) which may drive its ability to learn affective value representations (Padoa-Schioppa and Assad, 2006) which appear to modulate perception in a top-down manner (de Araujo et al, 2005); when affectively charged external labels are applied to percepts, OFC responses bias cingulate and striatal responses in the direction of the label (Grabenhorst et al., 2008). Furthermore, damage to the OFC can result in spontaneous confabulation, a delusion-like disorder in which patients confuse ongoing

reality with past experiences (Schnider, 2003). Thus, hyper-engagement of top-down attentional biases may contribute to the aberrant salience underpinning delusional beliefs (Kapur, 2003) as well as to their maintenance (Corlett *et al.*, 2009a; Corlett *et al.*, 2009c).

#### 2.4 Simple synaptic learning and memory mechanisms of belief govern

**2.4.1 Our sense of self, agency and free will**—Like beliefs, the self is difficult to define and multifaceted (Mishara, 2007). We will focus on one conception of self, that of an agent that is responsible for actions (Wegner, 2004). In this respect, excessive agency accounts of paranoia (Kaney and Bentall, 1992) may be enriched by a consideration of the phenomenon of superstitious instrumental conditioning (Skinner, 1948), in which spurious associations are learned between an action and a salient outcome and the action persists despite there being no causal connection between it and the salient outcome. An excessively noisy dopamine system would be fertile grounds for superstitions, which are essentially delusional associations that are reinforced between merely coincident thoughts or actions and environmental events (Shaner, 1999). According to action reselection hypotheses of dopaminergic prediction error signals (Redgrave and Gurney, 2006), inappropriate dopaminergic prediction error signals would confer a spurious sense of agency for events.

Initial lesion studies suggested that hippocampal damage increased superstitious learning in experimental animals (Devenport, 1979). However, more extensive investigations implicated the parietal cortex in superstitious responding, suggesting that collateral damage to this region of cortex may have occurred when the hippocampus was aspirated (Mittleman *et al.*, 1990). Elevated superstitious responding has been demonstrated in chronic ketamine users with delusion like ideation and perceptual aberrations (Freeman *et al.*, 2009) and patients with schizophrenia who have delusions (Roiser *et al.*, 2009), although the rate of superstitious responding in (presumably non-delusional) control subjects was high in both of these studies.

Lesions of the parietal cortex grossly alter bodily perception and representation, for example, hemi-spatial neglect involves a failure to appreciate half of the body, external world and mental images (Bisiach and Luzzatti, 1978). Perhaps another function of the parietal cortex in instrumental learning involves keeping track of the sense of self as agent in the environment (Farrer et al., 2008). Wegner and others hypothesize that a sense of self agency may be learned through experience; having an intention to act very frequently precedes the action itself and this contiguity binds intentions with actions through associative learning (Glymour, 2004; Hume, 1739/2007; Wegner, 2004). This system can be fooled using subliminal prime events that alter the contiguity between actions and outcomes (Aarts et al., 2005) and furthermore, subjects judge the time between performing an action and producing an outcome as shorter when the action was intentional, a process of actionoutcome binding (Moore et al., 2009). Schizophrenic patients with severe positive symptoms show a hyper-binding effect, an exaggerated binding between their actions and the outcomes they produce, consistent with a disturbed agency account of paranoia (Franck et al., 2005; Haggard et al., 2003). This process of learned intentionality has been modeled using Bayesian mechanisms; in essence, the task of inferring causal agency involves conditioning the evidence (whether the outcome occurred?) over the priors (was there an intention to act and would the outcome be consistent with the outcome performed? (Hendricks, 2007; Lau et al., 2007). Inappropriate engagement of this inference mechanism could account for excessive and inappropriate agency underpinning, for example, beliefs in telekinesis or telepathy, but what about delusions of passivity or external control?

The parietal cortex has also been implicated in passivity experiences through prediction error; in this case, the mismatch between expected and experienced consequences of movements (Schnell *et al.*, 2008). Producing movements over which we feel a sense of

Corlett et al.

agency also involves predictive learning and prediction error (Blakemore *et al.*, 2002). Again, a Bayesian mechanism may underlie motor control; an internal predictive model of motor commands which is used to predict the sensory consequences of movements and compare them with the actual sensory feedback during movement execution (Wolpert *et al.*, 1995; Wolpert and Miall, 1996). The cerebellum appears to store internal world models and compute discrepancies between predicted and experienced sensory consequences of actions (Blakemore *et al.*, 2001). Event related functional MRI studies of the period before a movement show that activations changes in the cerebellum and PFC occur several seconds before movement onset and the degree of cerebellar activation correlates with that in prefrontal and inferior parietal cortices (Allen *et al.*, 2005).

Internal 'forward' models use an efference copy of motor commands (Von Holst, 1954) to make a prediction about the sensory consequences of an action (Blakemore, 2003). This comparison can be used to cancel sensory effects of performing the action, compared with identical movements that are externally produced (Blakemore et al., 1999; Weiskrantz et al., 1971). An impairment in such a predictive system would result in a failure to attenuate the sensory consequences of self-produced actions, making them appear indistinguishable from externally generated sensations and engendering the inference that one's own movements were externally caused (Blakemore et al., 2002; Frith et al., 2000a). This theory provides an elegant explanation for why we can't tickle ourselves, since we cancel the predicted sensory consequences of the action (Blakemore et al., 2000b). However, patients experiencing passivity phenomena and hallucinations, in whom sensory cancellation is presumed to be impaired, rate self generated stimulation as ticklish (Blakemore et al., 2000a). Impaired cancellation of efference copies has likewise been implicated in the pathophysiology of hallucinations; here internally generated speech is misperceived as externally generated due to this impairment in the cancellation of forward model predictions (Ford and Mathalon, 2005; Ford et al., 2007).

There are some rare patients who call the proposed model of passivity into question; subjects who have suffered haptic deafferentiation and therefore do not perceive sensory feedback from the actions they perform (Fourneret *et al.*, 2002). Since a haptically deafferented subject does not suffer from delusions of passivity; some have argued that aberrant percepts of one's own action are not sufficient to explain passivity delusions; invoking a further belief evaluation dysfunction that is necessary for the delusional inference to occur (Coltheart, 2010). To clarify the prediction error based explanation of these phenomena; patients with passivity experiences do not use forward model predictions to cancel the predicted consequences of their movements so they experience the sensory consequences of their actions and therefore be protected from passivity experiences; since such experiences do not depend on absence of feedback but on inappropriately large or unexpected feedback. It is this persistence and unexpected nature of aberrant prediction error that engages delusion formation.

Parietal cortex receives inputs from the cerebellar internal model (Ito, 1993), possibly combining them with a multi-sensory salience map of the external world and the motor plans necessary to approach or avoid salient features (Mohanty *et al.*, 2009). Activity in the parietal operculum is also attenuated during self initiated movements compared with passive movements (Weiller *et al.*, 1996) and during self produced compared with external stimulation. Patients with lesion to the right hemisphere in white matter underlying the parietal operculum delusion that their limb belonged to their niece (Bottini *et al.*, 2002).

Even healthy individuals can be tricked into accepting that a false hand belongs to their own body (Botvinick and Cohen, 1998). If subjects perceive the false hand being stimulated at

the same time as they feel their own (occluded) hand receiving the same stimulation, they begin to feel that the false hand belongs to them, incorporating it into their body schema such that, when asked to estimate where their own hand is positioned, they point to a location closer to the false hand (Makin et al., 2008). Patients with schizophrenia are more susceptible to this illusion (Peled et al., 2003). It appears that the processes of multisensory integration involved in judging ownership of a body part involve synaptic learning via associative Hebbian mechanisms, representing the confluence of seeing a hand stimulated and feeling a hand stimulated (Keysers et al., 2004). Furthermore, top-down attentional biases seem to influence the illusion (Tsakiris and Haggard, 2005). These biases again emerge through associative learning and are subject to the same formal rules, a surprising mismatch between the expected confluence of sensation and vision weakens the illusion. Likewise the illusion does not occur for a stick: people perceive rubber hand illusions more readily than rubber object illusions (Press et al., 2008). Physiological noise in the multisensory integration process that confers bodily ownership may engender mutated prior expectations about the body which bias subsequent perception, resulting in somatoparophrenias, delusions of body representation and agency (Vallar and Ronchi, 2009).

#### 2.4.2 Social learning and therefore our beliefs about others—Social

neuroscientists also appreciate the power of prediction error and predictive coding (Behrens *et al.*, 2009; Kilner *et al.*, 2007a, b; Lee, 2008a). Reinforcement learning circuits are engaged when human subjects make social value judgments and a further network of brain regions is engaged when subjects make judgments about the intentions of others – including the superior temporal sulcus/temporoparietal junction (STS/TPJ) (Behrens *et al.*, 2009). These data build upon previous suggestions that associative principles like prediction error govern various social attribution processes (Miller, 1959). For example social attributions made about worker productivity are susceptible to associative learning phenomena like Kamin blocking (Cramer *et al.*, 2002).

fMRI studies of prediction error driven reinforcement learning usually require participants to learn which of two stimuli to choose in order to win the most points (Pessiglione *et al.*, 2006). In an extension to the standard paradigm, Behrens and colleagues gave subjects an additional source of information, the suggestion of a confederate who may or may not know the appropriate choice to make. Hence the subjects learned simultaneously whether to choose the blue or the green card and also whether they could trust the advice of the confederate. They were able to distinguish brain regions coding a mismatch between expected and experienced reward from brain regions coding a mismatch between expected and experienced truth. Intriguingly, these analyses revealed that adjacent but distinct regions of the anterior cingulate cortex coded reward and truth prediction error. The STS/TPJ also appeared to reflect social prediction errors about the truth of the confederate's advice (Behrens *et al.*, 2008).

The analysis of social learning in terms of prediction error has recently bridged theories of both reinforcement learning and predictive coding. Building upon the empirical Bayes model of brain function, this approach combines the forward model of intentional motor control (Blakemore, 2003; Blakemore *et al.*, 2001; Wolpert *et al.*, 1995; Wolpert and Miall, 1996) with the observations of social prediction errors in STS (Behrens *et al.*, 2008; Hampton *et al.*, 2008) to explain the function of the brain's mirror neurons system through its direct link between action and observation (Kilner *et al.*, 2007a, b). Here, the most likely cause of an observed action can be inferred by minimizing the prediction error across all levels in the cortical hierarchy that are engaged by that observation.

Observing, imagining, or in any way representing an action excites the motor program used to execute the same action (Jeannerod, 1994). Mirror Neurons discharge not only during action execution but also during action observation; they were identified in non-human primates, using neural recording, in area F5 and the inferior parietal lobule (Fogassi and Luppino, 2005; Gallese et al., 1996; Rizzolatti et al., 1996). Functional magnetic resonance imaging data have been used to infer the presence of mirror neurons in the human inferior parietal lobule (Chong et al., 2008) and inferior frontal gyrus (Kilner et al., 2009). However some have failed to find evidence of mirror neuron-like activations (Lingnau et al., 2009). Indeed, the spatial resolution of fMRI is such that it may be inappropriate to ascribe the response in a particular region to a specific population of cells. Furthermore, some have questioned the reified status of mirror neurons; that is, instead of being indivisible, they may simply reflect conditioning of an association between a motor program for an action and a visual representation of that action; learned by experience across the life course (Heyes, 2010). The present theory does not depend on the exact origin of mirror representations and, given that the regions in which mirror neurons have been identified with direct recording in non-human primates largely overlap with those regions that responded to action observation and execution in human subjects, we proceed by discussing the potential role of mirror neurons in human social cognition (Gallese et al., 2004).

Implicit in the description of mirror neurons is the idea that information is passed by forward connections from low level representations of the movement kinematics to high-level representations of the intentions subtending the action. Observation of an action activates the STS, which in turn drives the inferior parietal lobule which drives the inferior frontal gyrus. Formally this is a recognition model that operates by the inversion of a generative model (Kilner *et al.*, 2007a, b). A generative model will produce an estimate of the visual consequences of an executed action given the causes or goals of that action. By inverting the model it is possible to infer the cause or goal of an action given the visual input (Kilner *et al.*, 2007a, b).

Again, bottom-up or top-down biases in this inference process would lead to gross misrepresentations of other's intentions. Those biases may arise due to aberrant prediction error signals, forging maladaptive social expectations manifest phenomenologically as intense feelings of social uncertainty and ultimately paranoia. More recently, it has emerged that beliefs about somebody's mental experience can influence how we perceive their physical attributes (Teufel *et al.*, 2009). While the full connotations of this have yet to be explored, it seems that we may perceive someone's behavior depending on what we think that they are thinking.

#### 3. The fixity of delusions

By inappropriately updating subject's priors, delusions are applied to all subsequent experiences(Conrad, 1958b; Mishara, 2009). Why might this be? Indeed, if we are arguing that delusions form under the influence of inappropriate, uncertain and imprecise prediction error, why do delusions become so tenacious? Here we turn to a process that has received increasing empirical attention in recent years; memory reconsolidation (Misanin *et al.*, 1968; Nader *et al.*, 2000). We conceive of beliefs and delusions as a kind of memory (Eichenbaum, 2000), that is, a means through which past experiences and processing organize responses to current inputs. Memories serve a more dynamic function than simple storage; they can be recalled, returned to a labile state (Misanin *et al.*, 1968; Nader *et al.*, 2000), updated with new information (Estes, 1997) and strengthened (Lee, 2008b); a set of reconsolidation processes that appear to be engaged when unexpected events occur (Eisenhardt and Menzel, 2007). This updating process involves a streamlining or schematization of the representation (Stickgold and Walker, 2007). We have previously argued that, once delusions are formed,

Corlett et al.

future prediction errors engage a reactivation, reconsolidation and strengthening of the delusion; rendering it impervious to contradictory evidence; each time a delusion is deployed, it is reinforced further, conferring resistance to contradiction (Corlett et al., 2009c), rather like the formation of an instrumental habit with overtraining (Adams, 1981; Lee, 2008b; Stickgold and Walker, 2007). That is, when subsequent prediction errors occur, they are explicable in terms of the delusion and they serve to reinforce it, hence the paradoxical observation that challenging subjects' delusions can actually strengthen their conviction (Milton et al., 1978; Simpson and Done, 2002). Neurobiologically, this reconsolidation based strengthening would shift control of behavior toward the dorsal striatal habit system (see Figure 3) and would manifest as immutable prior expectancies in Bayesian cortical hierarchies(Corlett et al., 2009b; Corlett et al., 2009c; Mishara, 2009). Delusions may be maintained despite being fallacious through disruptions in frontostriatal synaptic metaplasticity, a form of 'plasticity of plasticity' (Abraham and Bear, 1996) that allows old associations to be overridden by new learning. Metaplasticity can be restored with n-acetyl-cysteiene, (Moussawi et al., 2009), a drug which increases the availability of glutamate in extrasynaptic spaces by stimulating the cysteine-glutamate antiporter (Baker et al., 2008).. This analysis of delusions, in terms of a shift away from computationally expensive prefrontal processing toward striatal habit (Daw et al., 2005; Mishara, 2009) may also explain the waxing and waning of delusional conviction and the paradoxical double book-keeping; patients endorse particular delusions but do not act as if they truly believe them (Bleuler, 1908; Sass, 2004); such situations would transpire if the goal-directed system occasionally won the competition for control of behavior, a state of the system that can be engendered by enhancing plasticity in prefrontal brain regions (Hitchcott et al., 2007; Moussawi et al., 2009).

Here we draw upon advances in the cognitive neuroscience of addiction to make our case about delusions. Like delusions, aberrant prediction error accounts have been outlined for the generation of addictive behaviors (Lapish *et al.*, 2006; Redish, 2004) as well as their maintenance as habits despite maladaptive consequences (Takahashi *et al.*, 2008). We posit that the inappropriate prediction error that occurs in endogenous psychosis is internally generated (rather than a plastic response to drug consumption, although see (Corlett *et al.*, 2009a) for a review of drug induced psychoses) and that they track merely coincident environmental stimuli rather than cues that predict access to drug and drug induced hedonic states. However, maladaptive prediction error responses in addiction and psychosis may be indicative of a fronto-striatal system that is sensitized toward aberrant learning and may therefore explain the strong co-morbidity between drug abuse and psychosis (Kalayasiri *et al.*, 2006; van Nimwegen *et al.*, 2005).

Reactivating a delusion (perhaps having a patient engage with and ruminate upon it) may drive its representation into a labile state; providing a novel therapeutic window in which to intervene and destabilize the delusion. This approach has been taken previously with some success (Rubin, 1976), however, future well-controlled investigations are essential.

# 4. One or Two Factors?

There are competing accounts of delusions in cognitive neuropsychiatry (Coltheart *et al.*, 2007; Freeman *et al.*, 2002; Garety, 1991; Garety and Freeman, 1999; Gerrans, 2002; Kinderman and Bentall, 1997; McKay *et al.*, 2007). Some argue that perceptual aberrations are all that is required for a delusion to form (Gerrans, 2002; Maher, 1974), others that delusions result from top-down reasoning impairments (Freeman *et al.*, 2002; Garety, 1991; Garety and Freeman, 1999), others still posit some combination of both factors, a two-factor approach in which perceptual and reasoning abnormalities combine (Coltheart *et al.*, 2007; McKay *et al.*, 2007). The latter derive from observations that neurological patients with

delusions often have two sites of damage; a lesion in a perceptual region (such as the fusiform face area) and an additional lesion in 'belief evaluation' regions, possibly in the right frontal cortex (Ramachandran, 1998). The first damage engenders odd percepts and the second generates bizarre explanations.

Prediction error driven Bayesian models of delusions (Corlett *et al.*, 2009a; Fletcher and Frith, 2009) subsume both factors into a single deficit in Bayesian inference; noise in predictive learning mechanisms engender inappropriate percepts which update future priors, leading to the formation and maintenance of delusions. Prediction error signals have been registered in right dorsolateral prefrontal cortex during causal learning (Corlett *et al.*, 2004; Fletcher *et al.*, 2001; Turner *et al.*, 2004), psychotogenic drug administration and endogenous psychosis are associated with inappropriate responding in this region, the magnitude of which was predictive of delusion severity (Corlett *et al.*, 2006; Corlett *et al.*, 2007b).

2-factor theorists have recently equated the inappropriate prediction error signals that we reported in dorsolateral prefrontal cortex with their aberrant belief evaluation process or factor 2 (Coltheart, 2010). However, a single deficit in Bayesian inference is able to explain more of what we know about the interactions between perception and belief-based expectation, the neurobiology of the delusions that occur in schizophrenia and the maintenance of delusions in the face of contradictory evidence. That is, unlike 2-factor theory, our model allows for dysfunctional prediction error to be calculated in PFC and imposed upon the rest of the brain or, alternatively for surprising perceptual inputs to arrive at PFC engaging surprise and demanding explanation. Both of these possibilities (bottom-up and top-down) are aberrations of a single factor; Bayesian inference.

We recognize the strong neurological evidence that perceptual aberration and delusional ideation are dissociable (Coltheart, 2010). However, we emphasize the potential consequences of prefrontal cortical damage alone (their Factor 2) as well peripheral perceptual dysfunction (their Factor 1); there are patients who suffer from delusion-like spontaneous confabulations following damage to ventromedial and lateral prefrontal cortex (Schnider, 2003; Turner *et al.*, 2004) and at least one patient in whom peripheral sensations are perturbed (following damage to the brachial plexus) who has somatic delusions in the absence of any apparent structural damage and by extension any deficit in factor 2 (Ghoreishi, 2010).

In short, the present model suggests that inappropriate mismatches between expectancy and experience engender prediction error where there ought to be none, driving new and aberrant learning directly and through the allocation of attention toward irrelevant but potentially explanatory cues (Corlett *et al.*, 2007a). This learning normally provides the basis for a variety of vital perceptual and cognitive functions that govern our interactions with the environment and other agents so when it malfunctions, gross misrepresentations of reality, delusions and perceptual aberrations, result.

## 5. A Neurodevelopmental Dimension?

Developmental studies suggest that children who go on to develop schizophrenia and therefore likely delusions (although not all patients with schizophrenia have delusions) have subtle neurological 'soft-signs' indicative of aberrant sensorimotor integration (Mohr *et al.*, 1996). In healthy individuals, there are relationships between motor developmental milestones, structural integrity of the frontal cortex, striatum and cerebellum and executive cognitive function, associations which are not present in patients with schizophrenia (Ridler *et al.*, 2006) suggesting impaired bootstrapping of cortical pathways into systems that can predict and respond to their inputs and thus, an impairment of adaptive interaction with the

environment and other agents; individuals with impaired sensorimotor integration throughout development would learn impoverished or maladaptive prior expectancies about the world (Hemsley and Garety, 1986b).

Different homeobox genes are responsible for controlling the development and patterning of the frontal cortex (Tabares-Seisdedos and Rubenstein, 2009), midbrain dopamine neurons (Maxwell and Li, 2005), the striatum (Long et al., 2009), the amygdala (Tole et al., 2005) and cerebellum (Sillitoe et al., 2008). Some of these genes and their expression products have been associated with psychotic symptoms, for example; DLX1 expression is decreased in the thalamus of individuals with psychosis compared with those without a history of psychosis and matched healthy controls (Kromkamp et al., 2003). Likewise, the homeogene Engrailed 2 which controls cerebellar development is associated with schizophrenia (Gourion et al., 2004). Knocking out FGF17, a gene that controls the patterning and organization of frontal cortical development, leads to profound deficits in social interaction in mice, perhaps indicative of a relationship to paranoia (Scearce-Levie et al., 2008). Indeed, a human genetic association study revealed a link between the chromosome region where FGF17 is found (8p13) and delusional beliefs (Chiu et al., 2002). We acknowledge that we are speculating here and we appreciate the dangers of anthropomorphizing social behaviors in rodents; future work should address the validity of FGF-knockout as a model of paranoia by exploring other prediction error related processes in these animals; do they have a deficit in conditioned avoidance learning, for example? We believe that the different themes of delusional beliefs entertained by different subjects may have their origins in subtle developmental dysfunctions in the circuits we have outlined, biasing prediction error driven deficits in glutamatergic and dopaminergic processing toward a particular set of experiences and a specific explanatory belief. Normal variation in these same genetic loci may underpin individual differences in perceptual aberration as well as the themes and severity of delusion-like ideation in the healthy population.

#### 6. Explaining Delusion Content

We now attempt to account for different kinds of delusion within this framework. While the scope of this section is by no means exhaustive, we believe that the range of delusions potentially accounted for within the framework is compelling (see Figure 4).

#### 6.1 Paranoia and delusions of reference

Referential delusions involve the belief that objects, events and agents in the environment are communicating specific and personal messages (Conrad, 1958b) ranging from the inanimate to animate, from newspapers, to television newsreaders (Startup and Startup, 2005) and even the fictional television detective Columbo (Chadwick, 2007). The psychotomimetic drug ketamine transiently induces delusions of reference in healthy volunteers (Krystal et al., 1994; Oye et al., 1992; Pomarol-Clotet et al., 2006). It blocks NMDA receptors (thus impairing the specification of top-down prior expectancies) while at the same time enhancing bottom-up AMPA signaling (Jackson et al., 2004) and engages acetylcholine release (Sarter et al., 2005). Low, sub psychotic doses of the drug engage the right frontostriatal prediction error signaling system in response to unsurprising and highly predictable events and the extent to which it does this showed a strong trend toward predicting the severity of heightened perception and delusional ideation (Corlett et al., 2006). We argue that delusions of reference form due to the attentional effects of aberrant prediction error (Pearce and Hall, 1980) mediated via surprise induced acetylcholine release from the nucleus basalis of meynert (Bao et al., 2001; Holland and Gallagher, 1993a, 1999a, 2006; Lee et al., 2005); subjects find their attention drawn toward irrelevant stimuli and events in the environment and impute personal meaning upon them, an experience that demands explanation, culminating in delusions of reference.

Corlett et al.

Paranoid ideation is associated with excessive fear or anxiety (Moutoussis *et al.*, 2007). In the context of the present analysis, paranoia would result when aberrant prediction error in frontostriatal learning systems engages the amygdala, engendering a feeling of fear and a state of hypervigilance. Relevant to this contention, delusions of reference and paranoid/ persecutory ideation tend to co-occur in patients with delusions (Startup and Startup, 2005), that is, hypervigilance and the perception of meaning in irrelevant and innocuous events may engender paranoia, since uncertainty and unpredictability are inherently fear inducing (Vinogradov *et al.*, 1992; Whitson and Galinsky, 2008). However, paranoid thoughts are commonly about other people (Melo *et al.*, 2006) and , as such, they may involve a prediction error driven dysfunction in the social learning mechanisms that we use to infer the intentions of others localized to frontostriatal and parietal circuits and the superior temporal sulcus/temporoparietal junction (Behrens *et al.*, 2009; Behrens *et al.*, 2008).

Physiological noise in this system, as a result of NMDA receptor hypofunction (which would disturb the specification of priors), AMPA receptor hyperfunction (which would signal prediction error where there should be none) and elevated dopamine levels within the mirror neuron circuit would impair the sufferer's ability to use what they have learned about their own actions and intentions to make inferences about other agents (Kilner et al., 2007a, b). Those disturbances in predicting and learning the consequences of our own actions may also have their origins in a disruption in the extended fronto-striatal-parietal reinforcement learning circuit; as we outlined, the midbrain dopamine neurons implicated in the pathophysiology of schizophrenia (Murray et al., 2008) may report an error in prediction, which is then processed in a circuit incorporating the frontal cortex, striatum and hippocampus (Redgrave and Gurney, 2006) as well as parietal cortex (Mittleman et al., 1990). This signal may be used to discern whether the organisms' actions caused a particular outcome, or whether the outcome happened due to external events (Redgrave and Gurney, 2006), while hypofunctioning of this circuit would lead to a decreased sense of agency for one's own actions, perhaps most relevant to delusions of external control (see below), we posit that the hyper engagement of this circuit could engender paranoia. That is, paranoid persecutory ideation is associated with superstitious biases in action-outcome learning (Kaney and Bentall, 1992). When playing rigged computer games paranoid individuals claimed to control both negative and positive outcomes when in fact there was no programmed contingency between their actions and the salient events.

Haggard and colleagues have reported an excessive binding between intentional actions and the outcomes they produce in patients with schizophrenia, however they did not relate this effect to delusions or paranoia in particular (Haggard et al., 2003). This maladaptive perception of contiguity between actions and outcomes would seem to offer an explanation for bizarre beliefs about telekinesis or enhanced predictive abilities, however in the context of the mirror neuron system account for computing and inferring the intentions of others (Kilner et al., 2007a, b), an individual who had learned spurious associations between their actions and salient environmental outcomes would also be expected to use those associations to infer the intentions of other agents. They would then ascribe supernatural abilities or excessively powerful status to individuals whom they encountered., In the context of prediction error induced amygdala responses, this inference would be affectively charged and result in a fear and distrust that is incommensurate with the current situation. This model makes some progress toward integrating neurobiology with psychodynamic explanations of paranoia (Kinderman and Bentall, 1996, 1997) in which attentional biases toward perceived threats are driven by mismatches between current self perceptions and how the patient believes they ought to be, focusing or projecting a threatening attributional bias onto external agents (Colby, 1977), patients with paranoia may attempt to avoid feelings of low self-esteem by attributing the cause of adverse experiences externally (Bentall et al., 2001). We believe that impairments in the brain's mirror neuron system and its ability to infer the

intentions of others based on inverting its own predictions (Kilner *et al.*, 2007a, b) may underpin these processes of inappropriate external projection of threat.

#### 6.2 Delusions of Motor Passivity

"My fingers pick up the pen, but I don't control them. What they do is nothing to do with me" (Mellor, 1970). It appears that these odd beliefs result from an impairment in the cancellation of predicted sensory consequences of motor behaviors (Blakemore, 2003; Blakemore *et al.*, 2003; Frith, 2005), involving a defect in the specification of motor predictions by the cerebellum which subsequently inappropriately engages parietal and frontal cortices (Frith, 2005; Schnell *et al.*, 2008; Spence *et al.*, 1997). An action produced without apparent forward model expectation is therefore ascribed to an external agent. A similar aberrant efference copy account has been made with respect to auditory hallucinations (Ford and Mathalon, 2005; Ford *et al.*, 2007; Heinks-Maldonado *et al.*, 2007).

However, some have criticized this model for failing to explain how patients with these delusions (and underlying brain pathology) can engage in any behavior at all. Having a sense of one's self as the source of our intentional actions may be essential for goal-directed instrumental learning (Glymour, 2004). This sense may be learned by the contiguous association between perceiving an intention to act, executing the motor program and encountering the consequences (Hume, 1739/2007; Wegner, 2004). Prediction errors due to physiological noise from dysregulated midbrain dopamine neurons projecting to prefrontal cortex could render those predictive associations unreliable (Corlett et al., 2007a). However, there is a less computationally intensive brain system that can control instrumental learning in the dorsolateral striatum. This system is said to mediate stimulus response habits (Daw et al., 2005; McDonald and Hong, 2004; Reading et al., 1991; Tang et al., 2007; Tricomi et al., 2009; Yin et al., 2004). The information used to guide behavior in this system is insensitive to the current value of the outcome (Daw et al., 2005). Habitual organisms behave reflexively, emitting motor responses to environmental cues irrespective of their consequences (Adams, 1981). The dorsal striatal habit system is believed to govern compulsive drug seeking and taking (Belin et al., 2009). The goal-directed and habit systems are conceived of as competitors for the control of behavior - the system that is least uncertain about the appropriate behavior given the context may win that competition (Daw et al., 2005). Competition between them can be biased towards the habit system by extended behavioral training (Adams, 1981); boosting synaptic dopamine levels in the striatum (Nelson and Killcross, 2006), or modulating AMPA receptor function (Bespalov et al., 2007). Goal directedness can be rescued by restoring dopamine induced plasticity in the prefrontal cortex (Hitchcott et al., 2007). It is possible that the habit system wins the competition in individuals with delusions (see below). Passivity experiences may therefore be explained as instrumental actions controlled by the habit system in the context of a noisy and inaccurate goal-directed system.

#### 6.3 Delusions of Parasitosis

Individuals with delusional parasitosis are convinced that small animals such as insects or lice are living on or within their skin (Berrios, 1982, 1985). This particular symptom highlights the overlap between delusions and hallucinations, perceptions and beliefs which calls in to question the strict clinical distinction (Corlett *et al.*, 2009a; Fletcher and Frith, 2009; Frith, 2000). Striatal lesions (Huber *et al.*, 2008), dopamine agonist medications (Charuvastra and Yaeger, 2006; Mitchell and Vierkant, 1991), cocaine (Mitchell and Vierkant, 1991; Siegel, 1978; Wallis, 1949) and amphetamine (Ellinwood, 1968; Ellinwood *et al.*, 1974) abuse can all engender delusions of parasitosis. Indeed, chronic treatment with dopamine antagonists can induce behaviors indicative of parasitosis in experimental animals (Ellison, 1994). In human stroke patients, delusions of parasitosis often occur following

lesions of right temporoparietal cortex, thalamus and putamen (Huber *et al.*, 2008). Putamen strongly influences visuotactile perception (Graziano and Gross, 1993; Ladavas *et al.*, 1998; Romo *et al.*, 1995; Yoo *et al.*, 2003), it contains bimodal cells with visual and tactile receptive fields, which help to encode the location of sensory stimuli mainly near the face. These cells project to parietal (ventral intraparietal cortex), primary somatosensory and premotor cortices (Graziano and Gross, 1993; Ladavas *et al.*, 1998).

We contend that sensations on the skin are a result of the same interaction between topdown and bottom-up mechanisms that we argue are crucial for visual perception. This is supported by the cutaneous rabbit illusion (Geldard and Sherrick, 1972) where simultaneous stimulation of two points on the skin gives rise to the percept of a rabbit 'hopping' between the two points; stimulation at a particular frequency is best explained by movement along a trajectory between the two points. There are Bayesian accounts of the illusion (Goldreich, 2007). Parasitosis may arise either due to bottom-up sensation that is normally ignored – for example a lack of adaptation of skin sensation over time or, alternatively, due to inappropriate top-down expectations – the power of cognition in cutaneous sensation is also underlined by contagious itch sensations experienced when subjects are exposed to conversations about insects on the skin (Heaven and McBrayer, 2000; Mitchell, 1995).

The same learning mechanisms that underpin the rubber hand illusion (Press *et al.*, 2008) might also be involved in parasitosis; a deficit in Bayesian multisensory integration would lead aberrant prediction error, driving attention toward potentially explanatory cues and forging inappropriate visuotactile associations. These associations, between sensation and a particular spatial location, might be represented by bimodal cells in the striatum, forming a new prior, a top-down bias in attention to the skin which would contribute to the maintenance of the delusion (Berrios, 1982; Corlett, 2009).

#### 6.4 Delusions of Misidentification

There are two main classes of misidentification delusion; Capgras; in which patients believe that their close family members have been replaced by imposters (Capgras, 1923), and Fregoli; in which patients believe that strangers that they encounter are their relatives in disguise (Courbon, 1927). Additionally, some patients have misidentification of their own home either feeling it is unfamiliar (Feinberg and Keenan, 2005; Fleminger, 1992) or that the hospital in which they find themselves is really their house hundreds of miles away (Schnider, 2001). Two factor models of these disorders assume a dual deficit, one in perception of affect, the other in belief evaluation (Coltheart et al., 2007). Instead, we argue that phenomenology of the percepts are such that bizarre beliefs are inevitable; surprising experiences demand surprising explanations (Kihlstrom, 1988). In our Bayesian, predictive learning scheme, Capgras results when patients experience an anomalous lack of affective responding when confronted with their relatives (Ellis and Young, 1990), the delusion constitutes a new prior driven by the experience, a means for explaining it away (Young, 2008). It is possible that the initial affective disturbance results from a failure to guide affect perception by prior experience, that is, just like sensory perception, emotions are predicted (Gilbert and Wilson, 2009); we have emotional priors, indeed, it is the prior expectancy of a familiar face combined with an emotional response (learned through experience) which breaks down in Capgras patients (Fleminger, 1992); fostering the misidentification of someone (or something) familiar as unfamiliar (Young, 2008). With the Fregoli delusion, it is a misplaced sense of familiarity (rather like a delusion of reference, specific to people) which guides patients to infer that people they do not know are actually their relatives in disguise.

In a meta-analysis of patients with delusional misidentification (Fregoli and Capgras delusions) about persons or objects, surveying 48 cases following neurological insult,

Feinberg et al found that the overwhelming majority had damage to the right hemisphere, commonly the frontal cortex. This observation is in line with our own work on prediction error during causal learning implicating a region of right dorsolateral prefrontal cortex in prediction error signaling (Corlett *et al.*, 2004; Fletcher *et al.*, 2001; Turner *et al.*, 2004) and implicating it in delusion formation (Corlett *et al.*, 2006; Corlett *et al.*, 2007b).

The laterality of damage that induces delusions seems replicable across studies of neurological patients with delusions (Devinsky, 2009). Spitzer and Walter (2003) speculate that this hemispheric bias can be explained by appealing to the different hemispheric modes of information processing (Kosslyn et al., 1992). Whereas the left hemisphere is characterized by smaller receptive fields resulting in focused, conjunctive coding, the right hemisphere is characterized by larger, overlapping receptive fields resulting in a coarse coding (Spitzer, 2003). In terms of Bayesian brain theory, receptive fields are related to the top-down specification of expected inputs (Rao and Ballard, 1999). Increasing dopamine levels may alter the signal to noise ratio of neurons, that is, it will increase the precision or certainty with which a prediction error is signaled (Friston et al., 2009) such that subjects respond to physiological noise as if it were meaningful signal (Grace, 1991). An increase in dopamine levels would serve to inappropriately increase confidence in noisy signals. It will therefore affect a system which relies on coarse coding, i.e. the right hemisphere, more prominently than a system which relies on conjunctive coding, i.e. the left hemisphere. That is, the right hemisphere is more susceptible to inappropriate optimization of prediction error because its predictions and prediction errors are inherently more noisy than the processing on the left hemisphere. Some speculate that, in response to right hemisphere error signals, the left hemisphere begins to construct explanations resulting in delusions (Devinsky, 2009), however the difficulty identifying and tracking delusions forming (Corlett et al., 2007a) means that this contention has not found empirical support.

When considering delusions of misidentification of neurological origin, it seems puzzling that damage in the same region could be associated with both an increase and a decrease in perceived familiarity. Two factor theorists would suggest that this is parsimoniously explained by ascribing the right frontal cortex the function of belief evaluation(Coltheart, 2010; Coltheart et al., 2007). However, we found that right frontal prediction error signal during causal learning was also related to ketamine induced perceptual aberrations (Corlett et al., 2006) and, furthermore, a study of individuals with lesions in the right dorsolateral prefrontal cortex suggested that lesion patients attended to and learned about irrelevant stimulus dimensions during a reward learning task (Hornak et al., 2004). It is possible that damage or dysfunction in prefrontal cortex could, paradoxically elevate activation in the remaining neurons since, in healthy individuals they provide a brake on subcortical dopamine nuclei through glutamatergic (Grace, 1991; Laruelle et al., 2003) and GABAergic mechanisms (Carlsson et al., 2001). Consequently, either due to a release from inhibition or an alteration of signal to noise properties, dopamine neurons projecting back from VTA to prefrontal cortex would increase in burst firing (Jackson et al., 2004) inducing rapid and random post-synaptic potentials in remaining functional cortical neurons (Lavin et al., 2005).

#### 6.5 Cotard Delusion

Perhaps one of the most bizarre delusions is the sufferer believing that they have died (Cotard, 1880), associated with claims that parts of them have "rotted away" or "disappeared" (Gerrans, 2002). It is possible that the same impoverished habitual mechanisms of instrumental action are engaged (see above) and the subject infers that the intentional agent that they were has disappeared, that is, the Cotard delusion may be a special case of passivity. Additionally, Some hypothesize that Capgras patients fail to recognize family members due a disconnection between face recognition units in the

fusiform face area and the ascription of emotional meaning in the limbic system, therefore, patients with Cotard delusion may have no connection at all between sensation and affective processing (Gerrans, 2002; Ramachandran, 1998). In this analysis, Cotard delusion is the converse of paranoia, instead of heightened and inappropriate emotional intensity it is a failure to ascribe emotional significance to any event (Gerrans, 2002; Ramachandran, 1998). Such a lack of emotional engagement with experiences would be surprising, engendering prediction error and sculpting the erroneous conclusion that the patient had died. Again, affective prediction fails, but instead of the rather specific effect in Capgras, it is a generalized failure in predicting the affective qualities of all sensory inputs. Like Capgras and Fregoli, this may involve a dysfunction in orbitofrontal cortex specifying top-down emotional predictions (Rolls and Grabenhorst, 2008). The delusion has been reported in a case study following right temporoparietal and bilateral frontal damage (Young et al., 1992), it also occurs in schizophrenia (Coltheart et al., 2007). This delusion involves both a deficit in affective forecasting (by the orbitofrontal cortex and amygdala), as well as (potentially) a deficit in motor forecasting (and thus sensory cancellation), with a diminished sense of self and emotional disengagement, the patient concludes that he/she is dead.

# 7. Why that odd belief? Individual differences in delusion susceptibility

While some psychotic patients get paranoid, others experience passivity, others still have multiple bizarre delusions. We posit a single factor, prediction error dysfunction for delusion formation and maintenance (Corlett *et al.*, 2009a; Corlett *et al.*, 2007a; Corlett, 2009; Fletcher and Frith, 2009). We have recently applied this single factor account to explain the range of phenomenological effects of pharmacologically distinct psychotomimetic drugs from dopamine agonist amphetamines, to NMDA antagonists, cannabinoids and serotonergic hallucinogens (Corlett *et al.*, 2009a). We believe the same explanation may be possible for the individual differences in susceptibility to different delusional themes observed in patients with schizophrenia.

Schizophrenia is a heritable but heterogeneous mental illness; its genetic inheritance appears to involve multiple genes of small effect (Tabares-Seisdedos and Rubenstein, 2009) or alternatively multiple rare genetic variants each with a large impact (Walsh et al., 2008). However, common to many of the identified risk genes for schizophrenia is a role in associative learning, prediction error signaling and NMDA receptor dependent synaptic plasticity (Hall et al., 2009; Stephan et al., 2006; Walsh et al., 2008). Some of the genes implicated in prediction error driven learning (Frank et al., 2007; Heyser et al., 2000) increase the risk for schizophrenia; the COMT val/met polymorphism may enhance maladaptive feedback between frontal cortex and subcortical dopamine neurons and is associated with risk for schizophrenia, aberrant salience and delusions (Bilder et al., 2004). In addition, PP1R1b, the gene coding for neostriatal signaling nexus DARPP-32 which integrates midbrain dopamine inputs with cortical glutamatergic signaling has been associated with prediction error driven learning (Frank et al., 2007; Heyser et al., 2000) frontostriatal structure and function as well as risk for schizophrenia (Meyer-Lindenberg et al., 2007). Variation in the function of these genes may explain inter-subject variability in susceptibility to delusions following psychotomimetic drug administration (Corlett et al., 2009a; Corlett et al., 2007a; Svenningsson et al., 2003).

However, different delusional themes are characteristic following the administration of different psychotomimetics; paranoia is more intense following cannabis administration (D'Souza *et al.*, 2009) whereas ketamine engenders delusions of reference (Krystal *et al.*, 1994; Oye *et al.*, 1992; Pomarol-Clotet *et al.*, 2006), although the two themes are by no means mutually exclusive (Startup and Startup, 2005). We believe that a second genetic insult may confer susceptibility to particular kinds of delusion in schizophrenia, an insult

involving disrupted cortical patterning and how the developing cortex interacts with environmental inputs in forming and maintaining cortical hierarchies (Sur and Rubenstein, 2005). Although this appears to be a two-factor theory, when we consider how Bayesian hierarchies like the brain develop into prediction engines (Friston, 2005b) through interactions between neural circuitry and incoming stimulation (Sur and Rubenstein, 2005), delusions really involve a singular dysfunction in predictive learning (i.e. an interaction between the two deficits which leads to (dys)interactions between poorly specified top-down predictions and noisy feedforward inputs; inducing aberrant and imprecise prediction errors (Corlett *et al.*, 2009a; Fletcher and Frith, 2009). The genes for building cortical hierarchies may also engender prediction error dysfunction irrespective of dopaminergic/glutamatergic 'prediction error' risk gene status and furthermore, the two insults may interact to produce more severe or varied delusions in the same patient.

Are there any empirical data to support of our contention that delusions with different themes are mediated by distinct (but overlapping) neural circuits? Patients with delusions secondary to neurological damage often have lesions in right frontal cortex but, according to two factor theories, the theme of the belief is conferred by damage to a second structure; for example the fusiform face area in Capgras delusion. Patients suffering from dementia with Lewy bodies experience delusions (Nagahama *et al.*, 2009; Nagahama *et al.*, 2007) like Capgras (Hirono and Cummings, 1999). Nagahama and colleagues used factor analysis to classify psychotic symptoms in dementia with Lewy bodies. They found that hallucinations, misidentification experiences and delusions were independent symptom domains (Nagahama *et al.*, 2007). More recently they replicated this factor structure in an independent group of patients and assessed the neural correlates of those factors by regressing factor scores onto resting state neuroimaging data across subjects (Nagahama *et al.*, 2009).

Patients suffering from misidentification had hypo-perfusion in left hippocampus, insula, inferior frontal gyrus and nucleus accumbens compared to patients without those symptoms. Individuals who had visual hallucinations of person or a feeling of presence had hypo-perfusion in bilateral parietal and left ventral occipital gyrus. Patients with persecutory delusions showed significant hyperactivity in right cingulate sulcus, bilateral middle frontal gyri, right inferior frontal gyrus, left medial superior frontal gyrus and left middle frontopolar gyrus. These distinct circuits tend to support our predicted delusion circuits (see Figure 4); that is, paranoia involves a frontal hyperactivity; delusions that potentially involve hyper salience of own body representations (e.g. hallucinations of people and feeling of presence) involve a parietal dysfunction and reduplications of person and place involve a predictive memory impairment; impaired familiarity processing and fronto-hippocampal as well as frontostriatal dysfunction.

Lewy bodies appear to accumulate in the space between bands of cortex; occupied by afferent or efferent connections with different cortical sites or with subcortical regions, that is, they have a laminar distribution (Armstrong *et al.*, 2001). Depending on which layer, they preferentially influence the feedforward (prediction error specifying) connections originating in laminae I-III and terminating in granular lamina IV of the adjacent lobe (Armstrong *et al.*, 2001). Alternatively, Lewy bodies may accumulate in the feedback fibers (responsible for specifying prior expectations and attentional modulation) which originate in laminae V and VI (and to some extent III) and terminate in lamina I (De Lacoste and White, 1993). Why the feedforward and feedback pathways of one particular circuit would be more sensitive to Lewy body inclusions than another circuit (conferring a particular delusion content) has yet to be determined, however, the disconnections that they engender within particular circuits is consistent with the putative disconnections invoked to explain the symptoms of schizophrenia (Friston, 2005a; Friston and Frith, 1995).

Finally, we turn to a rare but intriguing phenomenon, Folie a Deux (Lasegue, 1877), to evaluate our proposal that delusional themes are mediated by inherited biological processes. Folie a Deux (FD) is a psychotic disorder shared between two sufferers; an 'inducer' who initially develops the belief and the 'induced', an apparently otherwise healthy individual who comes to share the delusional belief. All kinds of rare delusional contents can be transmitted e.g. Cotard, Capgras, Fregoli (Wolff and McKenzie, 1994).

FD commonly occurs in persons who live close together, the delusion perhaps being transmitted through social learning processes. Additionally, if both patients are related, they may share the same genetically driven illness or predisposition. Monozygotic twins can share the same delusional themes (Lazarus, 1986); however, since they often share both genetic and environmental exposure, it is difficult to discern the unique contributions made by genes and environment. Scharfetter attempted to dissect these contributions by identifying dyads in whom there was no cosanginuity (e.g. husband and wife) then evaluating the risk for schizophrenia in each respective family. Incidence in both inducer and induced was very high (6.5% to 26.2%, compared with 1% population incidence), suggesting that a general predisposition toward delusions was necessary for accepting someone else's aberrant belief (Scharfetter, 1970). Future empirical research should investigate the personality, cognitive and neural functions of related and unrelated FD dyads to ascertain the roles of specific neural circuits in instantiating particular delusional beliefs.

# 8. Testing the hypothesis

Our sketch of the emerging neurobiology of delusional beliefs makes a number of testable predictions which will assess the validity of the venture:

- 1. We have argued that delusions arise and are maintained due to aberrations of glutamatergic synaptic plasticity, specifically chronically elevated synaptic glutamate which renders inappropriate salience and learning that engenders a limit on metaplasticity. Given its effectiveness against cocaine induced deficits in metaplasticity (Moussawi *et al.*, 2009), we predict that N-acetylcysteine should be an effective treatment for delusions.
- 2. Patients with delusional parasitosis and delusions of passivity should be more susceptible to the rubber hand illusion because of the dysinteraction between the bimodal cells in their striatum and parietal and cerebellar circuits responsible for coding top-down, motor expectancies and cancelling the sensory consequences of actions.
- **3.** Paranoia should be associated with prediction error dysfunction in mesocorticolimbic regions as well as the mirror neuron circuit, especially the superior temproral sulcus region involved in learning to infer the intentions of other agents (Behrens *et al.*, 2008).
- **4.** Given a large enough sample and phenomenologically rigorous assessment it should be possible to test our aetiological hypothesis about homeobox genes, development and the specification of priors; we predict that subtle variation in the gene coding for pax6 will alter amygdala development and therefore confer a risk for paranoia (Tole *et al.*, 2005); Engrailed 2 will be associated with an increased likelihood of cerebellar dysfunction (Sillitoe *et al.*, 2008) and as such will confer risk for passivity delusions.
- **5.** Reconsolidation processes should be enhanced in individuals with intractable delusions engaging and challenging their belief should increase its severity but treatments that block reconsolidation (such as the alpha adrenergic receptor

antagonist, propanalol) should ameliorate delusions (if they have been actively engaged).

- 6. Physical interventions that target reactivated representations of delusions should also have therapeutic benefits (Rubin, 1976), for example, it may be possible to disrupt the reconsolidation of delusions with transcranial magnetic stimulation (TMS) (Corlett, 2009). Based on the observed relationship between DLPFC dysfunction and delusional ideation (Corlett *et al.*, 2006; Corlett *et al.*, 2007b) as well as the role of DLPFC in controlled memory retrieval and updating (Fletcher and Henson, 2001) we suggest that specifically targeting that region with TMS following memory engagement may prove beneficial.
- 7. Individuals with high positive schizotypy or treated with psychotomimetic drugs should demonstrate aberrant prediction error signaling and therefore form learned habits more rapidly than controls.
- 8. If delusions are learned habits, then pharmacological interventions that restore goal directedness should be effective therapeutically; for example, antagonizing AMPA receptors (Bespalov *et al.*, 2007), boosting PFC dopamine levels (Hitchcott *et al.*, 2007) and attenuating striatal dopamine (Nelson and Killcross, 2006) should favor plasticity and goal directedness. The dopamine partial agonist Aripiprazole combines both antagonism of elevated striatal dopamine and an elevation of attenuated prefrontal dopamine and may specifically target aberrant prediction error signaling in midbrain dopamine neurons (Hamamura and Harada, 2007). It may be particularly effective against cognitive habits like delusions.

In order to complete a revolution of translation, having been inspired by the role of prediction error in associative learning in infrahuman species to develop our account of delusions, we should use invasive preclinical neuroscientific approaches in combination with associative learning phenomena to model delusion formation and maintenance in experimental animals. There are a number of potential opportunities here; combining acute psychotomimetic pharmacological models) to recapitulate putative neurobiological mechanisms of psychosis) with associative learning tasks that are sensitive to prediction error (to model delusion formation) or habit learning and memory reconsolidation (to model delusion maintenance).

# 9. Conclusion

We have outlined an account of delusional beliefs based on the tenets of animal learning theory and hierarchical Bayesian inference. We apply those tenets not only to explain dysfunctions in Pavlovian predictive learning (Corlett *et al.*, 2006; Corlett *et al.*, 2007b) and instrumental conditioning (Freeman *et al.*, 2009; Murray *et al.*, 2008; Roiser *et al.*, 2009; Schlagenhauf *et al.*, 2009), but also to account for the perceptual, affective and social disruptions that attend delusions (Bentall *et al.*, 2001; Maher, 1974; Vinogradov *et al.*, 1992).

In deluded individuals, the ability to use learned information to constrain current experience is impaired resulting aberrations of sensory and affective perception as well as cognition (Gray, 1991; Hemsley, 1994). Delusions may arise as an explanation for these odd happenings and they engage new learning (Kapur, 2003; Maher, 1974; McGhie and Chapman, 1961). They bring such relief that they are stamped into memory and become a new explanatory scheme for the sufferer (Jaspers, 1963), that is, delusions are elastic; they encompass new experiences and maintain a certain consistency of the world for the patient. In terms of the Bayesian model we outlined, delusions become the sufferer's new priors and they are used to predict and explain future experiences. We believe that the same prediction

error driven learning mechanisms can account for the fixity of delusional beliefs, since, now, when subsequent physiological noise elicits a reactivation of the delusion, it is reinforced and reconsolidated more strongly (Corlett *et al.*, 2009c). These hypotheses are readily testable in individuals suffering endogenous delusions, in healthy subjects exposed to psychotomimetic model psychoses and in preclinical models by focusing on the framework for translational cognitive neuroscience provided by formal associative learning theory, hierarchical Bayesian learning, predictive coding and information theory – the concept that intersects all of these is surprise or prediction error (Friston, 2010) and our model implicates aberrant prediction in the pathophysiology of delusions.

We have applied this model to various different kinds of delusions, examining beliefs that result from neurological damage as well as those that result from ingestion of psychotomimetic compounds and those that occur in schizophrenia, we feel, with some success. However, Brendan Maher, Emeritus Professor of the Psychology of Personality at Harvard, astutely aligned delusions with scientific theories (Maher, 1988), suggesting that scientists, like individuals with delusions, were extremely resistant to giving up their preferred theories even in the face damningly negative evidence. Like scientists, deluded individuals are confronted by surprising data which they explain away by abductive inference, generating hypotheses that explain away the surprise (Coltheart et al., 2010). Scientists (some of them at least) will engage in deductive inference to test the validity of their conclusions; whilst patients with delusions may not engage in this process (Miller, 1976), showing a bias against disconfirmatory evidence (Woodward et al., 2006). Furthermore, inductive inference, that is, reasoning from the specific to the general, has been invoked to explain the influence of prior experience over current perception (Barlow, 1990). We propose that the inductive process, reasoning beyond the data, may provide a mechanism through which delusions are maintained and pervade future experiences (Jaspers, 1963). Whilst the theory outlined in the present piece is our preferred explanation of delusions, we hope that we engender discussion, debate and investigation. As Maher says of science and psychosis: "Puzzles demand an explanation; the search for an explanation begins and continues until one has been devised'. We hope that this article might encourage others to join the search.

## References

- Aarts H, Custers R, Wegner DM. On the inference of personal authorship: enhancing experienced agency by priming effect information. Conscious Cogn. 2005; 14:439–458. [PubMed: 16091264]
- Abraham WC, Bear MF. Metaplasticity: the plasticity of synaptic plasticity. Trends Neurosci. 1996; 19:126–130. [PubMed: 8658594]
- Adams, CD.; Dickinson, A. Actions and Habits: variations in associative representations during instrumental learning. In: Spear, NE.; Miller, RR., editors. Information Processing in Animals: memory Mechanisms. Erlbaum; New Jersey: 1981.
- Allen G, McColl R, Barnard H, Ringe WK, Fleckenstein J, Cullum CM. Magnetic resonance imaging of cerebellar-prefrontal and cerebellar-parietal functional connectivity. Neuroimage. 2005; 28:39– 48. [PubMed: 16023375]
- Alloy LB, Tabachnik N. Assessment of covariation by humans and animals: the joint influence of prior expectations and current situational information. Psychol Rev. 1984; 91:112–149. [PubMed: 6571422]
- Angelucci A, Levitt JB, Lund JS. Anatomical origins of the classical receptive field and modulatory surround field of single neurons in macaque visual cortical area V1. Prog Brain Res. 2002a; 136:373–388. [PubMed: 12143395]
- Angelucci A, Levitt JB, Walton EJ, Hupe JM, Bullier J, Lund JS. Circuits for local and global signal integration in primary visual cortex. J Neurosci. 2002b; 22:8633–8646. [PubMed: 12351737]

- Angrilli A, Spironelli C, Elbert T, Crow TJ, Marano G, Stegagno L. Schizophrenia as failure of left hemispheric dominance for the phonological component of language. PLoS One. 2009; 4:e4507. [PubMed: 19223971]
- Ardid S, Wang XJ, Gomez-Cabrero D, Compte A. Reconciling coherent oscillation with modulation of irregular spiking activity in selective attention: gamma-range synchronization between sensory and executive cortical areas. J Neurosci. 2010; 30:2856–2870. [PubMed: 20181583]
- Armstrong RA, Cairns NJ, Lantos PL. What does the study of the spatial patterns of pathological lesions tell us about the pathogenesis of neurodegenerative disorders? Neuropathology. 2001; 21:1–12. [PubMed: 11304036]
- Atkinson JW, McClelland DC. The projective expression of needs. II. The effect of different intensities of the hunger drive on Thematic Apperception. Journal of Experimental Psychology. 1948; 38:643–658. [PubMed: 18893180]
- Baker DA, Madayag A, Kristiansen LV, Meador-Woodruff JH, Haroutunian V, Raju I. Contribution of cystine-glutamate antiporters to the psychotomimetic effects of phencyclidine. Neuropsychopharmacology. 2008; 33:1760–1772. [PubMed: 17728701]
- Balleine BW, Killcross S. Parallel incentive processing: an integrated view of amygdala function. Trends Neurosci. 2006; 29:272–279. [PubMed: 16545468]
- Bao S, Chan VT, Merzenich MM. Cortical remodelling induced by activity of ventral tegmental dopamine neurons. Nature. 2001; 412:79–83. [PubMed: 11452310]
- Barlow H. Conditions for versatile learning, Helmholtz's unconscious inference, and the task of perception. Vision Res. 1990; 30:1561–1571. [PubMed: 2288075]
- Bayes T. An essay towards solving a problem in the doctrine of chances. Philos Trans R Soc Lond. 1763; 53:370–418.
- Beck J, Beck J, Forstmeier W. Superstition and Belief as Inevitable ByProducts of an Adaptive Learning Strategy. Human Nature. 2007; 18:35.
- Behrens TE, Hunt LT, Rushworth MF. The computation of social behavior. Science. 2009; 324:1160–1164. [PubMed: 19478175]
- Behrens TE, Hunt LT, Woolrich MW, Rushworth MF. Associative learning of social value. Nature. 2008; 456:245–249. [PubMed: 19005555]
- Belin D, Jonkman S, Dickinson A, Robbins TW, Everitt BJ. Parallel and interactive learning processes within the basal ganglia: relevance for the understanding of addiction. Behav Brain Res. 2009; 199:89–102. [PubMed: 18950658]
- Belova MA, Paton JJ, Morrison SE, Salzman CD. Expectation modulates neural responses to pleasant and aversive stimuli in primate amygdala. Neuron. 2007; 55:970–984. [PubMed: 17880899]
- Bentall RP, Corcoran R, Howard R, Blackwood N, Kinderman P. Persecutory delusions: a review and theoretical integration. Clin Psychol Rev. 2001; 21:1143–1192. [PubMed: 11702511]
- Berke JD, Hetrick V, Breck J, Greene RW. Transient 23–30 Hz oscillations in mouse hippocampus during exploration of novel environments. Hippocampus. 2008; 18:519–529. [PubMed: 18398852]
- Berridge KC. The debate over dopamine's role in reward: the case for incentive salience. Psychopharmacology (Berl). 2007; 191:391–431. [PubMed: 17072591]
- Berrios GE. Tactile hallucinations: conceptual and historical aspects. J Neurol Neurosurg Psychiatry. 1982; 45:285–293. [PubMed: 7042917]
- Berrios GE. Delusional parasitosis and physical disease. Compr Psychiatry. 1985; 26:395–403. [PubMed: 4028691]
- Berrios GE. Delusions as "wrong beliefs": a conceptual history. Br J Psychiatry Suppl. 1991:6–13. [PubMed: 1840782]
- Bespalov AY, Harich S, Jongen-Relo AL, van Gaalen MM, Gross G. AMPA receptor antagonists reverse effects of extended habit training on signaled food approach responding in rats. Psychopharmacology (Berl). 2007; 195:11–18. [PubMed: 17634927]
- Bilder RM, Volavka J, Lachman HM, Grace AA. The catechol-O-methyltransferase polymorphism: relations to the tonic-phasic dopamine hypothesis and neuropsychiatric phenotypes. Neuropsychopharmacology. 2004; 29:1943–1961. [PubMed: 15305167]

- Binzegger T, Douglas RJ, Martin KA. A quantitative map of the circuit of cat primary visual cortex. J Neurosci. 2004; 24:8441–8453. [PubMed: 15456817]
- Bisiach E, Luzzatti C. Unilateral neglect of representational space. Cortex. 1978; 14:129–133. [PubMed: 16295118]
- Bissiere S, Humeau Y, Luthi A. Dopamine gates LTP induction in lateral amygdala by suppressing feedforward inhibition. Nat Neurosci. 2003; 6:587–592. [PubMed: 12740581]

Blakemore SJ. Deluding the motor system. Conscious Cogn. 2003; 12:647-655. [PubMed: 14656507]

- Blakemore SJ, Frith CD, Wolpert DM. Spatio-temporal prediction modulates the perception of selfproduced stimuli. J Cogn Neurosci. 1999; 11:551–559. [PubMed: 10511643]
- Blakemore SJ, Frith CD, Wolpert DM. The cerebellum is involved in predicting the sensory consequences of action. Neuroreport. 2001; 12:1879–1884. [PubMed: 11435916]
- Blakemore SJ, Oakley DA, Frith CD. Delusions of alien control in the normal brain. Neuropsychologia. 2003; 41:1058–1067. [PubMed: 12667541]
- Blakemore SJ, Smith J, Steel R, Johnstone CE, Frith CD. The perception of self-produced sensory stimuli in patients with auditory hallucinations and passivity experiences: evidence for a breakdown in self-monitoring. Psychol Med. 2000a; 30:1131–1139. [PubMed: 12027049]
- Blakemore SJ, Wolpert D, Frith C. Why can't you tickle yourself? Neuroreport. 2000b; 11:R11–16. [PubMed: 10943682]
- Blakemore SJ, Wolpert DM, Frith CD. Abnormalities in the awareness of action. Trends Cogn Sci. 2002; 6:237–242. [PubMed: 12039604]
- Bleuler E. Die Prognose der Dementia praecox (Schizophreniegruppe). Allgemeine Zeitschrift für Psychiatrie und psychischgerichtliche Medizin. 1908; 65:436–464.
- Bottini G, Bisiach E, Sterzi R, Vallar G. Feeling touches in someone else's hand. Neuroreport. 2002; 13:249–252. [PubMed: 11893919]
- Botvinick M, Cohen J. Rubber hands 'feel' touch that eyes see. Nature. 1998; 391:756. [PubMed: 9486643]
- Braver TS, Cohen JD. Dopamine, cognitive control, and schizophrenia: the gating model. Prog Brain Res. 1999; 121:327–349. [PubMed: 10551035]
- Bromberg-Martin ES, Hikosaka O. Midbrain dopamine neurons signal preference for advance information about upcoming rewards. Neuron. 2009; 63:119–126. [PubMed: 19607797]
- Bruner J, Bruner J, Postman L. Perception, cognition, and behavior. Journal of Personality. 1949; 18:14.
- Buhl EH, Tamas G, Fisahn A. Cholinergic activation and tonic excitation induce persistent gamma oscillations in mouse somatosensory cortex in vitro. J Physiol. 1998; 513 (Pt 1):117–126. [PubMed: 9782163]
- Buschman TJ, Miller EK. Top-down versus bottom-up control of attention in the prefrontal and posterior parietal cortices. Science. 2007; 315:1860–1862. [PubMed: 17395832]
- Buzsaki G. The structure of consciousness. Nature. 2007; 446:267. [PubMed: 17361165]
- Capgras J, Reboul-Lachaux J. L'illusion des "soises" dans un delire systematise. Bulletin de Society Clinique de Medicine Mentale. 1923; 11:6–16.
- Carlsson A, Waters N, Holm-Waters S, Tedroff J, Nilsson M, Carlsson ML. Interactions between monoamines, glutamate, and GABA in schizophrenia: new evidence. Annu Rev Pharmacol Toxicol. 2001; 41:237–260. [PubMed: 11264457]
- Carlsson M, Carlsson A. Schizophrenia: a subcortical neurotransmitter imbalance syndrome? Schizophr Bull. 1990; 16:425–432. [PubMed: 1981107]
- Cepeda C, Levine MS. Dopamine and N-methyl-D-aspartate receptor interactions in the neostriatum. Dev Neurosci. 1998; 20:1–18. [PubMed: 9600386]
- Chadwick PK. Peer-professional first-person account: schizophrenia from the inside--phenomenology and the integration of causes and meanings. Schizophr Bull. 2007; 33:166–173. [PubMed: 16973785]
- Charuvastra A, Yaeger D. Tactile hallucinations associated with therapeutic doses of bupropion in 2 patients. J Clin Psychiatry. 2006; 67:1820–1821. [PubMed: 17196068]

- Chiu YF, McGrath JA, Thornquist MH, Wolyniec PS, Nestadt G, Swartz KL, Lasseter VK, Liang KY, Pulver AE. Genetic heterogeneity in schizophrenia II: conditional analyses of affected schizophrenia sibling pairs provide evidence for an interaction between markers on chromosome 8p and 14q. Mol Psychiatry. 2002; 7:658–664. [PubMed: 12140791]
- Chong TT, Cunnington R, Williams MA, Kanwisher N, Mattingley JB. fMRI adaptation reveals mirror neurons in human inferior parietal cortex. Curr Biol. 2008; 18:1576–1580. [PubMed: 18948009]
- Clifford W. The ethics of belief. Contemporary Review. 1877
- Colby KM. Appraisal of four psychological theories of paranoid phenomena. J Abnorm Psychol. 1977; 86:54–59. [PubMed: 838949]
- Coltheart M. The neuropsychology of delusions. Ann N Y Acad Sci. 2010; 1191:16–26. [PubMed: 20392273]
- Coltheart M, Langdon R, McKay R. Schizophrenia and monothematic delusions. Schizophr Bull. 2007; 33:642–647. [PubMed: 17372282]
- Coltheart M, Menzies P, Sutton J. Abductive inference and delusional belief. Cogn Neuropsychiatry. 2010; 15:261–287. [PubMed: 20017038]
- Conrad, K. Die Beginnende Schizophrenie. G. Thieme; Stuttgart: 1958a.
- Conrad, K. Die BeginnendeSchizophrenie. G. Thieme; Stuttgart: 1958b.
- Corlett PR, Aitken MR, Dickinson A, Shanks DR, Honey GD, Honey RA, Robbins TW, Bullmore ET, Fletcher PC. Prediction error during retrospective revaluation of causal associations in humans: fMRI evidence in favor of an associative model of learning. Neuron. 2004; 44:877–888. [PubMed: 15572117]
- Corlett PR, Frith CD, Fletcher PC. From drugs to deprivation: a Bayesian framework for understanding models of psychosis. Psychopharmacology (Berl). 2009a
- Corlett PR, Frith CD, Fletcher PC. From drugs to deprivation: a Bayesian framework for understanding models of psychosis. Psychopharmacology (Berl). 2009b; 206:515–530. [PubMed: 19475401]
- Corlett PR, Honey GD, Aitken MR, Dickinson A, Shanks DR, Absalom AR, Lee M, Pomarol-Clotet E, Murray GK, McKenna PJ, Robbins TW, Bullmore ET, Fletcher PC. Frontal responses during learning predict vulnerability to the psychotogenic effects of ketamine: linking cognition, brain activity, and psychosis. Arch Gen Psychiatry. 2006; 63:611–621. [PubMed: 16754834]
- Corlett PR, Honey GD, Fletcher PC. From prediction error to psychosis: ketamine as a pharmacological model of delusions. J Psychopharmacol. 2007a; 21:238–252. [PubMed: 17591652]
- Corlett PR, Krystal JH, Taylor JR, Fletcher PC. Why do delusions persist? Front Hum Neurosci. 2009c; 3:12. [PubMed: 19636384]
- Corlett PR, Krystal JK, Taylor JR, Fletcher PC. Why do delusions persist? Frontiers in Human Neuroscience. 2009 In press.
- Corlett PR, Murray GK, Honey GD, Aitken MR, Shanks DR, Robbins TW, Bullmore ET, Dickinson A, Fletcher PC. Disrupted prediction-error signal in psychosis: evidence for an associative account of delusions. Brain. 2007b; 130:2387–2400. [PubMed: 17690132]
- Corlett PR, Simons JS, Pigott JS, Gardner JM, Murray GK, Krystal JH, Fletcher PC. Illusions and delusions: relating experimentally-induced false memories to anomalous experiences and ideas. Front Behav Neurosci. 2009d; 3:53. [PubMed: 19956402]
- Cotard J. Du délire hypocondriaque dans une forme grave de la melancolie anxieuse. Memoire lu à la Société médicopsychologique dans la Séance du 28 Juin 1880. Ann Medico-Psychol Med. 1880:168–174.
- Courbon P, Fail G. Syndrome d' "illusion de Frégoli" et schizophrénie. Bulletin de la Société Clinique de Médecine Mentale. 1927; 15:121–124.
- Courville AC, Daw ND, Touretzky DS. Bayesian theories of conditioning in a changing world. Trends Cogn Sci. 2006; 10:294–300. [PubMed: 16793323]
- Cramer RE, Weiss RF, William R, Reid S, Nieri L, Manning-Ryan B. Human agency and associative learning: Pavlovian principles govern social process in causal relationship detection. Q J Exp Psychol B. 2002; 55:241–266. [PubMed: 12188526]

- Critchley HD, Mathias CJ, Dolan RJ. Fear conditioning in humans: the influence of awareness and autonomic arousal on functional neuroanatomy. Neuron. 2002; 33:653–663. [PubMed: 11856537]
- D'Souza DC, Sewell RA, Ranganathan M. Cannabis and psychosis/schizophrenia: human studies. Eur Arch Psychiatry Clin Neurosci. 2009; 259:413–431. [PubMed: 19609589]
- Dakin S, Carlin P, Hemsley D. Weak suppression of visual context in chronic schizophrenia. Curr Biol. 2005; 15:R822–824. [PubMed: 16243017]
- Damasio, A. Thinking about Belief: Concluding Remarks. In: Schacter, DL.; Scarry, E., editors. Memory, Brain and Belief. Harvard University Press; Cambridge Massachussetts / London, England: 2000.
- Daw ND, Niv Y, Dayan P. Uncertainty-based competition between prefrontal and dorsolateral striatal systems for behavioral control. Nat Neurosci. 2005; 8:1704–1711. [PubMed: 16286932]
- De Clerambault GG. Les Psychoses Passionelles. Oeuvere Psychiatrique:Paris: Presses Universities, de France. 1942; 331:337–339. 357, 408.
- De Lacoste MC, White CL 3rd. The role of cortical connectivity in Alzheimer's disease pathogenesis: a review and model system. Neurobiol Aging. 1993; 14:1–16. [PubMed: 8450928]
- Delespaul P, van Os J. Jaspers was right after all--delusions are distinct from normal beliefs. Against. Br J Psychiatry. 2003; 183:286. [PubMed: 14575030]
- Delgado MR, Li J, Schiller D, Phelps EA. The role of the striatum in aversive learning and aversive prediction errors. Philos Trans R Soc Lond B Biol Sci. 2008a; 363:3787–3800. [PubMed: 18829426]
- Delgado MR, Nearing KI, Ledoux JE, Phelps EA. Neural circuitry underlying the regulation of conditioned fear and its relation to extinction. Neuron. 2008b; 59:829–838. [PubMed: 18786365]
- den Ouden HE, Daunizeau J, Roiser J, Friston KJ, Stephan KE. Striatal prediction error modulates cortical coupling. J Neurosci. 2010; 30:3210–3219. [PubMed: 20203180]
- den Ouden HE, Friston KJ, Daw ND, McIntosh AR, Stephan KE. A dual role for prediction error in associative learning. Cereb Cortex. 2009; 19:1175–1185. [PubMed: 18820290]
- Dennett, D. Do animals have beliefs. In: Roitblat, HL.; Meyer, JA., editors. Comparative Approaches to Cognitive Science. MIT Press; Cambrudge Massachusetts; London, England: 1995.
- Devenport LD. Superstitious bar pressing in hippocampal and septal rats. Science. 1979; 205:721–723. [PubMed: 462183]
- Devinsky O. Delusional misidentifications and duplications: right brain lesions, left brain delusions. Neurology. 2009; 72:80–87. [PubMed: 19122035]
- Dickinson A. The 28th Bartlett Memorial Lecture. Causal learning: an associative analysis. Q J Exp Psychol B. 2001; 54:3–25. [PubMed: 11216300]
- Dima D, Roiser JP, Dietrich DE, Bonnemann C, Lanfermann H, Emrich HM, Dillo W. Understanding why patients with schizophrenia do not perceive the hollow-mask illusion using dynamic causal modelling. Neuroimage. 2009; 46:1180–1186. [PubMed: 19327402]
- Dommett E, Coizet V, Blaha CD, Martindale J, Lefebvre V, Walton N, Mayhew JE, Overton PG, Redgrave P. How visual stimuli activate dopaminergic neurons at short latency. Science. 2005; 307:1476–1479. [PubMed: 15746431]
- Eichenbaum, H.; Bodkin, JA. Belief and knowledge as distinct forms of memory. In: Schacter, DL.; Scarry, E., editors. Memory Brain and Belief. Harvard University Press; Cambridge, Massachusetts/London England: 2000.
- Eisenhardt D, Menzel R. Extinction learning, reconsolidation and the internal reinforcement hypothesis. Neurobiol Learn Mem. 2007; 87:167–173. [PubMed: 17079171]
- Ellinwood EH Jr. Amphetamine psychosis. II. Theoretical implications. Int J Neuropsychiatry. 1968; 4:45–54. [PubMed: 5645819]
- Ellinwood EH Jr, Sudilovsky A, Nelson LM. Behavior and EEG analysis of chronic amphetamine effect. Biol Psychiatry. 1974; 8:169–176. [PubMed: 4858385]
- Ellis HD, Young AW. Accounting for delusional misidentifications. Br J Psychiatry. 1990; 157:239– 248. [PubMed: 2224375]
- Ellison DG. Hallucinations produced by sensory conditioning. Journal of Experimental Psychology. 1941; 28:1–20.

- Ellison G. Stimulant-induced psychosis, the dopamine theory of schizophrenia, and the habenula. Brain Res Brain Res Rev. 1994; 19:223–239. [PubMed: 7914793]
- Emrich HM. A three-component-system hypothesis of psychosis. Impairment of binocular depth inversion as an indicator of a functional dysequilibrium. Br J Psychiatry Suppl. 1989:37–39. [PubMed: 2690888]
- Estes WK. Processes of memory loss, recovery, and distortion. Psychol Rev. 1997; 104:148–169. [PubMed: 9009883]
- Farrer C, Frey SH, Van Horn JD, Tunik E, Turk D, Inati S, Grafton ST. The angular gyrus computes action awareness representations. Cereb Cortex. 2008; 18:254–261. [PubMed: 17490989]
- Feinberg TE. The nested heirarchy of consciousness: A neurobiological soultion to the problem of mental unity. Neurocase. 2000; 6:75–81.
- Feinberg TE, Keenan JP. Where in the brain is the self? Conscious Cogn. 2005; 14:661–678. [PubMed: 16325140]
- Fiorillo CD. Towards a general theory of neural computation based on prediction by single neurons. PLoS ONE. 2008; 3:e3298. [PubMed: 18827880]
- Fiorillo CD, Tobler PN, Schultz W. Discrete coding of reward probability and uncertainty by dopamine neurons. Science. 2003; 299:1898–1902. [PubMed: 12649484]
- Fiser J. Perceptual learning and representational learning in humans and animals. Learn Behav. 2009; 37:141–153. [PubMed: 19380891]
- Fiser J, Berkes P, Orban G, Lengyel M. Statistically optimal perception and learning: from behavior to neural representations. Trends Cogn Sci. 2010; 14:119–130. [PubMed: 20153683]
- Fleminger S. Seeing is believing: the role of 'preconscious' perceptual processing in delusional misidentification. Br J Psychiatry. 1992; 160:293–303. [PubMed: 1562856]
- Fletcher PC, Anderson JM, Shanks DR, Honey R, Carpenter TA, Donovan T, Papadakis N, Bullmore ET. Responses of human frontal cortex to surprising events are predicted by formal associative learning theory. Nat Neurosci. 2001; 4:1043–1048. [PubMed: 11559855]
- Fletcher PC, Frith CD. Perceiving is believing: a Bayesian approach to explaining the positive symptoms of schizophrenia. Nat Rev Neurosci. 2009; 10:48–58. [PubMed: 19050712]
- Fletcher PC, Henson RN. Frontal lobes and human memory: insights from functional neuroimaging. Brain. 2001; 124:849–881. [PubMed: 11335690]
- Flint AJ. Delusions in dementia: a review. J Neuropsychiatry Clin Neurosci. 1991; 3:121–130. [PubMed: 1687962]
- Fogassi L, Luppino G. Motor functions of the parietal lobe. Curr Opin Neurobiol. 2005; 15:626–631. [PubMed: 16271458]
- Ford JM, Mathalon DH. Corollary discharge dysfunction in schizophrenia: can it explain auditory hallucinations? Int J Psychophysiol. 2005; 58:179–189. [PubMed: 16137779]
- Ford JM, Roach BJ, Faustman WO, Mathalon DH. Synch before you speak: auditory hallucinations in schizophrenia. Am J Psychiatry. 2007; 164:458–466. [PubMed: 17329471]
- Fourneret P, Paillard J, Lamarre Y, Cole J, Jeannerod M. Lack of conscious recognition of one's own actions in a haptically deafferented patient. Neuroreport. 2002; 13:541–547. [PubMed: 11930177]
- Franck N, Posada A, Pichon S, Haggard P. Altered subjective time of events in schizophrenia. J Nerv Ment Dis. 2005; 193:350–353. [PubMed: 15870620]
- Frank MJ, Moustafa AA, Haughey HM, Curran T, Hutchison KE. Genetic triple dissociation reveals multiple roles for dopamine in reinforcement learning. Proc Natl Acad Sci U S A. 2007; 104:16311–16316. [PubMed: 17913879]
- Freeman D, Garety PA, Kuipers E, Fowler D, Bebbington PE. A cognitive model of persecutory delusions. Br J Clin Psychol. 2002; 41:331–347. [PubMed: 12437789]
- Freeman D, Garety PA, Kuipers E, Fowler D, Bebbington PE, Dunn G. Acting on persecutory delusions: the importance of safety seeking. Behav Res Ther. 2007; 45:89–99. [PubMed: 16530161]

- Freeman TP, Morgan CJ, Klaassen E, Das RK, Stefanovic A, Brandner B, Curran HV. Superstitious conditioning as a model of delusion formation following chronic but not acute ketamine in humans. Psychopharmacology (Berl). 2009
- Friston K. Disconnection and cognitive dysmetria in schizophrenia. Am J Psychiatry. 2005a; 162:429– 432. [PubMed: 15741456]
- Friston K. A theory of cortical responses. Philos Trans R Soc Lond B Biol Sci. 2005b; 360:815–836. [PubMed: 15937014]
- Friston K. The free-energy principle: a rough guide to the brain? Trends Cogn Sci. 2009; 13:293–301. [PubMed: 19559644]
- Friston K. The free-energy principle: a unified brain theory? Nat Rev Neurosci. 2010; 11:127–138. [PubMed: 20068583]
- Friston KJ. Hallucinations and perceptual inherence. Behavioral and Brain Sciences. 2005c; 28:764–766.
- Friston KJ, Daunizeau J, Kiebel SJ. Reinforcement learning or active inference? PLoS One. 2009; 4:e6421. [PubMed: 19641614]
- Friston KJ, Frith CD. Schizophrenia: a disconnection syndrome? Clin Neurosci. 1995; 3:89–97. [PubMed: 7583624]
- Frith C. The neural basis of hallucinations and delusions. C R Biol. 2005; 328:169–175. [PubMed: 15771003]
- Frith CD, Blakemore S, Wolpert DM. Explaining the symptoms of schizophrenia: abnormalities in the awareness of action. Brain Res Brain Res Rev. 2000a; 31:357–363. [PubMed: 10719163]
- Frith CD, Blakemore SJ, Wolpert DM. Abnormalities in the awareness and control of action. Philos Trans R Soc Lond B Biol Sci. 2000b; 355:1771–1788. [PubMed: 11205340]
- Frith, CD.; Dolan, RJ. The role of memory in the delusions associated with schizophrenia. In: Schacher, D.; Scarry, E., editors. Memory, Brain and Belief. Harvard University Press; 2000.
- Fuster JM. The prefrontal cortex--an update: time is of the essence. Neuron. 2001; 30:319–333. [PubMed: 11394996]
- Gallese V, Fadiga L, Fogassi L, Rizzolatti G. Action recognition in the premotor cortex. Brain. 1996; 119 ( Pt 2):593–609. [PubMed: 8800951]
- Gallese V, Keysers C, Rizzolatti G. A unifying view of the basis of social cognition. Trends Cogn Sci. 2004; 8:396–403. [PubMed: 15350240]
- Garety P. Reasoning and delusions. Br J Psychiatry Suppl. 1991:14–18. [PubMed: 1840774]
- Garety PA, Freeman D. Cognitive approaches to delusions: a critical review of theories and evidence. Br J Clin Psychol. 1999; 38 ( Pt 2):113–154. [PubMed: 10389596]
- Geldard FA, Sherrick CE. The cutaneous "rabbit": a perceptual illusion. Science. 1972; 178:178–179. [PubMed: 5076909]
- Gerrans P. A one-stage explanation of the Cotard delusion. Philosophy, Psychiatry & Psychology. 2002; 9:47–53.
- Geyer MA, Vollenweider FX. Serotonin research: contributions to understanding psychoses. Trends Pharmacol Sci. 2008; 29:445–453. [PubMed: 19086254]
- Ghoreishi A. A somatic type delusional disorder secondary to peripheral neuropathy: a case report. Psychiatria Danubina. 2010; 20:85–87. [PubMed: 18376336]
- Gilbert DT, Wilson TD. Why the brain talks to itself: sources of error in emotional prediction. Philos Trans R Soc Lond B Biol Sci. 2009; 364:1335–1341. [PubMed: 19528015]
- Glymour C. We believe in freedom of the will so we can learn [Comment on Wegner, Precis of the Illusion of Conscious Will. Behavioral and Brain Sciences. 2004; 27:661–662.
- Goldreich D. A Bayesian perceptual model replicates the cutaneous rabbit and other tactile spatiotemporal illusions. PLoS One. 2007; 2:e333. [PubMed: 17389923]
- Goldstone R. Effects of categorization on color perception. Psychological Science. 1995; 6:298-304.
- Goto Y, O'Donnell P. Network synchrony in the nucleus accumbens in vivo. J Neurosci. 2001; 21:4498–4504. [PubMed: 11404437]

Corlett et al.

- Gourion D, Leroy S, Bourdel MC, Goldberger C, Poirier MF, Olie JP, Krebs MO. Cerebellum development and schizophrenia: an association study of the human homeogene Engrailed 2. Psychiatry Res. 2004; 126:93–98. [PubMed: 15123388]
- Grabenhorst F, Rolls ET, Bilderbeck A. How cognition modulates affective responses to taste and flavor: top-down influences on the orbitofrontal and pregenual cingulate cortices. Cereb Cortex. 2008; 18:1549–1559. [PubMed: 18056086]
- Grace AA. Phasic versus tonic dopamine release and the modulation of dopamine system responsivity: a hypothesis for the etiology of schizophrenia. Neuroscience. 1991; 41:1–24. [PubMed: 1676137]
- Grace AA, Floresco SB, Goto Y, Lodge DJ. Regulation of firing of dopaminergic neurons and control of goal-directed behaviors. Trends Neurosci. 2007; 30:220–227. [PubMed: 17400299]
- Gray JA. On biology, phenomenology, and pharmacology in schizophrenia. Am J Psychiatry. 2004; 161:377. author reply 377–378. [PubMed: 14754801]
- Gray JA, Feldon J, Rawlins JNP, Hemsley D, Smith AD. The Neuropsychology of Schizophrenia. Behav Brain Sci. 1991; 14:1–84.
- Graziano MS, Gross CG. A bimodal map of space: somatosensory receptive fields in the macaque putamen with corresponding visual receptive fields. Exp Brain Res. 1993; 97:96–109. [PubMed: 8131835]
- Grossberg S. Processing of expected and unexpected events during conditioning and attention: a psychophysiological theory. Psychol Rev. 1982; 89:529–572. [PubMed: 7178332]
- Grossberg S. How hallucinations may arise from brain mechanisms of learning, attention, and volition. J Int Neuropsychol Soc. 2000; 6:583–592. [PubMed: 10932478]
- Grossberg S. Cortical and subcortical predictive dynamics and learning during perception, cognition, emotion and action. Philos Trans R Soc Lond B Biol Sci. 2009; 364:1223–1234. [PubMed: 19528003]
- Haber SN. The primate basal ganglia: parallel and integrative networks. J Chem Neuroanat. 2003; 26:317–330. [PubMed: 14729134]
- Haber SN, Kim KS, Mailly P, Calzavara R. Reward-related cortical inputs define a large striatal region in primates that interface with associative cortical connections, providing a substrate for incentive-based learning. J Neurosci. 2006; 26:8368–8376. [PubMed: 16899732]
- Haggard P, Martin F, Taylor-Clarke M, Jeannerod M, Franck N. Awareness of action in schizophrenia. Neuroreport. 2003; 14:1081–1085. [PubMed: 12802207]
- Haider B, Duque A, Hasenstaub AR, McCormick DA. Neocortical network activity in vivo is generated through a dynamic balance of excitation and inhibition. J Neurosci. 2006; 26:4535– 4545. [PubMed: 16641233]
- Hall J, Romaniuk L, McIntosh AM, Steele JD, Johnstone EC, Lawrie SM. Associative learning and the genetics of schizophrenia. Trends Neurosci. 2009; 32:359–365. [PubMed: 19427043]
- Halligan PW, David AS. Cognitive neuropsychiatry: towards a scientific psychopathology. Nat Rev Neurosci. 2001; 2:209–215. [PubMed: 11256082]
- Hamamura T, Harada T. Unique pharmacological profile of aripiprazole as the phasic component buster. Psychopharmacology (Berl). 2007; 191:741–743. [PubMed: 17205315]
- Hampton AN, Bossaerts P, O'Doherty JP. Neural correlates of mentalizing-related computations during strategic interactions in humans. Proc Natl Acad Sci U S A. 2008; 105:6741–6746. [PubMed: 18427116]
- Hasselmo ME, Rolls ET, Baylis GC. The role of expression and identity in the face-selective responses of neurons in the temporal visual cortex of the monkey. Behav Brain Res. 1989a; 32:203–218. [PubMed: 2713076]
- Hasselmo ME, Rolls ET, Baylis GC, Nalwa V. Object-centered encoding by face-selective neurons in the cortex in the superior temporal sulcus of the monkey. Exp Brain Res. 1989b; 75:417–429. [PubMed: 2721619]
- Heaven L, McBrayer D. External motivators of self-touching behavior. Percept Mot Skills. 2000; 90:338–342. [PubMed: 10769920]
- Hebb, DO. The Organization of Behavior. John Wiley; 1949a.
- Hebb, DO. The organization of behaviour: a neuropsychological theory. Wiley; New York: 1949b.

- Heinks-Maldonado TH, Mathalon DH, Houde JF, Gray M, Faustman WO, Ford JM. Relationship of imprecise corollary discharge in schizophrenia to auditory hallucinations. Arch Gen Psychiatry. 2007; 64:286–296. [PubMed: 17339517]
- von Helmholtz, H. The Facts of Perception. In: Kahl, R., editor. Selected Writings of Herman von Helmholtz. Weslyan University Press; 1878/1971.
- Hemsley, DR. Perceptual and cognitive abnormalities as the basis for schizophrenic symptoms. In: David, AS.; Cutting, JC., editors. The Neuropsychology of Schizophrenia. Laurence Erlbaum Associates; Hove, UK: 1994. p. 97-118.
- Hemsley DR, Garety PA. The formation and maintenance of delusions: a Bayesian analysis. Br J Psychiatry. 1986a; 149:51–56. [PubMed: 3779313]
- Hemsley DR, Garety PA. The formation of maintenance of delusions: a Bayesian analysis. Br J Psychiatry. 1986b; 149:51–56. [PubMed: 3779313]
- Hendricks, KV.; Wiggers, P.; Jonker, CM.; Haselager, WF. Towards a computational model of the self-attribution of agency. In: Oliver, P.; Kray, C., editors. the artificial intelligence and simulation of behaviour annual convention. 2007. p. 350-356.
- Herrero JL, Roberts MJ, Delicato LS, Gieselmann MA, Dayan P, Thiele A. Acetylcholine contributes through muscarinic receptors to attentional modulation in V1. Nature. 2008; 454:1110–1114. [PubMed: 18633352]
- Heyes C. Mesmerising mirror neurons. Neuroimage. 2010; 51:789-791. [PubMed: 20167276]
- Heyser CJ, Fienberg AA, Greengard P, Gold LH. DARPP-32 knockout mice exhibit impaired reversal learning in a discriminated operant task. Brain Res. 2000; 867:122–130. [PubMed: 10837805]
- Hikosaka O, Bromberg-Martin E, Hong S, Matsumoto M. New insights on the subcortical representation of reward. Curr Opin Neurobiol. 2008a; 18:203–208. [PubMed: 18674617]
- Hikosaka O, Sesack SR, Lecourtier L, Shepard PD. Habenula: crossroad between the basal ganglia and the limbic system. J Neurosci. 2008b; 28:11825–11829. [PubMed: 19005047]
- Hirono N, Cummings JL. Neuropsychiatric aspects of dementia with Lewy bodies. Curr Psychiatry Rep. 1999; 1:85–92. [PubMed: 11122909]
- Hitchcott PK, Quinn JJ, Taylor JR. Bidirectional modulation of goal-directed actions by prefrontal cortical dopamine. Cereb Cortex. 2007; 17:2820–2827. [PubMed: 17322558]
- Hoffman RE, Dobscha SK. Cortical pruning and the development of schizophrenia: a computer model. Schizophr Bull. 1989; 15:477–490. [PubMed: 2814376]
- Holland PC, Gallagher M. Amygdala central nucleus lesions disrupt increments, but not decrements, in conditioned stimulus processing. Behav Neurosci. 1993a; 107:246–253. [PubMed: 8484890]
- Holland PC, Gallagher M. Effects of amygdala central nucleus lesions on blocking and unblocking. Behav Neurosci. 1993b; 107:235–245. [PubMed: 8484889]
- Holland PC, Gallagher M. Amygdala circuitry in attentional and representational processes. Trends Cogn Sci. 1999; 3:65–73. [PubMed: 10234229]
- Holland PC, Gallagher M. Different roles for amygdala central nucleus and substantia innominata in the surprise-induced enhancement of learning. J Neurosci. 2006; 26:3791–3797. [PubMed: 16597732]
- Holt DJ, Lebron-Milad K, Milad MR, Rauch SL, Pitman RK, Orr SP, Cassidy BS, Walsh JP, Goff DC. Extinction Memory Is Impaired in Schizophrenia. Biol Psychiatry. 2008
- Hornak J, O'Doherty J, Bramham J, Rolls ET, Morris RG, Bullock PR, Polkey CE. Reward-related reversal learning after surgical excisions in orbito-frontal or dorsolateral prefrontal cortex in humans. J Cogn Neurosci. 2004; 16:463–478. [PubMed: 15072681]
- Houran J, Houran J. Preliminary study of tolerance of ambiguity of individuals reporting paranormal experiences. Psychological Reports. 1998; 82:183. [PubMed: 9520551]
- Howes OD, Montgomery AJ, Asselin MC, Murray RM, Valli I, Tabraham P, Bramon-Bosch E, Valmaggia L, Johns L, Broome M, McGuire PK, Grasby PM. Elevated striatal dopamine function linked to prodromal signs of schizophrenia. Arch Gen Psychiatry. 2009; 66:13–20. [PubMed: 19124684]

- Huber M, Karner M, Kirchler E, Lepping P, Freudenmann RW. Striatal lesions in delusional parasitosis revealed by magnetic resonance imaging. Prog Neuropsychopharmacol Biol Psychiatry. 2008; 32:1967–1971. [PubMed: 18930778]
- Hudson LA, Rollins YD, Anderson CA, Johnston-Brooks C, Tyler KL, Filley CM. Reduplicative paramnesia in Morvan's syndrome. J Neurol Sci. 2008; 267:154–157. [PubMed: 17928004]
- Hume, D. A treatise of human nature. Oxford University Press; Oxford: 1739/2007.
- Ito M. Movement and thought: identical control mechanisms by the cerebellum. Trends Neurosci. 1993; 16:448–450. discussion 453–444. [PubMed: 7507615]
- Jackson ME, Homayoun H, Moghaddam B. NMDA receptor hypofunction produces concomitant firing rate potentiation and burst activity reduction in the prefrontal cortex. Proc Natl Acad Sci U S A. 2004; 101:8467–8472. [PubMed: 15159546]
- Jaspers, K. General Psychopathology. Manchester University Press; 1963.
- Jeannerod M. The hand and the object: the role of posterior parietal cortex in forming motor representations. Can J Physiol Pharmacol. 1994; 72:535–541. [PubMed: 7954083]
- Joliot M, Ribary U, Llinas R. Human oscillatory brain activity near 40 Hz coexists with cognitive temporal binding. Proc Natl Acad Sci U S A. 1994; 91:11748–11751. [PubMed: 7972135]
- Jones H. Defining delusion. Br J Psychiatry. 2004; 185:354–355. [PubMed: 15459001]
- Kalayasiri R, Sughondhabirom A, Gueorguieva R, Coric V, Lynch WJ, Morgan PT, Cubells JF, Malison RT. Self-reported paranoia during laboratory "binge" cocaine self-administration in humans. Pharmacol Biochem Behav. 2006; 83:249–256. [PubMed: 16549106]
- Kandel ER. A new intellectual framework for psychiatry. Am J Psychiatry. 1998; 155:457–469. [PubMed: 9545989]
- Kaney S, Bentall RP. Persecutory delusions and the self-serving bias. Evidence from a contingency judgment task. J Nerv Ment Dis. 1992; 180:773–780. [PubMed: 1469376]
- Kapur S. Psychosis as a state of aberrant salience: a framework linking biology, phenomenology, and pharmacology in schizophrenia. Am J Psychiatry. 2003; 160:13–23. [PubMed: 12505794]
- Karson CN. A new look at delusions of grandeur. Compr Psychiatry. 1980; 21:62–69. [PubMed: 7357864]
- Keinan G, Keinan G. Effects of stress and tolerance of ambiguity on magical thinking. Journal of Personality and Social Psychology. 1994; 67:48.
- Keysers C, Wicker B, Gazzola V, Anton JL, Fogassi L, Gallese V. A touching sight: SII/PV activation during the observation and experience of touch. Neuron. 2004; 42:335–346. [PubMed: 15091347]
- Kihlstrom, JF.; Hoyt, IP. Hypnosis and the Psychology of Delusions. In: Oltmanns, TF.; Maher, BA., editors. Delusional Beliefs. Wiley; New York: 1988.
- Kilner JM, Friston KJ, Frith CD. The mirror-neuron system: a Bayesian perspective. Neuroreport. 2007a; 18:619–623. [PubMed: 17413668]
- Kilner JM, Friston KJ, Frith CD. Predictive coding: an account of the mirror neuron system. Cogn Process. 2007b; 8:159–166. [PubMed: 17429704]
- Kilner JM, Neal A, Weiskopf N, Friston KJ, Frith CD. Evidence of mirror neurons in human inferior frontal gyrus. J Neurosci. 2009; 29:10153–10159. [PubMed: 19675249]
- Kinderman P, Bentall RP. Self-discrepancies and persecutory delusions: evidence for a model of paranoid ideation. J Abnorm Psychol. 1996; 105:106–113. [PubMed: 8666699]
- Kinderman P, Bentall RP. Causal attributions in paranoia and depression: internal, personal, and situational attributions for negative events. J Abnorm Psychol. 1997; 106:341–345. [PubMed: 9131855]
- King R, Barchas JD, Huberman BA. Chaotic behavior in dopamine neurodynamics. Proc Natl Acad Sci U S A. 1984; 81:1244–1247. [PubMed: 6583705]
- Knobel A, Heinz A, Voss M. Imaging the deluded brain. Eur Arch Psychiatry Clin Neurosci. 2008; 258(Suppl 5):76–80. [PubMed: 18985300]
- Koechlin E, Anton JL, Burnod Y. Dynamical computational properties of local cortical networks for visual and motor processing: a bayesian framework. J Physiol Paris. 1996; 90:257–262. [PubMed: 9116679]

Konorski, J. Conditioned reflexes and neuron organization. Cambridge University Press; 1948.

- Kosslyn SM, Chabris CF, Marsolek CJ, Koenig O. Categorical versus coordinate spatial relations: computational analyses and computer simulations. J Exp Psychol Hum Percept Perform. 1992; 18:562–577. [PubMed: 1593235]
- Kot T, Serper M. Increased susceptibility to auditory conditioning in hallucinating schizophrenic patients: a preliminary investigation. J Nerv Ment Dis. 2002; 190:282–288. [PubMed: 12011606]
- Kraeplin, E. Clinical Psychiatry. MacMillan; New York: 1902.
- Kromkamp M, Uylings HB, Smidt MP, Hellemons AJ, Burbach JP, Kahn RS. Decreased thalamic expression of the homeobox gene DLX1 in psychosis. Arch Gen Psychiatry. 2003; 60:869–874. [PubMed: 12963668]
- Krystal JH, Karper LP, Seibyl JP, Freeman GK, Delaney R, Bremner JD, Heninger GR, Bowers MB Jr, Charney DS. Subanesthetic effects of the noncompetitive NMDA antagonist, ketamine, in humans. Psychotomimetic, perceptual, cognitive, and neuroendocrine responses. Arch Gen Psychiatry. 1994; 51:199–214. [PubMed: 8122957]
- Kurylo DD, Gazes Y. Effects of Ketamine on perceptual grouping in rats. Physiol Behav. 2008; 95:152–156. [PubMed: 18571682]
- Ladavas E, Zeloni G, Farne A. Visual peripersonal space centred on the face in humans. Brain. 1998; 121 (Pt 12):2317–2326. [PubMed: 9874482]
- Lange R, Lange R, Houran J. Delusions of the paranormal: A haunting question of perception. Journal of Nervous and Mental Disease. 1998; 186:637. [PubMed: 9788641]
- Lapish CC, Seamans JK, Chandler LJ. Glutamate-dopamine cotransmission and reward processing in addiction. Alcohol Clin Exp Res. 2006; 30:1451–1465. [PubMed: 16930207]
- Laruelle M, Kegeles LS, Abi-Dargham A. Glutamate, dopamine, and schizophrenia: from pathophysiology to treatment. Ann N Y Acad Sci. 2003; 1003:138–158. [PubMed: 14684442]
- Lasegue C, Falret J. La folie a deux (ou folie communiquee). Ann Med Psychol. 1877; 18:321-355.
- Lau HC, Rogers RD, Passingham RE. Manipulating the experienced onset of intention after action execution. J Cogn Neurosci. 2007; 19:81–90. [PubMed: 17214565]
- Lavin A, Nogueira L, Lapish CC, Wightman RM, Phillips PE, Seamans JK. Mesocortical dopamine neurons operate in distinct temporal domains using multimodal signaling. J Neurosci. 2005; 25:5013–5023. [PubMed: 15901782]
- Laviolette SR, Grace AA. The roles of cannabinoid and dopamine receptor systems in neural emotional learning circuits: implications for schizophrenia and addiction. Cell Mol Life Sci. 2006; 63:1597–1613. [PubMed: 16699809]
- Lazarus A. Folie a deux in identical twins: interaction of nature and nurture. Br J Psychiatry. 1986; 148:324–326. [PubMed: 3719226]
- Lecourtier L, Defrancesco A, Moghaddam B. Differential tonic influence of lateral habenula on prefrontal cortex and nucleus accumbens dopamine release. Eur J Neurosci. 2008; 27:1755–1762. [PubMed: 18380670]
- Lee D. Game theory and neural basis of social decision making. Nat Neurosci. 2008a; 11:404–409. [PubMed: 18368047]
- Lee HJ, Groshek F, Petrovich GD, Cantalini JP, Gallagher M, Holland PC. Role of amygdalo-nigral circuitry in conditioning of a visual stimulus paired with food. J Neurosci. 2005; 25:3881–3888. [PubMed: 15829640]
- Lee JL. Memory reconsolidation mediates the strengthening of memories by additional learning. Nat Neurosci. 2008b; 11:1264–1266. [PubMed: 18849987]
- Lingnau A, Gesierich B, Caramazza A. Asymmetric fMRI adaptation reveals no evidence for mirror neurons in humans. Proc Natl Acad Sci U S A. 2009; 106:9925–9930. [PubMed: 19497880]
- Lisman J, Buzsaki G. A neural coding scheme formed by the combined function of gamma and theta oscillations. Schizophr Bull. 2008; 34:974–980. [PubMed: 18559405]
- Lisman J, Redish AD. Prediction, sequences and the hippocampus. Philos Trans R Soc Lond B Biol Sci. 2009; 364:1193–1201. [PubMed: 19528000]
- Lisman JE, Grace AA. The hippocampal-VTA loop: controlling the entry of information into longterm memory. Neuron. 2005; 46:703–713. [PubMed: 15924857]

- Llinas RR, Roy S. The 'prediction imperative' as the basis for self-awareness. Philos Trans R Soc Lond B Biol Sci. 2009; 364:1301–1307. [PubMed: 19528011]
- Lodge DJ, Grace AA. The hippocampus modulates dopamine neuron responsivity by regulating the intensity of phasic neuron activation. Neuropsychopharmacology. 2006a; 31:1356–1361. [PubMed: 16319915]
- Lodge DJ, Grace AA. The laterodorsal tegmentum is essential for burst firing of ventral tegmental area dopamine neurons. Proc Natl Acad Sci U S A. 2006b; 103:5167–5172. [PubMed: 16549786]
- Loh M, Rolls ET, Deco G. A dynamical systems hypothesis of schizophrenia. PLoS Comput Biol. 2007; 3:e228. [PubMed: 17997599]
- Long JE, Swan C, Liang WS, Cobos I, Potter GB, Rubenstein JL. Dlx1&2 and Mash1 transcription factors control striatal patterning and differentiation through parallel and overlapping pathways. J Comp Neurol. 2009; 512:556–572. [PubMed: 19030180]
- Ma WJ, Beck JM, Latham PE, Pouget A. Bayesian inference with probabilistic population codes. Nat Neurosci. 2006; 9:1432–1438. [PubMed: 17057707]
- Mackintosh NJ. A theory of attention: Variations in the associability of stimuli with reinforcement. Psychological Review. 1975:82.
- Maher BA. Delusional thinking and perceptual disorder. J Individ Psychol. 1974; 30:98–113. [PubMed: 4857199]
- Maher, BA. Delusions as normal theories. Wiley; New York: 1988.
- Makin TR, Holmes NP, Ehrsson HH. On the other hand: dummy hands and peripersonal space. Behav Brain Res. 2008; 191:1–10. [PubMed: 18423906]
- Matsumoto M, Hikosaka O. Two types of dopamine neuron distinctly convey positive and negative motivational signals. Nature. 2009; 459:837–841. [PubMed: 19448610]
- Maxwell SL, Li M. Midbrain dopaminergic development in vivo and in vitro from embryonic stem cells. J Anat. 2005; 207:209–218. [PubMed: 16185245]
- McCurdy HG. Coin perception studies and the concept of schemata. Psychological Review. 1956; 63:160–168. [PubMed: 13323171]
- McDonald RJ, Hong NS. A dissociation of dorso-lateral striatum and amygdala function on the same stimulus-response habit task. Neuroscience. 2004; 124:507–513. [PubMed: 14980722]
- McGhie A, Chapman J. Disorders of attention and perception in early schizophrenia. Br J Med Psychol. 1961; 34:103–116. [PubMed: 13773940]
- McKay R, Langdon R, Coltheart M. Models of misbelief: Integrating motivational and deficit theories of delusions. Conscious Cogn. 2007; 16:932–941. [PubMed: 17331741]
- McLaren IP, Dickinson A. The conditioning connection. Philos Trans R Soc Lond B Biol Sci. 1990; 329:179–186. [PubMed: 1978363]
- Mellor CS. First rank symptoms of schizophrenia. I. The frequency in schizophrenics on admission to hospital. II. Differences between individual first rank symptoms. Br J Psychiatry. 1970; 117:15– 23. [PubMed: 5479324]
- Melo SS, Taylor JL, Bentall RP. 'Poor me' versus 'bad me' paranoia and the instability of persecutory ideation. Psychol Psychother. 2006; 79:271–287. [PubMed: 16774723]
- Mesulam M. Representation, inference, and transcendent encoding in neurocognitive networks of the human brain. Ann Neurol. 2008; 64:367–378. [PubMed: 18991346]
- Meyer-Lindenberg A, Straub RE, Lipska BK, Verchinski BA, Goldberg T, Callicott JH, Egan MF, Huffaker SS, Mattay VS, Kolachana B, Kleinman JE, Weinberger DR. Genetic evidence implicating DARPP-32 in human frontostriatal structure, function, and cognition. J Clin Invest. 2007; 117:672–682. [PubMed: 17290303]
- Milad MR, Quirk GJ, Pitman RK, Orr SP, Fischl B, Rauch SL. A role for the human dorsal anterior cingulate cortex in fear expression. Biol Psychiatry. 2007; 62:1191–1194. [PubMed: 17707349]
- Milad MR, Vidal-Gonzalez I, Quirk GJ. Electrical stimulation of medial prefrontal cortex reduces conditioned fear in a temporally specific manner. Behav Neurosci. 2004; 118:389–394. [PubMed: 15113265]

- Miller, NE. Liberalization of basic S-R concepts: Extensions to conflict behaviour, motivation and social learning. In: Kock, S., editor. Psychology: A study of a science. McGraw-Hill; New York: 1959. p. 196-292.
- Miller R. Schizophrenic psychology, associative learning and the role of forebrain dopamine. Med Hypotheses. 1976; 2:203–211. [PubMed: 9558]
- Miltner WH, Braun C, Arnold M, Witte H, Taub E. Coherence of gamma-band EEG activity as a basis for associative learning. Nature. 1999; 397:434–436. [PubMed: 9989409]
- Milton F, Patwa VK, Hafner RJ. Confrontation vs. belief modification in persistently deluded patients. Br J Med Psychol. 1978; 51:127–130. [PubMed: 646958]
- Misanin JR, Miller RR, Lewis DJ. Retrograde amnesia produced by electroconvulsive shock after reactivation of a consolidated memory trace. Science. 1968; 160:554–555. [PubMed: 5689415]
- Mishara AL. Is minimal self preserved in schizophrenia? A subcomponents view. Conscious Cogn. 2007; 16:715–721. [PubMed: 17920523]
- Mishara AL, Corlett PR. Are delusions biologically adaptive? Salvaging the doxastic shear pin. Behavioral and Brain Sciences. 2009; 32:530–531.
- Mitchell CW. Effects of subliminally presented auditory suggestions of itching on scratching behavior. Percept Mot Skills. 1995; 80:87–96. [PubMed: 7624224]
- Mitchell J, Vierkant AD. Delusions and hallucinations of cocaine abusers and paranoid schizophrenics: a comparative study. J Psychol. 1991; 125:301–310. [PubMed: 1880755]
- Mittleman G, Whishaw IQ, Jones GH, Koch M, Robbins TW. Cortical, hippocampal, and striatal mediation of schedule-induced behaviors. Behav Neurosci. 1990; 104:399–409. [PubMed: 2354035]
- Mohanty A, Egner T, Monti JM, Mesulam MM. Search for a threatening target triggers limbic guidance of spatial attention. J Neurosci. 2009; 29:10563–10572. [PubMed: 19710309]
- Mohr F, Hubmann W, Cohen R, Bender W, Haslacher C, Honicke S, Schlenker R, Wahlheim C, Werther P. Neurological soft signs in schizophrenia: assessment and correlates. Eur Arch Psychiatry Clin Neurosci. 1996; 246:240–248. [PubMed: 8863002]
- Montague PR, Dayan P, Sejnowski TJ. A framework for mesencephalic dopamine systems based on predictive Hebbian learning. J Neurosci. 1996; 16:1936–1947. [PubMed: 8774460]
- Moore JW, Wegner DM, Haggard P. Modulating the sense of agency with external cues. Conscious Cogn. 2009
- Moritz S, Woodward TS, Chen E. Investigation of metamemory dysfunctions in first-episode schizophrenia. Schizophr Res. 2006; 81:247–252. [PubMed: 16256310]
- Morris JS, deBonis M, Dolan RJ. Human amygdala responses to fearful eyes. Neuroimage. 2002; 17:214–222. [PubMed: 12482078]
- Moussawi K, Pacchioni A, Moran M, Olive MF, Gass JT, Lavin A, Kalivas PW. N-Acetylcysteine reverses cocaine-induced metaplasticity. Nat Neurosci. 2009; 12:182–189. [PubMed: 19136971]
- Moutoussis M, Williams J, Dayan P, Bentall RP. Persecutory delusions and the conditioned avoidance paradigm: towards an integration of the psychology and biology of paranoia. Cogn Neuropsychiatry. 2007; 12:495–510. [PubMed: 17978936]
- Murray GK, Corlett PR, Clark L, Pessiglione M, Blackwell AD, Honey G, Jones PB, Bullmore ET, Robbins TW, Fletcher PC. Substantia nigra/ventral tegmental reward prediction error disruption in psychosis. Mol Psychiatry. 2008; 13:239, 267–276. [PubMed: 17684497]
- Nader K, Schafe GE, Le Doux JE. Fear memories require protein synthesis in the amygdala for reconsolidation after retrieval. Nature. 2000; 406:722–726. [PubMed: 10963596]
- Nagahama Y, Okina T, Suzuki N, Matsuda M. Neural correlates of psychotic symptoms in dementia with Lewy bodies. Brain. 2009
- Nagahama Y, Okina T, Suzuki N, Matsuda M, Fukao K, Murai T. Classification of psychotic symptoms in dementia with Lewy bodies. Am J Geriatr Psychiatry. 2007; 15:961–967. [PubMed: 17974867]
- Nelson A, Killcross S. Amphetamine exposure enhances habit formation. J Neurosci. 2006; 26:3805–3812. [PubMed: 16597734]

- O'Donnell P. Dopamine gating of forebrain neural ensembles. Eur J Neurosci. 2003; 17:429–435. [PubMed: 12581161]
- Owen G, Harland R, Antonova E, Broome M. Jaspers' concept of primary delusion. Br J Psychiatry. 2004; 185:77–78. [PubMed: 15231560]
- Oye I, Paulsen O, Maurset A. Effects of ketamine on sensory perception: evidence for a role of Nmethyl-D-aspartate receptors. J Pharmacol Exp Ther. 1992; 260:1209–1213. [PubMed: 1312163]
- Pally R. The predicting brain: unconscious repetition, conscious reflection and therapeutic change. Int J Psychoanal. 2007; 88:861–881. [PubMed: 17681897]
- Pan WX, Schmidt R, Wickens JR, Hyland BI. Tripartite mechanism of extinction suggested by dopamine neuron activity and temporal difference model. J Neurosci. 2008; 28:9619–9631. [PubMed: 18815248]
- Parthasarathi UD, Harrower T, Tempest M, Hodges JR, Walsh C, McKenna PJ, Fletcher PC. Psychiatric presentation of voltage-gated potassium channel antibody-associated encephalopathy. Case report. Br J Psychiatry. 2006; 189:182–183. [PubMed: 16880491]
- Passie T, Karst M, Borsutzky M, Wiese B, Emrich HM, Schneider U. Effects of different subanaesthetic doses of (S)-ketamine on psychopathology and binocular depth inversion in man. J Psychopharmacol. 2003; 17:51–56. [PubMed: 12680739]
- Paton JJ, Belova MA, Morrison SE, Salzman CD. The primate amygdala represents the positive and negative value of visual stimuli during learning. Nature. 2006; 439:865–870. [PubMed: 16482160]
- Pearce JM, Hall G. A model for Pavlovian learning: variations in the effectiveness of conditioned but not of unconditioned stimuli. Psychol Rev. 1980; 87:532–552. [PubMed: 7443916]
- Peled A, Pressman A, Geva AB, Modai I. Somatosensory evoked potentials during a rubber-hand illusion in schizophrenia. Schizophr Res. 2003; 64:157–163. [PubMed: 14613680]
- Pennartz CM, McNaughton BL, Mulder AB. The glutamate hypothesis of reinforcement learning. Prog Brain Res. 2000; 126:231–253. [PubMed: 11105650]
- Pessiglione M, Seymour B, Flandin G, Dolan RJ, Frith CD. Dopamine-dependent prediction errors underpin reward-seeking behaviour in humans. Nature. 2006; 442:1042–1045. [PubMed: 16929307]
- Phillips WA, Silverstein SM. Convergence of biological and psychological perspectives on cognitive coordination in schizophrenia. Behav Brain Sci. 2003; 26:65–82. discussion 82–137. [PubMed: 14598440]
- Pomarol-Clotet E, Honey GD, Murray GK, Corlett PR, Absalom AR, Lee M, McKenna PJ, Bullmore ET, Fletcher PC. Psychological effects of ketamine in healthy volunteers. Phenomenological study. Br J Psychiatry. 2006; 189:173–179. [PubMed: 16880489]
- Press C, Heyes C, Haggard P, Eimer M. Visuotactile learning and body representation: an ERP study with rubber hands and rubber objects. J Cogn Neurosci. 2008; 20:312–323. [PubMed: 18275337]
- Preuschoff K, Bossaerts P, Quartz SR. Neural differentiation of expected reward and risk in human subcortical structures. Neuron. 2006; 51:381–390. [PubMed: 16880132]
- Ramachandran, V.; Blakeslee, S. Phantoms in the Brain: Probing the mysteries of the human mind. William Morrow; New York: 1998.
- Rao RP, Ballard DH. Predictive coding in the visual cortex: a functional interpretation of some extraclassical receptive-field effects. Nat Neurosci. 1999; 2:79–87. [PubMed: 10195184]
- Ravina B, Marder K, Fernandez HH, Friedman JH, McDonald W, Murphy D, Aarsland D, Babcock D, Cummings J, Endicott J, Factor S, Galpern W, Lees A, Marsh L, Stacy M, Gwinn-Hardy K, Voon V, Goetz C. Diagnostic criteria for psychosis in Parkinson's disease: report of an NINDS, NIMH work group. Mov Disord. 2007; 22:1061–1068. [PubMed: 17266092]
- Reading PJ, Dunnett SB, Robbins TW. Dissociable roles of the ventral, medial and lateral striatum on the acquisition and performance of a complex visual stimulus-response habit. Behav Brain Res. 1991; 45:147–161. [PubMed: 1789923]
- Redgrave P, Gurney K. The short-latency dopamine signal: a role in discovering novel actions? Nat Rev Neurosci. 2006; 7:967–975. [PubMed: 17115078]
- Redish AD. Addiction as a computational process gone awry. Science. 2004; 306:1944–1947. [PubMed: 15591205]

- Rescorla, RA.; Wagner, AR. A theory of Pavlovian conditioning: Variations in the effectiveness of reinforcement and non-reinforcement. In: Black, AH.; Prokasy, WF., editors. Classical conditioning II: Current research and theory. Appleton-Century-Crofts; New York: 1972.
- Ridler K, Veijola JM, Tanskanen P, Miettunen J, Chitnis X, Suckling J, Murray GK, Haapea M, Jones PB, Isohanni MK, Bullmore ET. Fronto-cerebellar systems are associated with infant motor and adult executive functions in healthy adults but not in schizophrenia. Proc Natl Acad Sci U S A. 2006; 103:15651–15656. [PubMed: 17028177]
- Rizzolatti G, Fadiga L, Gallese V, Fogassi L. Premotor cortex and the recognition of motor actions. Brain Res Cogn Brain Res. 1996; 3:131–141. [PubMed: 8713554]
- Roiser JP, Stephan KE, den Ouden HE, Barnes TR, Friston KJ, Joyce EM. Do patients with schizophrenia exhibit aberrant salience? Psychol Med. 2009; 39:199–209. [PubMed: 18588739]
- Rolls ET. The representation of information about faces in the temporal and frontal lobes. Neuropsychologia. 2007; 45:124–143. [PubMed: 16797609]
- Rolls ET, Grabenhorst F. The orbitofrontal cortex and beyond: from affect to decision-making. Prog Neurobiol. 2008; 86:216–244. [PubMed: 18824074]
- Rolls ET, Grabenhorst F, Margot C, da Silva MA, Velazco MI. Selective attention to affective value alters how the brain processes olfactory stimuli. J Cogn Neurosci. 2008; 20:1815–1826. [PubMed: 18370603]
- Romo R, Merchant H, Ruiz S, Crespo P, Zainos A. Neuronal activity of primate putamen during categorical perception of somaesthetic stimuli. Neuroreport. 1995; 6:1013–1017. [PubMed: 7632884]
- Ros H, Sachdev RN, Yu Y, Sestan N, McCormick DA. Neocortical networks entrain neuronal circuits in cerebellar cortex. J Neurosci. 2009; 29:10309–10320. [PubMed: 19692605]
- Rubin RD. Clinical use of retrograde amnesia produced by electroconvulsive shock. A conditioning hypothesis. Can Psychiatr Assoc J. 1976; 21:87–90. [PubMed: 1277097]
- Sanchez-Vives MV, McCormick DA. Cellular and network mechanisms of rhythmic recurrent activity in neocortex. Nat Neurosci. 2000; 3:1027–1034. [PubMed: 11017176]
- Sander D, Grafman J, Zalla T. The human amygdala: an evolved system for relevance detection. Rev Neurosci. 2003; 14:303–316. [PubMed: 14640318]
- Sarter M, Nelson CL, Bruno JP. Cortical cholinergic transmission and cortical information processing in schizophrenia. Schizophr Bull. 2005; 31:117–138. [PubMed: 15888431]
- Sass LA. Some Reflections on the (Analytic) Philosophical Approach to Delusion. Philosophy, Psychiatry & Psychology. 2004; 11:71–80.
- Scearce-Levie K, Roberson ED, Gerstein H, Cholfin JA, Mandiyan VS, Shah NM, Rubenstein JL, Mucke L. Abnormal social behaviors in mice lacking Fgf17. Genes Brain Behav. 2008; 7:344– 354. [PubMed: 17908176]
- Scharfetter C. On the hereditary aspects of symbiontic psychoses. A contribution towards the understanding of the schizophrenia-like psychoses. Psychiatr Clin (Basel). 1970; 3:145–152. [PubMed: 5422677]
- Schiller D, Levy I, Niv Y, LeDoux JE, Phelps EA. From fear to safety and back: reversal of fear in the human brain. J Neurosci. 2008; 28:11517–11525. [PubMed: 18987188]
- Schlagenhauf F, Sterzer P, Schmack K, Ballmaier M, Rapp M, Wrase J, Juckel G, Gallinat J, Heinz A. Reward feedback alterations in unmedicated schizophrenia patients: relevance for delusions. Biol Psychiatry. 2009; 65:1032–1039. [PubMed: 19195646]
- Schneider, K. Clinical Psychopathology. Grune & Stratton; New York: 1959.
- Schnell K, Heekeren K, Daumann J, Schnell T, Schnitker R, Moller-Hartmann W, Gouzoulis-Mayfrank E. Correlation of passivity symptoms and dysfunctional visuomotor action monitoring in psychosis. Brain. 2008; 131:2783–2797. [PubMed: 18713781]
- Schnider A. Spontaneous confabulation, reality monitoring, and the limbic system--a review. Brain Res Brain Res Rev. 2001; 36:150–160. [PubMed: 11690611]
- Schnider A. Spontaneous confabulation and the adaptation of thought to ongoing reality. Nat Rev Neurosci. 2003; 4:662–671. [PubMed: 12894241]

- Schobel SA, Lewandowski NM, Corcoran CM, Moore H, Brown T, Malaspina D, Small SA. Differential targeting of the CA1 subfield of the hippocampal formation by schizophrenia and related psychotic disorders. Arch Gen Psychiatry. 2009; 66:938–946. [PubMed: 19736350]
- Schultz W. Predictive reward signal of dopamine neurons. J Neurophysiol. 1998; 80:1–27. [PubMed: 9658025]
- Schultz W, Apicella P, Ljungberg T. Responses of monkey dopamine neurons to reward and conditioned stimuli during successive steps of learning a delayed response task. J Neurosci. 1993; 13:900–913. [PubMed: 8441015]
- Schultz W, Dayan P, Montague PR. A neural substrate of prediction and reward. Science. 1997; 275:1593–1599. [PubMed: 9054347]
- Schultz W, Dickinson A. Neuronal coding of prediction errors. Annu Rev Neurosci. 2000; 23:473– 500. [PubMed: 10845072]
- Schultz W, Dickinson A. Neural coding of prediction errors. Annual Review of Neuroscience. 2000:473–500.
- Seamans JK, Yang CR. The principal features and mechanisms of dopamine modulation in the prefrontal cortex. Prog Neurobiol. 2004; 74:1–58. [PubMed: 15381316]
- Seashore CE. Measurements of illusions and hallucinations in normal life. Studies from the Yale Psychological Laboratory. 1895; 3:1–67.
- Sesack SR, Grace AA. Cortico-Basal Ganglia reward network: microcircuitry. Neuropsychopharmacology. 2010; 35:27–47. [PubMed: 19675534]
- Shaner A. Delusions, superstitious conditioning and chaotic dopamine neurodynamics. Med Hypotheses. 1999; 52:119–123. [PubMed: 10340292]
- Shanks DR. Learning: from association to cognition. Annu Rev Psychol. 2010; 61:273–301. [PubMed: 19575617]
- Sharp FR, Tomitaka M, Bernaudin M, Tomitaka S. Psychosis: pathological activation of limbic thalamocortical circuits by psychomimetics and schizophrenia? Trends Neurosci. 2001; 24:330– 334. [PubMed: 11356504]
- Shepard PD, Holcomb HH, Gold JM. Schizophrenia in translation: the presence of absence: habenular regulation of dopamine neurons and the encoding of negative outcomes. Schizophr Bull. 2006; 32:417–421. [PubMed: 16717257]
- Sherman SM, Guillery RW. On the actions that one nerve cell can have on another: distinguishing "drivers" from "modulators". Proc Natl Acad Sci U S A. 1998; 95:7121–7126. [PubMed: 9618549]
- Siegel RK. Cocaine hallucinations. Am J Psychiatry. 1978; 135:309–314. [PubMed: 626219]
- Sillitoe RV, Stephen D, Lao Z, Joyner AL. Engrailed homeobox genes determine the organization of Purkinje cell sagittal stripe gene expression in the adult cerebellum. J Neurosci. 2008; 28:12150– 12162. [PubMed: 19020009]
- Silverstein S, Uhlhaas PJ, Essex B, Halpin S, Schall U, Carr V. Perceptual organization in first episode schizophrenia and ultra-high-risk states. Schizophr Res. 2006; 83:41–52. [PubMed: 16497484]
- Simons JS, Henson RN, Gilbert SJ, Fletcher PC. Separable forms of reality monitoring supported by anterior prefrontal cortex. J Cogn Neurosci. 2008; 20:447–457. [PubMed: 18004946]
- Simpson J, Done DJ. Elasticity and confabulation in schizophrenic delusions. Psychol Med. 2002; 32:451–458. [PubMed: 11989990]
- Sirota A, Montgomery S, Fujisawa S, Isomura Y, Zugaro M, Buzsaki G. Entrainment of neocortical neurons and gamma oscillations by the hippocampal theta rhythm. Neuron. 2008; 60:683–697. [PubMed: 19038224]
- Skinner BF. Superstition" in the pigeon. Journal of Experimental Psychology. 1948; 38:168–172. [PubMed: 18913665]

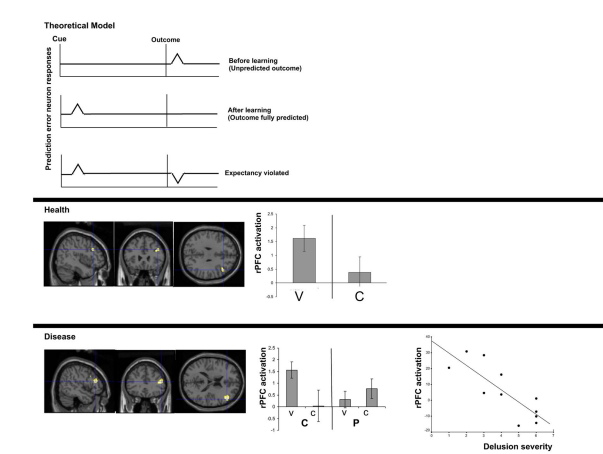
Sokolov, EN. Neuronal models and the orienting reflex. Josiah Macy Jr Foundation; New York: 1960.

Soltani A, Wang XJ. Synaptic computation underlying probabilistic inference. Nat Neurosci. 2010; 13:112–119. [PubMed: 20010823]

- Spence SA, Brooks DJ, Hirsch SR, Liddle PF, Meehan J, Grasby PM. A PET study of voluntary movement in schizophrenic patients experiencing passivity phenomena (delusions of alien control). Brain. 1997; 120 (Pt 11):1997–2011. [PubMed: 9397017]
- Sperry, RW. Forebrain Commisurectomy and consciuos awareness. In: Trevarthen, C., editor. Brain Circuits and the Mind. Cambridge University Press; New York: 1990.
- Spitzer M. A neurocomputational approach to delusions. Compr Psychiatry. 1995; 36:83–105. [PubMed: 7758303]
- Spitzer, M.; Walter, H. The cognitive neuroscience of agency in schizophrenia. In: David, A.; Kircher, T., editors. The self in neuroscience and psychiatry. Cambridge University Press; Cambridge: 2003. p. 436-444.
- Startup M, Startup S. On two kinds of delusion of reference. Psychiatry Res. 2005; 137:87–92. [PubMed: 16226316]
- Stephan KE, Baldeweg T, Friston KJ. Synaptic plasticity and dysconnection in schizophrenia. Biol Psychiatry. 2006; 59:929–939. [PubMed: 16427028]
- Steriade M, Dossi RC, Pare D, Oakson G. Fast oscillations (20–40 Hz) in thalamocortical systems and their potentiation by mesopontine cholinergic nuclei in the cat. Proc Natl Acad Sci U S A. 1991; 88:4396–4400. [PubMed: 2034679]
- Stickgold R, Walker MP. Sleep-dependent memory consolidation and reconsolidation. Sleep Med. 2007; 8:331–343. [PubMed: 17470412]
- Sur M, Rubenstein JL. Patterning and plasticity of the cerebral cortex. Science. 2005; 310:805–810. [PubMed: 16272112]
- Sutton, RS.; Barto, AG. Reinforcement Learning: An Introduction. MIT Press; 1998.
- Svenningsson P, Tzavara ET, Carruthers R, Rachleff I, Wattler S, Nehls M, McKinzie DL, Fienberg AA, Nomikos GG, Greengard P. Diverse psychotomimetics act through a common signaling pathway. Science. 2003; 302:1412–1415. [PubMed: 14631045]
- Tabares-Seisdedos R, Rubenstein JL. Chromosome 8p as a potential hub for developmental neuropsychiatric disorders: implications for schizophrenia, autism and cancer. Mol Psychiatry. 2009; 14:563–589. [PubMed: 19204725]
- Takahashi Y, Schoenbaum G, Niv Y. Silencing the critics: understanding the effects of cocaine sensitization on dorsolateral and ventral striatum in the context of an actor/critic model. Front Neurosci. 2008; 2:86–99. [PubMed: 18982111]
- Takahashi YK, Roesch MR, Stalnaker TA, Haney RZ, Calu DJ, Taylor AR, Burke KA, Schoenbaum G. The orbitofrontal cortex and ventral tegmental area are necessary for learning from unexpected outcomes. Neuron. 2009; 62:269–280. [PubMed: 19409271]
- Tang C, Pawlak AP, Prokopenko V, West MO. Changes in activity of the striatum during formation of a motor habit. Eur J Neurosci. 2007; 25:1212–1227. [PubMed: 17331217]
- Teufel C, Alexis DM, Todd H, Lawrance-Owen AJ, Clayton NS, Davis G. Social cognition modulates the sensory coding of observed gaze direction. Curr Biol. 2009; 19:1274–1277. [PubMed: 19559619]
- Thiebierge G. Les acaraphobes. Annales de Dermatologie et de Syphiligraphie. 1894; 3:730–736. Thorndike, EL. Animal Intelligence. MacMillan; New York: 1911.
- Tole S, Remedios R, Saha B, Stoykova A. Selective requirement of Pax6, but not Emx2, in the specification and development of several nuclei of the amygdaloid complex. J Neurosci. 2005; 25:2753–2760. [PubMed: 15758185]
- Tolman, EC. Purposive behaviour in animals and men. Century; New York: 1932.
- Tricomi E, Balleine BW, O'Doherty JP. A specific role for posterior dorsolateral striatum in human habit learning. Eur J Neurosci. 2009; 29:2225–2232. [PubMed: 19490086]
- Tsakiris M, Haggard P. The rubber hand illusion revisited: visuotactile integration and self-attribution. J Exp Psychol Hum Percept Perform. 2005; 31:80–91. [PubMed: 15709864]
- Turner DC, Aitken MR, Shanks DR, Sahakian BJ, Robbins TW, Schwarzbauer C, Fletcher PC. The role of the lateral frontal cortex in causal associative learning: exploring preventative and superlearning. Cereb Cortex. 2004; 14:872–880. [PubMed: 15054060]

- Uhlhaas PJ, Haenschel C, Nikolic D, Singer W. The role of oscillations and synchrony in cortical networks and their putative relevance for the pathophysiology of schizophrenia. Schizophr Bull. 2008; 34:927–943. [PubMed: 18562344]
- Uhlhaas PJ, Linden DE, Singer W, Haenschel C, Lindner M, Maurer K, Rodriguez E. Dysfunctional long-range coordination of neural activity during Gestalt perception in schizophrenia. J Neurosci. 2006a; 26:8168–8175. [PubMed: 16885230]
- Uhlhaas PJ, Mishara AL. Perceptual anomalies in schizophrenia: integrating phenomenology and cognitive neuroscience. Schizophr Bull. 2007; 33:142–156. [PubMed: 17118973]
- Uhlhaas PJ, Phillips WA, Mitchell G, Silverstein SM. Perceptual grouping in disorganized schizophrenia. Psychiatry Res. 2006b; 145:105–117. [PubMed: 17081620]
- Uhlhaas PJ, Singer W. Abnormal neural oscillations and synchrony in schizophrenia. Nat Rev Neurosci. 11:100–113. [PubMed: 20087360]
- Vallar G, Ronchi R. Somatoparaphrenia: a body delusion. A review of the neuropsychological literature. Exp Brain Res. 2009; 192:533–551. [PubMed: 18813916]
- van Nimwegen L, de Haan L, van Beveren N, van den Brink W, Linszen D. Adolescence, schizophrenia and drug abuse: a window of vulnerability. Acta Psychiatr Scand Suppl. 2005:35– 42. [PubMed: 15877720]
- Vernon D, Haenschel C, Dwivedi P, Gruzelier J. Slow habituation of induced gamma and beta oscillations in association with unreality experiences in schizotypy. Int J Psychophysiol. 2005; 56:15–24. [PubMed: 15725486]
- Vickery TJ, Jiang YV. Associative grouping: perceptual grouping of shapes by association. Atten Percept Psychophys. 2009; 71:896–909. [PubMed: 19429967]
- Vinogradov S, King RJ, Huberman BA. An associationist model of the paranoid process: application of phase transitions in spreading activation networks. Psychiatry. 1992; 55:79–94. [PubMed: 1557472]
- Von Holst E. Relations between the central nervous system and the peripheral organs. British Journal of Animal Behaviour. 1954; 2:89–94.
- Waelti P, Dickinson A, Schultz W. Dopamine responses comply with basic assumptions of formal learning theory. Nature. 2001; 412:43–48. [PubMed: 11452299]
- Waldmann, MR.; Martignon, L. A Bayesian network model of causal learning. In: Gernsbacher, MA.; Derry, SJ., editors. Proceedings of the Twentieth Annual Conference of the Cognitive Science Society. Mahwah, NJ: Earlbaum; 1998. p. 1102-1107.
- Wallis GG. A case of hallucinosis due to cocaine. J R Nav Med Serv. 1949; 35:112. [PubMed: 18127069]
- Walsh T, McClellan JM, McCarthy SE, Addington AM, Pierce SB, Cooper GM, Nord AS, Kusenda M, Malhotra D, Bhandari A, Stray SM, Rippey CF, Roccanova P, Makarov V, Lakshmi B, Findling RL, Sikich L, Stromberg T, Merriman B, Gogtay N, Butler P, Eckstrand K, Noory L, Gochman P, Long R, Chen Z, Davis S, Baker C, Eichler EE, Meltzer PS, Nelson SF, Singleton AB, Lee MK, Rapoport JL, King MC, Sebat J. Rare structural variants disrupt multiple genes in neurodevelopmental pathways in schizophrenia. Science. 2008; 320:539–543. [PubMed: 18369103]
- Waltz JA, Frank MJ, Robinson BM, Gold JM. Selective reinforcement learning deficits in schizophrenia support predictions from computational models of striatal-cortical dysfunction. Biol Psychiatry. 2007; 62:756–764. [PubMed: 17300757]
- Wang J, O'Donnell P. D(1) dopamine receptors potentiate nmda-mediated excitability increase in layer V prefrontal cortical pyramidal neurons. Cereb Cortex. 2001; 11:452–462. [PubMed: 11313297]
- Wang XJ. Neurophysiological and Computational Principles of Cortical Rhythms in Cognition. Physiological Reviews. (In Press).
- Wegner DM. Precis of the illusion of conscious will. Behav Brain Sci. 2004; 27:649–659. discussion 659–692. [PubMed: 15895616]
- Weiller C, Juptner M, Fellows S, Rijntjes M, Leonhardt G, Kiebel S, Muller S, Diener HC, Thilmann AF. Brain representation of active and passive movements. Neuroimage. 1996; 4:105–110. [PubMed: 9345502]

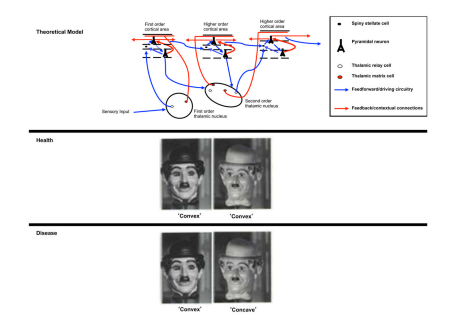
- Weiskrantz L, Elliott J, Darlington C. Preliminary observations on tickling oneself. Nature. 1971; 230:598–599. [PubMed: 4928671]
- Whalen PJ, Rauch SL, Etcoff NL, McInerney SC, Lee MB, Jenike MA. Masked presentations of emotional facial expressions modulate amygdala activity without explicit knowledge. J Neurosci. 1998; 18:411–418. [PubMed: 9412517]
- Whalen PJ, Shin LM, McInerney SC, Fischer H, Wright CI, Rauch SL. A functional MRI study of human amygdala responses to facial expressions of fear versus anger. Emotion. 2001; 1:70–83. [PubMed: 12894812]
- Whitson JA, Galinsky AD. Lacking control increases illusory pattern perception. Science. 2008; 322:115–117. [PubMed: 18832647]
- Winterer G. Cortical microcircuits in schizophrenia--the dopamine hypothesis revisited. Pharmacopsychiatry. 2006; 39(Suppl 1):S68–71. [PubMed: 16508900]
- Wolff G, McKenzie K. Capgras, Fregoli and Cotard's syndromes and Koro in folie a deux. Br J Psychiatry. 1994; 165:842. [PubMed: 7881797]
- Wolpert DM, Ghahramani Z, Jordan MI. An internal model for sensorimotor integration. Science. 1995; 269:1880–1882. [PubMed: 7569931]
- Wolpert DM, Miall RC. Forward Models for Physiological Motor Control. Neural Netw. 1996; 9:1265–1279. [PubMed: 12662535]
- Woodward TS, Moritz S, Chen EY. The contribution of a cognitive bias against disconfirmatory evidence (BADE) to delusions: a study in an Asian sample with first episode schizophrenia spectrum disorders. Schizophr Res. 2006; 83:297–298. [PubMed: 16513331]
- Yin HH, Knowlton BJ, Balleine BW. Lesions of dorsolateral striatum preserve outcome expectancy but disrupt habit formation in instrumental learning. Eur J Neurosci. 2004; 19:181–189. [PubMed: 14750976]
- Yoo SS, Freeman DK, McCarthy JJ 3rd, Jolesz FA. Neural substrates of tactile imagery: a functional MRI study. Neuroreport. 2003; 14:581–585. [PubMed: 12657890]
- Young AW, Robertson IH, Hellawell DJ, de Pauw KW, Pentland B. Cotard delusion after brain injury. Psychol Med. 1992; 22:799–804. [PubMed: 1410102]
- Young G. Capgras delusion: an interactionist model. Conscious Cogn. 2008; 17:863–876. [PubMed: 18314350]
- Yu AJ, Dayan P. Uncertainty, neuromodulation, and attention. Neuron. 2005; 46:681–692. [PubMed: 15944135]



## Figure 1. Neural Instantiation of predictive learning and belief

*Theoretical model*: Schematic of reward prediction error signals before learning, following learning and during extinction

*Health*: Right DLPFC prediction error response during casual learning in healthy subjects (Corlett et al, 2004) – V: Violation of expectancy, C: Confirmation of expectancy *Disease*: Aberrant right frontal prediction error response in patients with first episode psychosis. The more profound the disruption, the more severe the delusions (Corlett et al, 2007b) - C: Controls, P: Patients with Psychosis

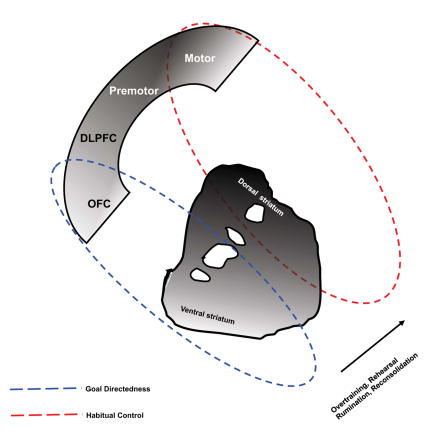


## Figure 2. Learning Memory and Belief Alter Perception

Theoretical model: Feedforward and feedback thalamocortical projections (adapted from http://wiki.tkk.fi/display/SYNB/Neocortex).

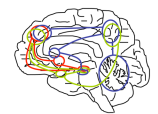
Health: The rotating hollow mask is continuously perceived as convex due to our consistent experience of faces as convex.

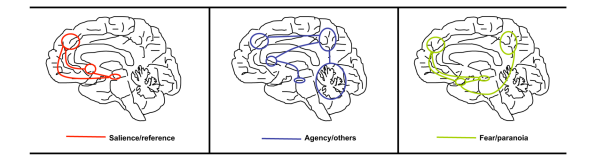
Disease: Individuals prone to or experiencing psychosis report the hollow mask as a hollow percept (Emrich et al, 1988).



## Figure 3. Neural Circuitry of Goal Directedness (knowledge) and habit (belief)

With repetition, rumination and reconsolidation, the control of behavior shifts from flexible goal-directed ventral cortcostriatal control toward control by the inflexible dorsal striatum and motor cortex.





## **Figure 4. Putative Delusion Circuits**

*Salience/Reference*: A circuit incorporating the midbrain dopaminergic nuclei, the associative striatum and frontal cortex. Aberrant prediction errors in midbrain update expectancies in the frontal cortex leading to aberrantly salient percepts.

*Agency/others*: The midbrain, PFC, Parietal cortex and cerebellum as well as the bimodal cells of the putamen. This circuit describes forward model predictions used to discern whether sensory stimulation was internally of externally generated. A breakdown in this predictive mechanism would manifest as hallucinatory tactile percepts and inferences of external control of intentional action.

*Fear/Paranoia*: A circuit incorporating the midbrain, amygdala, frontal and parietal cortices. Here, neutral or irelevent stimuli, thoughts and percepts come to engender fear and anxiety. A dysfunction in frontoparietal circuitry engenders inappropriate social predictions and maladaptive inferences about the intentions of others.

*Interaction between circuits:* These circuits interact and likely mutually reinforce one another.