A Neuro-Computational Model of Economic Decisions

Aldo Rustichini¹ and Camillo Padoa-Schioppa²

¹ Department of Economics, University of Minnesota, 1925 4th Street South 4-101, Minneapolis, MN 55455. ² Departments of Anatomy and Neurobiology, Economics, and Biomedical Engineering, Washington University in St Louis, 660 South Euclid Avenue, St Louis, MO 63110

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Correspondence to: A.R. (rusti001@umn.edu) or C.P.-S. (camillo@wustl.edu)

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Abstract

Neuronal recordings and lesion studies indicate that key aspects of economic decisions take place in the orbitofrontal cortex (OFC). Previous work identified in this area three groups of neurons encoding the offer value, the chosen value and the identity of the chosen good. An important and open question is whether and how decisions could emerge from a neural circuit formed by these three populations. Here we adapted a biophysically realistic neural network previously proposed for perceptual decisions (Wang 2002; Wong and Wang 2006). The domain of economic decisions is significantly broader than that for which the model was originally designed: yet the model performed remarkably well. The input and output nodes of the network were naturally mapped onto two groups of cells in OFC. Surprisingly, the activity of interneurons in the network closely resembled that of the third group of cells, namely chosen value cells. The model reproduced several phenomena related to the neuronal origins of choice variability. It also generated testable predictions on the excitatory/inhibitory nature of different neuronal populations and on their connectivity. Some aspects of the empirical data were not reproduced, but simple extensions of the model could overcome these limitations. These results render a biologically credible model for the neuronal mechanisms of economic decisions. They demonstrate that choices could emerge from the activity of cells in the OFC, suggesting that chosen value cells directly participate in the decision process. Importantly, Wang's model provides a platform to investigate the implications of neuroscience results for economic theory.

Introduction

Economic choices are thought to entail two mental stages: subjective values are first assigned to the available options, and decisions are made by comparing values. Evidence from lesion studies (Fellows 2011; Rudebeck and Murray 2014), functional imaging (Bartra et al. 2013; Clithero and Rangel 2013) and neurophysiology (Mainen and Kepecs 2009; Padoa-Schioppa 2011; Wallis 2011) indicates that choices, in particular choices between goods, involve the OFC. Neuronal recordings in primates choosing between different juices identified three groups of neurons in this area: offer value cells encoding the value of individual juices, chosen juice cells encoding the
binary decision outcome, and *chosen value* cells encoding the value of the chosen juice (Padoa-Schioppa 2013; Padoa-Schioppa and Assad 2006). Prima facie, these groups of neurons appear sufficient to characterize – and possibly generate – a decision. Indeed, *offer value* cells capture the input to the decision process, while *chosen juice* and *chosen value* cells capture the identity and value of the chosen good, and thus the decision outcome. A primary goal for decision neuroscience is to formalize this intuition by building a biologically realistic model in which the groups of cells found in OFC form a circuit that generates decisions. Ideally, such a model would encompass all that is known about these neurons and, concurrently, make new and testable predictions.

Our thinking about the mechanisms of economic decisions is influenced by work on motion perception (perceptual decisions). In a simplified scheme, two brain regions play a primary role: the middle temporal (MT) area, where neurons encode the momentary evidence, and the lateral intraparietal (LIP) area, where cells represent the decision outcome in the form of a planned saccade (Gold and Shadlen 2007; Parker and Newsome 1998). Notably, there is a natural analogy between neurons in MT and *offer value* cells, and between neurons in LIP and *chosen juice* cells. In contrast, *chosen value* cells do not have a known correspondent in perceptual decisions.

Several models have been proposed to describe the neuronal mechanisms of perceptual decisions (Bogacz et al. 2006; Drugowitsch and Pouget 2012; Gold and Shadlen 2007). At the biophysical level, a leading proposal is Wang's model, in which decisions emerge from a balance of recurrent excitation and pooled inhibition. Different variants of the model account for perceptual decisions (Wang 2002; Wong and Wang 2006), similarity judgments (Engel and Wang 2011), probabilistic inference (Soltani and Wang 2010), behavior in a competitive game (Soltani et al. 2006) and flexible sensori-motor mapping (Fusi et al. 2007). More recently, the model has been adapted to describe the activity in LIP during foraging tasks (Soltani and Wang 2006) and to fit aggregate neural activity during value-based decisions (Hunt et al. 2012; Jocham et al. 2012). However, a precise mapping between Wang's model and the activity of neurons in OFC (or any brain area) during economic decisions has not yet been attempted. In this study, we examined the extent to which Wang's model can reproduce neuronal activity in the OFC. From a modeling perspective, this is a challenging test because – as we argue in detail below – the domain of economic
decisions is significantly broader than that for which the model was originally designed, and
because neuronal activity in the OFC during economic decisions only partly resembles that in
area LIP during perceptual decisions. Our deliberate goal was to test the model without changing
its structure. Importantly, all the parameters in the model represent biophysical quantities such as
synaptic efficacies and time constants and are derived, at least approximately, from empirical
measures (Wang 2002). In our investigation, we used the same parameters previously used to
model perceptual decisions (Wong and Wang 2006).

In most respects, Wang's model reproduced the activity of different groups of cells in OFC
remarkably well. The two layers corresponding to the input and to the categorical output of the
network were identified with offer value cells and chosen juice cells, respectively. Most
surprisingly, the activity of inhibitory interneurons in the network closely resembled that of the
third group of neurons found in OFC, namely chosen value cells. The model also reproduced
several phenomena related to the neuronal origins of choice variability, namely choice hysteresis,
the "predictive activity" of chosen juice cells and the "activity overshooting" of chosen value
cells (Padoa-Schioppa 2013). Two aspects of the empirical data were not reproduced. First, the
model did not include neurons with negative encoding. Second, a significant baseline in the
activity of offer value cells introduced distortions in the behavior of the network. However,
simple extensions of the model could overcome these limitations. The results of this study render
a biologically credible model for the neuronal mechanisms of economic decisions.

Materials and Methods

Structure of the model

In its extended form (Brunel and Wang 2001; Wang 2002), the model is a recurrent network of
2,000 spiking neurons, of which 80% \( N_E \) are excitatory pyramidal cells and 20% \( N_I \) are
inhibitory interneurons. All neurons in the network are leaky integrate-and-fire cells endowed
with biophysically realistic parameters. Two external stimuli provide the primary input, and each
stimulus activates a fraction \( f = 0.15 \) of pyramidal cells. The remaining \((1-2f)\) \( N_E \) pyramidal cells
are non-responsive (untuned). The synaptic input to each neuron is both excitatory and inhibitory.
Excitatory inputs are through AMPA- and NMDA-mediated synapses, while inhibitory inputs are through GABA_A-mediated synapses. For each neuron, the input has an external component and a recurrent component. For the two groups of selective pyramidal cells, the external component includes the external stimulus. In addition, each neuron in the network receives an external background noise distributed as a Poisson process. Stimulus current and external background are through AMPA-mediated synapses. The recurrent component is provided by other neurons in the network. In analogy to the Hebbian rule, synapses between neurons in a given group (which fire together) are potentiated by a factor $w_+ > 1$, whereas synapses between neurons in different groups are depressed by a factor $w_- < 1$. The condition $w_- = 1 - f(w_+ - 1)/(1 - f)$ ensures that all excitatory cells have the same spontaneous firing rate.

Using a mean-field approach (Renart et al. 2003; Wong and Wang 2006), the network can be reduced to a dynamic system of 11 variables (see below). In the following, we refer to this version of the model as W11. Under several additional assumptions and approximations, the model can be further reduced to a dynamic system of 2 variables (henceforth, W2). The advantage of the 2-variable formulation is that it is easily tractable – for example, it is possible to examine the dynamics of the system in a phase plane. However, two important reasons motivated us to examine W11. First, W11 is expressed in terms of neuronal firing rates, which allows a direct comparison between the activity of units in the model and that of cells recorded experimentally. In contrast, W2 is expressed in terms of gating variables, which represent the fraction of open channels and are not directly accessible in our experiments. Second, W11 expresses explicitly the activity of each group of cells including inhibitory interneurons. In contrast, interneurons in W2 exist only implicitly and their dynamics is not easily recovered. In summary, using W11 allowed us to contrast directly the predictions of Wang's model with the activity of neurons in the OFC.

Many previous studies examined the extended, spiking version of the model or W2. In contrast, W11 was discussed only briefly by Wong and Wang (2006). We thus recapitulate it here for convenience. A schematic representation of the model is provided in Fig.2. In preliminary tests, we observed that the activity of inhibitory interneurons in the model examined as a function of the offer type resemble that of chosen value cells. Throughout the paper, we refer to neurons in OFC as offer value, chosen juice and chosen value cells, and to nodes in the model as OV, CJ
and CV cells, respectively. The three groups of pyramidal cells correspond to CJA cells (group 1), CJB cells (group 2) and non-selective cells (group 3). Unless otherwise indicated, all the parameters used in our simulations were set identical to those used in the original W11 (Wong and Wang 2006). Their values are indicated in Table 1.

The dynamic system is defined by the following 11 equations:

\[ \tau_{\text{AMPA}} \frac{dr_i}{dt} = -r_i + \phi(I_{\text{syn},i}) \quad i = 1, 2, 3 \] (1)

\[ \tau_{\text{GABA}} \frac{dr_i}{dt} = -r_i + \phi(I_{\text{syn},i}) \] (2)

\[ \frac{dS_{\text{AMPA},i}}{dt} = -\frac{S_{\text{AMPA},i}}{\tau_{\text{AMPA}}} + r_i \quad i = 1, 2, 3 \] (3)

\[ \frac{dS_{\text{NMDA},i}}{dt} = -\frac{S_{\text{NMDA},i}}{\tau_{\text{NMDA}}} + (1 - S_{\text{NMDA},i})\gamma r_i \quad i = 1, 2, 3 \] (4)

\[ \frac{dS_{\text{GABA}}}{dt} = -\frac{S_{\text{GABA}}}{\tau_{\text{GABA}}} + r_i \] (5)

In Eq. 1 and below, \( i = 1, 2, 3 \) refers to the three groups of pyramidal cells and \( r_i \) indicates the firing rate. In Eq. 2 and below, \( I \) indicates interneurons. For each group of cells \( i \) and for each receptor type \( R, S_{R,i} \) is the corresponding gating variable, defined as the fraction of open ionic channels. The parameters \( \tau \) are time constants and \( \gamma \) in Eq. 4 is a constant (see Table 1). Note that firing rates and gating variables are all time dependent. The input-output relation for a leaky integrate-and-fire cell is given by the simplified formula of Abbott and Chance (2005):

\[ \phi(I_{\text{syn}}) = \frac{c_{E,i}I_{\text{syn}} - I_{E,i}}{1 - \exp\left(-g_{E,i}\left(c_{E,i}I_{\text{syn}} - I_{E,i}\right)\right)} \] (6)

Note that Eq. 6 is written separately for excitatory pyramidal cells (\( E \)) and inhibitory interneurons (\( I \)). In this equation, \( I_{\text{syn}} \) is the total synaptic input to the cell, and the parameters \( I_{E,i}, c_{E,i} \) and \( g_{E,i} \) are, respectively, the threshold current, the gain factor and the noise factor (see Table 1).
For each group of cells, the input current $I_{syn}$ includes several components:

$$I_{syn} = I_{AMPA,ext} + I_{AMPA,rec} + I_{NMDA,rec} + I_{GABA,rec} + I_{stim}$$ \hfill (7)

In Eq.7 and below, indices $ext$ and $rec$ refer to external and recurrent currents. Currents depend on the gating variables through the following equations:

$$I_{AMPA,ext,i} = -J_{AMPA,ext,pyr} \tau_{AMPA} C_{ext} r_{ext} + I_{\eta,i} \quad i = 1, 2, 3$$ \hfill (8)

$$I_{AMPA,rec,i} = -N_E f J_{AMPA,pyr} \left(w_+ S_{AMPA,i} + w_- S_{AMPA,j\neq i}\right) - N_E (1 - 2f) J_{AMPA,pyr} w_- S_{AMPA,3} \quad i, j = 1, 2$$ \hfill (9)

$$I_{AMPA,rec,3} = -N_E f J_{AMPA,pyr} \left(S_{AMPA,1} + S_{AMPA,2}\right) - N_E (1 - 2f) J_{AMPA,pyr} S_{AMPA,3}$$ \hfill (10)

$$I_{NMDA,rec,i} = -N_E f J_{NMDA,pyr} \delta J_{NMDA,i} \left(w_+ S_{NMDA,i} + w_- S_{NMDA,j\neq i}\right) - N_E (1 - 2f) J_{NMDA,pyr} w_- S_{NMDA,3} \quad i, j = 1, 2$$ \hfill (11)

$$I_{NMDA,rec,3} = -N_E f J_{NMDA,pyr} \left(S_{NMDA,1} + S_{NMDA,2}\right) - N_E (1 - 2f) J_{NMDA,pyr} S_{NMDA,3}$$ \hfill (12)

$$I_{GABA,rec,i} = -N_E f J_{GABA,pyr} \delta J_{GABA,i} S_{GABA} \quad i = 1, 2, 3$$ \hfill (13)

The corresponding equations written for interneurons are:

$$I_{AMPA,ext,1} = -J_{AMPA,ext,in} \tau_{AMPA} C_{ext} r_{ext} + I_{\eta,1}$$ \hfill (14)

$$I_{AMPA,rec,1} = -N_E f J_{AMPA,in} \left(S_{AMPA,1} + S_{AMPA,2}\right) - N_E (1 - 2f) J_{AMPA,in} S_{AMPA,3}$$ \hfill (15)

$$I_{NMDA,rec,1} = -N_E f J_{NMDA,rec,in} \left(S_{NMDA,1} + S_{NMDA,2}\right) - N_E (1 - 2f) J_{NMDA,rec,in} S_{NMDA,3}$$ \hfill (16)

$$I_{GABA,rec,1} = -N_E f J_{GABA,in} S_{GABA}$$ \hfill (17)

In these equations, $C_{ext}$ is the total number of external synapses per cell and the parameters $J$ are synaptic efficacies, whose values are derived from empirical measures (Wang 2002; Wong and Wang 2006). The two parameters $\delta J_{NMDA,i}$ and $\delta J_{GABA,i}$, which were not present in the original W11, were introduced here in certain simulations to examine the effects of synaptic imbalance (see section Imposing non-trivial relative values). In the initial simulations, $\delta J_{NMDA,i}$ and $\delta J_{GABA,i}$ were set equal to 1.
Finally, $I_{\eta,i}$ in Eq.8 and $I_{\eta,i}$ in Eq.14 are noise terms. Neurons in the extended, spiking network (Wang 2002) are noisy Poisson processes. In the derivation of W11, Wong and Wang (2006) first removed the time-dependent noise during the mean-field approximation and then reintroduced a white noise in the form of a Ornstein-Uhlenbeck process. Thus $I_{\eta}$ is according to the following equation:

$$\tau_{AMPA} \frac{dI_{\eta,i}}{dt} = -I_{\eta,i} + \eta(t)\sqrt{\tau_{AMPA} \sigma_{\eta}^2} \quad j = 1, 2, 3, I \quad (18)$$

where $\eta(t)$ is a white noise with unit variance (Renart et al. 2003). The parameter $\sigma_{\eta}$ represents the amount of noise (see Table 1).

**Modeling offer value cells**

The last term in Eq.7, namely $I_{stim}$, is the primary input, which equals zero for non-selective pyramidal cells and for interneurons. For CJA and CJB cells, we set $I_{stim}$ as follows:

$$I_{stim,i} = -J_{AMPA, input} \delta J_{HL,i} \delta J_{stim,i} \tau_{AMPA} r_{OV,i} \quad i = 1, 2 \quad (19)$$

where $r_{OV}$ is the firing rate of OV cells. The synaptic efficacy $J_{AMPA, input}$ reflects the number of connections between OV cells and CJ cells. In the original W11, $J_{AMPA, input}$ was set equal to $J_{AMPA, ext, pyr}$. Unless otherwise indicated, in all our simulations we set $J_{AMPA, input} = 30 J_{AMPA, ext, pyr}$. This adjustment provided a sizable dynamic range for CV cells and reflected the fact that offer value cells and chosen juice cells are found in close proximity in the same brain region, while cells in MT and LIP are connected only long-distance.

In Eq.19, the parameter $\delta J_{stim,i}$ was introduced to impose non-trivial relative values (see section *Imposing non-trivial relative values*). In the initial simulations, we set $\delta J_{stim} = (2, 1)$. Conversely, the parameter $\delta J_{HL,i}$ accounted for the Hebbian learning taking place after range adaptation (see section *Range adaptation, context-dependent preferences and Hebbian learning*). We generally set $\delta J_{HL} = (\Delta A/\Delta B, 1)$. Unless otherwise indicated, the two value ranges used in the simulations were equal and $\delta J_{HL} = (1, 1)$. 


For the sake of simplicity, the value encoded by offer value cells is treated here as equivalent to the corresponding juice quantity (In reality, these two variables are distinct (Raghuraman and Padoa-Schioppa 2014).) For any session and for any juice \( X \), \(#X\) indicates the quantity of \( X \) offered in the current trial, \(#X_{\text{min}}\) and \(#X_{\text{max}}\) indicate the minimum and maximum quantities of \( X \) offered in the session, and \( \Delta X = [\#X_{\text{min}}, \#X_{\text{max}}]\) is the value range. The encoding of values in OFC undergoes range adaptation (Kobayashi et al. 2010; Padoa-Schioppa 2009). In other words, the activity of offer value cells on any given trial is a linear function of the value rank \( x \):

\[
x = \left( \frac{\#X - \#X_{\text{min}}}{\#X_{\text{max}} - \#X_{\text{min}}} \right)
\]

(20)

The activity profile of OV cells was modeled as follows:

\[
r_{\text{OV},i}(n,t) = r_0 + \Delta r \cdot f(t) \cdot x_i
\]

\[i = 1, 2\]

(21)

\[
f(t) = g(t) / \max(g(t))
\]

(22)

\[
g(t) = \frac{1}{1 + \exp\left(-\frac{t - a}{b}\right)} \cdot \frac{1}{1 + \exp\left(-\frac{t - c}{d}\right)}
\]

(23)

where \( n \) is the trial number, \( t \) is time within a trial, and \( x_i \) is the rank for juice \( X_i \). Juices A and B correspond to \( X_1 \) and \( X_2 \), respectively.

The baseline activity \( (r_0) \), dynamic range \( (\Delta r) \) and time constants \( (a, b, c, d) \) used in the simulations are indicated in Table 1. Unless otherwise indicated, in all the simulations we set the same value range for the two juices, namely \( \Delta A = \Delta B = [0, 20] \). This large range allowed to generate choice patterns and neuronal activity at high resolution.

Simulations

Simulations were ran in Matlab (MathWorks Inc). Unless otherwise specified, each session included 4,000 trials with both offers randomly selected on each trial from the range \([0, 20]\). Offers 0A:0B were excluded. The network dynamics was generated with a resolution \( dt = 0.5 \) ms and then examined averaging over 5 ms time bins. For each trial, the choice outcome was identified by comparing the activity of CJA and CJB cells in the interval 400-600 ms after the offer (similar results were obtained using different time windows). A trial type was defined by
two offers and a choice (e.g., [2A:5B, B]). For the study of activity profiles (e.g., Fig.4a), trials were divided in groups according to the relevant criterion and the activity was averaged across trials for each group. For the study of tuning functions (e.g., Fig.4bcd), we focused on specific time windows, namely 0-500 ms after the offer for OV cells and CV cells, and 500-1000 ms after the offer for CJ cells. However, qualitatively similar results were obtained using different time windows. Firing rates were averaged across trials for each trial type. Choice patterns (e.g., Fig.5a) were analyzed using logistic regressions (see Results). The code is available upon request.

Results

Summary of experimental observations

Fig.1 summarizes the primary experimental results that we sought to reproduce with Wang's model. In the experiments (Padoa-Schioppa and Assad 2006; 2008), monkeys chose between two juices labeled A and B, with A preferred. On any given trial, the offers appeared on a computer monitor on the two sides of a center fixation point. After the offer, the animals maintained center fixation for a randomly variable delay that lasted 1-2 s, after which the center fixation was dimmed (go signal). The animal indicated their choice with a saccade. Juice quantities varied from trial to trial and choice patterns typically presented a quality-quantity trade-off. For example in the session shown in Fig.1A, the monkey was roughly indifferent between 1A and 3B. Recordings were performed in central OFC. In the initial analysis, we defined several time windows aligned with different behavioral events and a large number of variables that neurons could conceivably encode. A variable selection analysis indicated that (1) neurons in OFC encoded one of three variables, namely offer value, chosen juice and chosen value. Further analyses showed that (2) the encoding of these variables was roughly linear and that (3) each cell encoded one variable or another variable, but not a mixture of variables. We also found that (4) any given cell generally encoded the same variable in different time windows. Thus neurons encoding the three variables were conceptualized as forming different groups of cells (Padoa-Schioppa 2013). For each variable (5) the encoding could be either positive (higher activity for higher values) or negative (lower activity for higher value). Importantly, experiments in which
animals chose between three juices offered pairwise showed that (6) cells encoding the *offer value* of one particular juice did not depend on the juice offered in alternative (menu invariance).

Last but not least, (7) the activity of both *offer value* and *chosen value* cells adapted to the range of values available in any given session (Padoa-Schioppa 2009).

Three examples of neurons encoding the *offer value* $B$, the *chosen juice* $A$ and the *chosen value* are illustrated in Fig.1. The three right panels depict the population activity profile recorded for each group of cells (only the populations of cells positive with encoding are shown). Fig.1d and behavioral measures (Padoa-Schioppa, unpublished observations) indicate that decisions were made within a few hundred milliseconds after the offer.

**Wang's model for economic decisions**

Fig.2 illustrates the structure of the model. We identified the input node (OV cells, corresponding to MT in perceptual decisions) with *offer value* cells. In the original W11, the input current has the form

$$I_{stim, i} = - J_{AMPAs, ext} \tau_{AMPAs} \mu_0 (1 \pm coh) \quad i = 1, 2 \quad (24)$$

where $\mu_0 = 40$ sp/s, $coh \in [-1, 1]$, and the ± signs refer to pools A and B, respectively. The fact that the range of possible inputs is bounded to the interval [0, 80] sp/s is essential for the network to operate properly. As we move from perceptual to economic decisions, we have to consider several factors.

First, unlike motion coherence, values are not bounded to a finite range. In principle, this fact could pose a challenge for the network. In reality, however, this challenge does not arise thanks to the phenomenon of range adaptation (Kobayashi et al. 2010; Padoa-Schioppa 2009). After neurons have adapted to the range of values available in the behavioral context (current session), their activity on any given trial is a linear function of the value rank, which varies in the interval [0, 1] (see Eq.20). We discuss below ways in which range adaptation presents a challenge for Wang's model. However, at this stage range adaptation makes it easy to identify *offer value* cells with the input node of the model.
Second, in the random dot task (Newsome 1997), the two inputs are perfectly anti-correlated. In this sense, the stimulus is intrinsically one-dimensional (and indeed it is parameterized by the uni-dimensional parameter $coh$). Thus if we consider the plane formed by $I_{stim, 1}$ and $I_{stim, 2}$, the inputs for the random dot task lie on the diagonal with slope -1, and the data point corresponding to $coh = 0$ is in the center of the diagram (Fig.3a). In contrast, the two offers in economic choice tasks (Padoa-Schioppa 2011) can vary independently of one another and assume any value within the range spanned in the behavioral session. Thus the input to the model can lie anywhere on the plane formed by $I_{stim, 1}$ and $I_{stim, 2}$ (Fig.3b). As a consequence, there are many sets of offers that induce behavioral indifference. In practice, offers used in the experimental sessions did not cover the plane densely, because in most trials one of the two juices was offered in quantity 1 (Padoa-Schioppa and Assad 2006; 2008). However, in the present study, we simulated offers covering the full plane (Fig.3b). In other words, we tested the neuronal network well beyond its original domain of definition.

Third, in the original W11, the time profile of the input currents is a box car, which mimics the fact that neurons in MT respond with good time fidelity to the momentary motion of the visual stimulus. In contrast, the time profile of offer value cells in OFC is more complex (Fig.1b). In particular, we note several salient aspects. (1) There is a baseline of ~6 sp/s and a dynamic range of ~10 sp/s. Importantly, the ratio between dynamic range and baseline is modest compared to that typically reported for MT and modeled in W11, where the baseline is negligible and the dynamic range is roughly 80 sp/s. (2) Compared to baseline, the modulation due to value is all in the direction of increased firing rates. In other words, focusing on the 500 ms following the offer, the baseline-subtracted mean firing rate of offer value cells ranges roughly between 0 and 10 sp/s, depending on the value offered in the trial. Thus in our simulations, we used a semi-realistic time profile to model the activity of offer value cells. In the initial simulations we set the baseline to zero. The effects of introducing a nonzero baseline will be examined in the section Baseline activity for OV cells.

Fourth, the input from MT to LIP is via long-distance connections. In contrast, offer value cells and chosen juice cells are in the same anatomical region and thus they presumably enjoy the density typical of local connections. We thus increased the connectivity between OV cells and CJ cells (Eq.19). In the simulations, we set $J_{AMPA, input} = 30 J_{AMPA, ext, pyr}$. This value was chosen to
obtain a sizable dynamic range for CV cells. Notably, even considering the fact that the dynamic range of offer value cells is much lower than that of neurons in MT, this value made the input current higher than that used in the original W11.

Fifth, the activation of chosen juice cells in OFC is largely phasic (Fig.1b). Their activity profile increases shortly after the offer, peaks ~350 ms after the offer, and decays rapidly in the following 300-400 ms, even though the offers are still on the monitor and the animal has not yet revealed its choice. The traces corresponding to the two choice outcomes remain separated, but that signal is relatively small until after the animal has performed its saccade. Furthermore, if one compares the traces for easy and split decisions (see below), it appears clear that the activity of chosen juice cells does not resemble a race-to-threshold. This largely-phasic activity profile differs from that reported for LIP in the random dot task, and this difference is relevant to the model. Indeed, the precursor to Wang's model discussed here was a model designed to describe the persistent delay activity observed in lateral prefrontal cortex and other association areas (Brunel and Wang 2001; Compte et al. 2000). Reverberation mediated by NMDA receptors, which have a slow dynamics, is a characteristic trait of this model. Working memory activity generally increases with $w_+$, and the original W11 set $w_+ = 1.80$ (Wong and Wang 2006). However, the authors noted that depending on $w_+$, the model could perform decisions with or without working memory. In our initial simulations we set $w_+ = 1.75$, which provided a largely-phasic (as opposed to a working memory-like) time profile for chosen juice cells. The dependence of the results on $w_+$ is described later in the section Dependence on the strength of recurring synapses.

Initial simulations

Fig.4 shows the results of one simulation. The top row illustrates the activity of OVB cells.

Fig.4a depicts the activity profile, splitting trials depending on whether the encoded value was low, medium or high. Fig4b depicts the full 2-dimensional tuning curve. The x- and y-axes represent the quantities of juice A and juice B offered and the z-axis represents the firing rate. Each data point represents one trial type, and trial types are coded depending on whether the model chose juice A (red diamonds) or juice B (blue circles). To visualize the tuning of OV cells in a 1-dimensional format similar to that of Fig.1a, we down-sampled the tuning curve selecting
a subset of offer types analogous to those typically employed in the experiments (Fig.4c). Finally, Fig.4d illustrates the activity of OVB cells plotted against the encoded variable (offer value B).

The center row illustrates the activity of CJB cells in a similar format. In Fig.4e, trials were divided depending on whether the model chose juice A or juice B. Fig.4f depicts the full tuning curve, with colors indicating the chosen juice. Fig.4g shows a down-sampled version of the tuning curve, and Fig.4h displays the activity of CJB cells as a function of the chosen juice.

The bottom row illustrates the activity of CV cells. To generate these plots, we analyzed the network's choices and derived the relative value of the two juices from the indifference point. The details of this computation is described in the next section. In essence, the relative value reconstructed from choices ($\rho = 2.03$) was very close to the ratio $\delta J_{stim, 1}/\delta J_{stim, 2} = 2$. Following the approach used in the analysis of empirical data (Padoa-Schioppa and Assad 2006), we could thus express quantities of either juice on a single value scale, and we conventionally used units of juice B (uB). For each trial type, we could thus calculate the chosen value. In Fig.4i, trials were divided depending on whether the chosen value was low, medium or high. Fig.4j depicts the full tuning curve, with colors indicating the chosen juice. Fig.4k shows a down-sampled version of the tuning curve, and Fig.4l displays the activity of CV cells as a function of the chosen value.

Several points are noteworthy. First, for both CJ cells and CV cells, the activity profile (Fig.4e, Fig.4i) reproduces fairly well the corresponding activity profile recorded experimentally (Fig.1b, Fig.1c). Second, for CJ cells, the tuning curve clearly separates between the two choice outcomes – the activity of CJB is higher (lower) when the network chooses juice B (juice A) (Fig.4f). At the same time, the tuning is not quite binary. For given choice, the activity of CJB cells increases (decreases) with the quantity of juice B (juice A) offered. This characteristic was already present in the extended version of the model describing perceptual decision (see Fig.4 in Wang (2002)). Third, the tuning curve obtained for CV cells (Fig.4k) closely resembles those obtained experimentally for chosen value cells. Inspection of Fig.4l reveals that, for given chosen value (x-axis), the activity of CV cells is essentially identical when the network chooses juice A (red diamonds) and when it chooses juice B (blue circles). This fact means that CV cells indeed capture the relative value of the two juices ($\rho$).
The resemblance between CV cell and chosen value cells was unexpected and quite remarkable if one considers the fact that interneurons were included in the model for biological realism and stability (Amit and Brunel 1997) – not to reproduce empirical observations analogous to chosen value cells. An intuition for this result can be gained noting that CV cells receive an input proportional to the activity sum of CJA and CJB cells (Eq.15, 17). By virtue of the decision, this sum eventually equals the activity of the winning pool. In turn, CJA and CJB cells receive an input proportional to the value of the corresponding juice weighted by the relative value \( \rho \).

Consequently, the input to – and thus the activity of – CV cells is linearly related to the value of the chosen juice, namely the chosen value. Important in this respect is the fact – rarely noted but clearly apparent in Wang (2002) – that the network attractors depend on the input currents. Fig.41 reveals that the activity of CV cells as a function of the chosen value is slightly sub-linear. To some extent, this is also true in the empirical population data (Padoa-Schioppa, unpublished observations), although the departure from linearity at the level of individual cells rarely reaches significance. As described below, the magnitude of these effects depends on the parameter \( w_+ \).

Wang’s network also includes a population of non-selective (NS) pyramidal cells, which are interconnected with both groups of CJ cells and receive inhibition from CV cells (Fig.2). For the reasons discussed above, this pattern of connectivity suggests that the activity of NS cells should resemble that of chosen value cells in OFC, and this is indeed what we found (not shown). Apart from the excitatory/inhibitory nature of the neurons, one difference between CV cells and NS cells is that the latter group presented a lower dynamic range (~1 sp/s for the simulation of Fig.4) and a higher trial-by-trial variability (relative to the dynamic range). Importantly, while inhibitory interneurons (CV cells) are critical to the decision mechanism, NS cells do not play a significant role in the decision (Amit and Brunel 1997; Wong and Wang 2006). Thus we will not discuss them further.

**Imposing non-trivial relative values**

Consider a subject choosing between quantities of goods A and B. If value functions are linear (i.e., if the subjective value assigned to each good increases as a linear function of the good’s quantity), the relative value between the two goods, namely \( \rho \), is defined as the quantity that makes the subject indifferent between 1A and \( \rho \)B. A hallmark of economic decisions is the fact
that relative values are subjective and even variable over time. In fact, relative values capture the quintessence of economic decisions, namely the fact that subjective value provides a common unit of measure to compare qualitatively different goods. In general, goods A and B have different physical dimensions, and $\rho$ has the physical dimensions necessary to convert one unit of good A into one unit of good B. A fundamental but often under-appreciated issue concerns the neural origins of relative values. In essence, the question is: *How is $\rho$ determined in the brain?*

To appreciate this issue, consider in our experiments choices between two juices offered in equal ranges. In this case, $\rho$ is a number. In general, $\rho$ can assume any value and we can conventionally set $\rho \geq 1$ (i.e., A is preferred to B). In principle, the relative value between two goods could be induced by differences in the activity of different groups of *offer value* cells. For example, if the animal choosing between juices A and B is indifferent between 1A and 3B, one might expect to observe that the firing rate of *offer value A* cells measured when the animal is offered 1A is 3 times as large as that of *offer value B* cells measured when the animal is offered 1B. However, experimental results indicate that this is not the case – in fact, the activity range of any given cell was found to be independent of the relative value of two juices (Padoa-Schioppa 2009). In other words, relative values do not simply reflect differences in the activity of *offer value* cells. This fact highlights an important point, namely that economic decisions cannot be conceptualized as simple comparisons of neuronal firing rates. So how are relative values determined in the brain?

We examined this issue in the framework of Wang's model. The original W11 is symmetric in A and B. If the activity of OV cells does not depend on the intrinsic preference for the corresponding juice, and if the two value ranges are equal, the symmetry of the network implies $\rho = 1$. However, non-trivial relative values (i.e., $\rho > 1$) can be imposed introducing an imbalance in the synaptic efficacies linking the various pools of neurons associated with the two juices. We experimented introducing such imbalance at different stages of the network. Fig. 5a illustrates the choices patterns obtained in the same simulation illustrated in Fig.4. The x- and y-axes represent the quantities of juice A and B offered, and the z-axis represents the proportion of trials in which juice B was chosen. Each gray dot represents one trial type, and the color surface shows the result of a 2-dimensional logistic regression. Specifically, to examine departures from linearity, we used a logistic model including all second-order terms:
choice \( B = \frac{1}{1 + e^{-X}} \)

\[ X = a_0 + a_1 \#A \, a_2 \#B + a_3 \#A^2 + a_4 \#B^2 + a_5 \#A \, \#B \]  

(25)

In Eq.25, the variable choice \( B \) is the proportion of B choices; \( \#A \) and \( \#B \) are the quantities of juice A and juice B offered, respectively. Fig.5b shows the same surface shown in Fig.5a seen from the z-axis. In this simulation, the synaptic imbalance was introduced at the level of the input current: referring to Eq.19, we set \( \delta J_{\text{stim}} = (2, 1) \). The ensemble of offers for which the model was indifferent between the two juices is termed the indifference function. Notably, the indifference function in this simulation was a straight line through the origin (Fig.5b).

Furthermore, the slope of the indifference function was essentially equal to the synaptic weight ratio = 2. A simplified logistic model including only first-order terms provided the measure \( \rho = a_1/a_2 = 2.03 \). (This value was used to generate the third row of Fig.4.)

In another simulation, we introduced the synaptic imbalance in the recurrent, NMDA-mediated self-excitation of CJ cells (Fig.5c). In this case, referring to Eq.11, we set \( \delta J_{\text{NMDA}} = (1.05, 1) \). (These values were chosen such that the indifference function would cross offers [10A:20B].) Notably the indifference function was no longer a straight line and, more importantly, it was no longer homogenous (it did not cross the origin). This behavior is not realistic because it amounts to stating that the model consistently chooses no juice over small quantities of juice B. We obtained similar results when we imposed the imbalance in the inhibition from interneurons to CJ cells (Fig.5d). In this case, referring to Eq.13, we set \( \delta J_{\text{GABA}} = (1, 1.02) \). Again, the indifference function was non-linear and, most importantly, non-homogenous. We concluded that within Wang's model non-trivial relative values emerge from an imbalance in the input synaptic ratio.

**Dependence on the strength of recurring synapses**

Wang's model is biophysically realistic in the sense that all the parameters represent biophysical quantities (synaptic efficacies, time constants, etc.) and their values are derived from or constrained by experimental measures (with some tuning). In this study, we used the same parameters previously set for perceptual decisions (Wong and Wang 2006). The only (partially) free parameter was the relative strength of recurring synapses \( w^+ \). This parameter is also thought
to characterize different brain regions (Murray et al. 2014). We thus examined how the results described above depended on $w_+$.

In the simulations described so far, we set $w_+ = 1.75$. Fig.6 summarizes the results of three additional simulations, in which we set $w_+ = 1.55$ (Fig.6, top row), $w_+ = 1.70$ (Fig.6, center row) and $w_+ = 1.85$ (Fig.6, bottom row). For each simulation, the two left panels illustrate the activity of CJB cells, and the two right panels illustrate the activity of CV cells. The results obtained for the three simulations are qualitatively similar, but a few quantitative trends can be observed. First, as $w_+$ increased, the sustained, working memory-like delay activity of CJB cells increased (Fig.6a, e, i). Concurrently, the steady state activity of these cells became more binary (Fig.6b, f, j). As described in the next section, the "predictive activity" of CJ cells increased as a function of $w_+$. Finally, the relation between the activity of CV cells and the chosen value was closer to linear for lower values of $w_+$ (Fig.6d, h, l). These trends notwithstanding, the primary observation of these analyses is that the results presented in previous sections held essentially true in a fairly wide range of $w_+$.

**Choice hysteresis and the predictive activity of chosen juice cells**

We next examined whether Wang's model could reproduce a series of empirical phenomena related to the origins of choice variability. Relevant to all these phenomena is the distinction between easy and split decisions. Consider the behavior shown in Fig.1e. For many offer types, away from the indifference point, the animal chooses consistently the same juice (A or B). These decisions are referred to as "easy". For other offer types, closer to the indifference point, the animal split its choices between the two juices. These decisions are referred to as "split".

All other things equal, monkeys in our experiments had the tendency to choose on any given trial the same juice chosen (and received) in the previous trial (Fig.7b). This phenomenon is termed "choice hysteresis" (Padoa-Schioppa 2013). Other analyses showed that the activity of *chosen juice* cells prior to the offer correlated with the eventual decision of the animal – an effect termed "predictive activity" (Padoa-Schioppa 2013). The predictive activity can be observed in Fig.7a. Trials were divided in 4 groups depending on whether the animal chose juice A or juice B and on whether decisions were easy or split. For easy trials, the activity of *chosen juice* cells recorded...
before the offer did not correlate with the eventual decision. In contrast, the pre-offer activity
recorded in split trials correlated with the eventual decision of the animal. Importantly, a large
component of the predictive activity observed in Fig.7a was tail activity from the previous trial.
In a conservative interpretation, this tail activity was correlated with the current decision because
the decision on any given trial was mildly correlated with that in the previous trial. In other
words, a large component of the predictive activity was closely related to choice hysteresis.
However, the predictive activity had also a smaller, "residual" component that did not depend on
the outcome of the previous trial (Padoa-Schioppa 2013).

We now examined whether W11 reproduced these phenomena. First, we considered the same
simulation depicted in Fig.4 and conducted on the activity of CJ cells the same analysis as that
conducted on neuronal data. As illustrated in Fig.7c, we found a consistent predictive activity.
Interestingly, this predictive activity increased as a function of $w_+$ (Fig.6aei). This trends reflects
the fact that baseline fluctuations increase with $w_+$. Also, for higher $w_+$ the decision is made
more rapidly (the network is more "impulsive") and thus the initial bias becomes more relevant
(see below).

The predictive activity in Fig.7c (W11) is noticeably modest compared to that in Fig.7a
(experimental data). Importantly, in all the simulations discussed so far, the initial conditions of
the network were reset at the beginning of each trial. This reset effectively prevented any choice
hysteresis (Fig.7d). As a consequence, the predictive activity observed in Fig.7c corresponds
exclusively to the residual component of the predictive activity observed empirically. To further
investigate choice hysteresis and the predictive activity within W11, we ran a series of
simulations in which the initial conditions of the network were set, in each trial, equal to the final
conditions at the end of the preceding trial. For any value of $w_+$ this policy induced some choice
hysteresis and enhanced the predictive activity. However, these effects were more pronounced
for higher values of $w_+$ (because of the larger tail activity). Fig.7e and Fig.7f illustrate a
simulation in which we set $w_+ = 1.82$. It can be noted that there is a robust choice hysteresis and
that the predictive activity is clearly enhanced (note the different y-axis scales in the inserts of
Fig.7e and Fig.7c).
In conclusion, W11 reproduces both components of the predictive activity, as well as choice hysteresis. These results suggest the following interpretation. The pre-offer activity represents the state of neural assembly prior to the offer, which varies to some extent from trial to trial due to history (tail activity from the previous trial) or stochastic fluctuations. On any given trial, the initial state introduces a small bias. When one of the two offered values dominates (easy trials), the initial bias is irrelevant and the higher value is always chosen. When the two offer values are close (split trials), the initial bias contributes significantly to the decision (predictive activity). This contribution determines a correlation between the pre-offer activity and the decision. Within W11, the tail component of the predictive activity causes choice hysteresis. While consistent with current data, this causal link remains to be tested empirically.

The overshooting of chosen value cells

Another phenomenon observed in the empirical data pertains to chosen value cells. In essence, all other things equal, the activity of these neurons presented a transient but robust "overshooting" when the decision was more difficult (Padoa-Schioppa 2013). For the analysis of neuronal data, we focused on trials in which the animal chose 1A and divided them in easy and split (Fig.8a). In first approximation, the activity recorded for the two groups of trials was the same (same chosen value). However, the activity recorded in split trials presented a transient overshooting in the time window extending 150-400 ms after the offer.

We now repeated this analysis on the activity of CV cells in W11. We examined the same simulation as in Fig. 4 and we focused on trials in which 1A was offered. We divided trials in easy and split, and we computed the two activity profiles. We then repeated this calculation for each quantity of A that induced some split decisions (offer value A = 1...10) and we averaged the activity profiles obtained for different values of A offered. As illustrated in Fig.8b, the model reproduced the activity overshooting. Notably, this result provides a novel interpretation for the activity overshooting. We previously showed that the activity overshooting of chosen value cells could be explained if the relative value of the two juices ($\rho$) fluctuated from trial-to-trial (Padoa-Schioppa 2013). However, the activity overshooting for CV cells was obtained here without introducing any variability in the synaptic efficacies. Of course, this observation does not
exclude that synaptic efficacies do fluctuate from trial to trial, which would provide an additional
correction to the overshooting.

**Range adaptation, context-dependent preferences and Hebbian learning**

The issue of range adaptation and Hebbian learning was addressed elsewhere (Padoa-Schioppa
and Rustichini 2014) and we discuss it here briefly for the sake of completeness.

As described above, the activity of *offer value* cells adapts to the range of values available in any
behavioral context (Padoa-Schioppa 2009). In principle, range adaptation ensures an efficient
representation. At the same time, range adaptation also poses a computational challenge, in the
following sense. First, note that in W11 – and in other decision models (Bogacz et al. 2006;
Krajbich et al. 2010) – decisions are ultimately comparisons of firing rates. Now consider a
session in which offers of juice A and juice B vary in ranges $\Delta A = [0, 3]$ and $\Delta B = [0, 6]$,
respectively, and assume that the relative value is such that $1A = 2B$. If decisions are made by
comparing firing rates, the animal chooses juice A whenever the activity of *offer value A* cells is
higher than that of *offer value B* cells. Now imagine to run a second session in which the range of
juice A remains unchanged ($\Delta A = [0, 3]$) while the range of juice B is doubled ($\Delta B = [0, 12]$).
The activity of *offer value B* cells will adapt to the new value range. If decisions are made by
comparing firing rates, range adaptation will inevitably induce a change of preferences in the
new session such that $1A = 4B$. Interestingly, framing effects described in behavioral economics
show that preferences can depend on the behavioral context in ways qualitatively similar to that
described here (Ariely et al. 2003; Savage 1972; Tversky and Kahneman 1981). At the same
time, it would seem puzzling if preferences could be manipulated so easily and so arbitrarily by
modifying the range of options. Indeed, an experiment in which monkeys chose between two
juices in two subsequent blocks showed that relative values were fairly stable even when the
ratio of offer value ranges ($\Delta A/\Delta B$) was varied by a factor 2 (Conen et al. unpublished
observations). These considerations raise one question: How can the network achieve
(reasonably) stable preferences under varying ranges of offer values?

The issue of preference stability in the presence of range adaptation is closely related to that non-
trivial relative values ($\rho>1$) discussed above. In essence, we propose that the network responds to
the changes in value ranges by altering the synaptic efficacies between OV cells and CJ cells
(Eq.19). As previously shown, this synaptic plasticity can be achieved with a mechanism of
Hebbian learning (Padoa-Schioppa and Rustichini 2014). Framing effects may be reproduced if
this Hebbian learning lags range adaptation of OV cells.

Stable points of the dynamical system

One important question concerns the number steady states of the model. Wong and Wang (2006)
provided a bifurcation diagram, in which the number of attractors and the working-memory
regime were examined as a function of the input firing rate ($\mu_0$) and the parameter $w_+$. Their
analysis was done for the reduced version of the model (W2) and for $coh = 0$. Generating a
bifurcation diagram for W11 is more difficult because the system is time dependent and high
dimensional. As a first step, we conducted a series of simulations to examine the end state of the
network for different sets of offer values (Fig.9). For these simulations we set $\Delta A = \Delta B = [0 15]$
and $\delta J_{stim} = (1, 1)$, such that $\rho = 1$. For comparison, although $w_+$ defined in W11 is not identical
to $w_+$ defined in W2, the value of $w_+$ used here is comparable to values examined by Wong and
Wang (2006). In contrast, stimulus currents are substantially higher in our case. Specifically, in
our simulations we normally used $\Delta J = J_{AMPA, input} / J_{AMPA, ext, pyr} = 30$ (see Table 1). In such
conditions and for the data point at the center of Fig.3 (right panel), the product $J_{AMPA, input} r_{OV}$ is
roughly equal to $100 J_{AMPA, ext, pyr}$ (see Eq.19). In contrast, in the original W11, the equivalent
product was set to $40 J_{AMPA, ext, pyr}$ when $coh = 0$.

Consider Fig.9, panel a2. In this scatter plot, x- and y-axes represent, respectively, the "final"
activity of CJA and CJB cells as measured in the time window 400-600 ms after stimulus on.
Each symbol + represents one trial and symbols are color coded according to the offer type. The
color legend is indicated in the leftmost panel in the same row. Rows 2 and 3 of the figure
illustrate the same data focusing on a subset of trials, with the color legend indicated in the
leftmost column. Several points can be noted. First, the network clearly separated between the
two end points when the value difference was sufficiently high (purple, blue and green symbols
in panels b1 and b2). It did not clearly separate between the end points when offer values were
close (panel b3). In separate simulations, we observed that extending the stimulus in time for 1 s
or increasing the levels of noise in the system did not affect these results (not shown). The
simulation in panels b1-b3 were based on our normal parameters $w_+ = 1.75$ and $\Delta J = 30$. We then repeated the simulations using parameters $w_+ = 1.85$ and $\Delta J = 30$ (panels c1-c3), and $w_+ = 1.75$ and $\Delta J = 15$ (panels d1-d3). The results obtained were qualitatively similar, although the separation for difficult decisions tended to increase when $w_+$ was raised (panel c2) and to decrease when $\Delta J$ was reduced (panel d2).

The formulation of W2 for economic decisions and a more formal analysis of the steady states will be presented elsewhere.

The baseline activity of offer value cells

As illustrated in Fig.1a, offer value cells presented a baseline activity of ~6 sp/s, which is smaller but comparable to the value-related dynamic range (~10 sp/s). Experimental work showed that this baseline activity does not depend on the value range, on the juice preference or, for given juice pair, on the relative value of the juices (Padoa-Schioppa 2009). As we examine the effects of introducing a baseline in the activity of OV cells, two premises are in order. First, the original W11 set the baseline equal to zero – a reasonable approximation since the baseline activity of MT cells is indeed low. Second, the presence of a substantial baseline changes dramatically the input to the network.

In a series of simulations, we tested W11 introducing a realistic baseline. If we consider the situation in which all synaptic efficacies are balanced, the network is robust to the introduction of a baseline. In this condition, even using the same parameters as in the initial simulations, we obtained for the indifference function a straight line through the origin (not shown). Small adjustments to the parameters provided realistic profiles for CJ cells and for CV cells (Fig.10a-f). However, problems arise when one considers the fact that the relative value $\rho$ between two goods should be free to assume any value (see above section Imposing non-trivial relative values) and/or the fact that (only) the value-dependent component of offer value cells activity adapts to the range of offer values (see above section Range adaptation, context-dependent preferences and Hebbian learning). The approach used in the baseline-subtracted case, namely to introduce an imbalance in the synaptic efficacies no longer works. In other words, the indifference function becomes non-homogenous even if the synaptic imbalance is limited to the input (Fig.10g-l). This
is because the synaptic efficacies $\delta J_{stim}$ multiply the entire activity of OV cells, including the
baseline. As a consequence, when $\delta J_{stim, 1} > \delta J_{stim, 2}$, the network chooses zero quantities of juice A
over small quantities of juice B. In conclusion, the baseline activity of offer value cells poses a
challenge for W11. Possible ways to address this issue are discussed below.

**Discussion**

We examined a biophysically realistic model previously proposed to describe the activity of area
LIP during perceptual decisions, namely W11, and we tested the extent to which it could
reproduce the activity recorded in the OFC during economic decisions. Our analysis represented
a challenging test for the model for at least four reasons. (1) Stimuli (i.e., offers) in economic
decisions are intrinsically two-dimensional, whereas stimuli in perceptual decisions are one-
dimensional. (2) The relative value between two given goods is arbitrary. Thus to accommodate
subjective preferences, the network must be flexible. (3) Offer value cells undergo range
adaptation but decisions should, in first approximation, not depend on the range of offer values.
In other words, the network must be adaptable. (4) The activity of neurons in the OFC only
partially resembles that of neurons in area LIP, for which the model was designed. Specifically,
in addition to offer value cells (analogous to MT) and chosen juice cells (analogous to LIP), there
are chosen value cells, which have no known correspondent in perceptual decisions. Moreover,
even the resemblance between chosen juice cells and neurons in LIP is weak, because the
representation of chosen juice cells is categorical while that of LIP is continuous, and because
chosen juice cells do not present the race-to-threshold and the working memory activity the
model was originally designed to reproduce. In spite of all these challenges, the spirit of this
study was not to design a new model. Rather, we tested Wang's model as much as possible off-
the-shelf, without modifying its structure or even adjusting its parameters. We did, however,
adapt the network in two important ways. First, we strengthened the connections between OV
cells and CJ cells (reflecting the fact that offer value cells and chosen juice cells are found in the
same area). Second, we introduced mechanisms of synaptic plasticity to account for the required
flexibility and adaptability of the model.
Several traits make W11 a particularly attractive model. First, the model is biophysically realistic. Elements in the extended, spiking network are neurons endowed with realistic synaptic connections and time constants, and the mean-field approach preserves the realism of the model. Second, the model is nearly without free parameters. In other words, the model is very constrained, especially when tested outside its original domain. Third, the model makes testable predictions on the excitatory/inhibitory nature of different neuronal populations and on their connectivity. These traits set W11 apart from other computational models of economic decisions including the drift-diffusion models, mutual inhibition models and the leaky competing accumulator model (Bogacz et al. 2006; Hare et al. 2011; Krajbich et al. 2010; Usher and McClelland 2001). Furthermore, the explicit inclusion of inhibitory interneurons in W11 provides an account for chosen value cells, which are not explained in more schematic models including the drift-diffusion model and the leaky competing accumulator model. Other studies examined Wang's model in the context of value-based decisions. In particular, Behrens and colleagues used the simplified W2 version to generate aggregate regressors for the analysis of MEG data (Hunt et al. 2012; Jocham et al. 2012). In this respect, we note that the tests ran here were significantly more stringent because we matched each node in the network with a specific group of neurons, because the original analysis of neuronal data (Padoa-Schioppa and Assad 2006) tested a large number of variables and not only variables generated from the model, and because we reproduced a variety of empirical findings.

We found that W11 provides a remarkably accurate account for the activity of neurons in the OFC. The input node of the model (OV cells) can be identified with offer value cells. The model naturally generates binary decisions, and the activity profile of the output node (CJ cells) is fairly similar to that of chosen juice cells. Perhaps most surprisingly, the activity of inhibitory interneurons (CV cells) is very similar to that of chosen value cells (more on this below). In addition, W11 reproduces several phenomena related to the neuronal origins of choice variability, namely choice hysteresis, the predictive activity of chosen juice cells and the activity overshooting of chosen value cells. We examined what changes in the network are necessary to ensure that relative values are arbitrary and to accommodate the fact that offer value cells undergo range adaptation. We found that these two requirements are addressed only when synaptic plasticity is introduced in the connections between OV cells and CJ cells. We also found
that W11 falls short of the experimental data in at least two ways. First, the model includes only
neurons with positive encoding (i.e., higher firing rate for higher values). In contrast, for each of
the three variables, a substantial fraction of the neurons in OFC presents negative encoding (i.e.,
higher firing rate for lower values) (Padoa-Schioppa 2013; 2009). Second, the model can provide
the flexibility necessary to accommodate any relative value, or it can handle a significant
baseline activity in the input node (OV cells), but it cannot do both of these things at the same
time. As discussed below, these limitations appear conceptually surmountable, although more
theoretical work is necessary in this respect.

Chosen value signals and interneurons in the OFC

One of the main results of this study is that interneurons of W11 were found to encode the
chosen value. This observation was unforeseen and somewhat extraordinary. Interneurons were
included in the progenitor models of W11 for biological realism and network stability (Amit and
Brunel 1997) – not to reproduce empirical observations analogous to chosen value cells.
Furthermore, chosen value cells have no known correspondent in perceptual decision tasks. In
contrast, chosen value signals have been observed in numerous studies of value-based decisions
and in multiple brain regions (Amemori and Graybiel 2012; Cai et al. 2011; Cai and Padoa-
Schioppa 2014; 2012; Grabenhorst et al. 2012; Lau and Glimcher 2008; Lee et al. 2012; Padoa-
Schioppa and Assad 2006; Roesch et al. 2009; Strait et al. 2014; Sul et al. 2010; Wunderlich et al.
2010). While it is clear that chosen value signals could inform a variety of mental functions
including associative learning, visual attention, emotion and others, the possible contributions of
chosen value signals to economic decisions have remained mysterious. The present results
suggest that chosen value cells in the OFC may be directly involved in the decision. This
suggestion is bolstered by the fact that CV cells in the model reproduce the activity overshooting
seen in chosen value cells. Importantly, the hypothesis that chosen value cells in the OFC
participate in the decision does not imply that all chosen value signals in the brain are the
signature of a decision process. Indeed, once computed and explicitly represented by a neuronal
population, the variable chosen value could be transmitted broadly to other brain regions and
thus facilitate various mental functions.
Considering the similarity between CV cells and *chosen value* cells from a different angle, W11 makes the strong prediction that all *chosen value* cells in the decision circuit are interneurons and that all *chosen juice* cells are pyramidal cells. Future experiments will test these predictions. Importantly, the present model provides an account for the decision circuit. However, OFC likely comprises additional neuronal populations that do not directly intervene in the decision but receive input from the decision circuit. Thus testing the predictions of the model will require distinguishing between neurons that contribute to the decisions and other neurons within OFC. Notably, the hypothesis that the variable *chosen value* is first computed in OFC and then transmitted to other brain regions implies that this variable is passed from the decision circuit to pyramidal cells in layers 5 and 6 of OFC, where cortico-cortical projections originate. In principle, NS pyramidal cells in W11 (whose activity resembles that of *chosen value* cells with low dynamic range) could serve that purpose, although alternate schemes are also possible.

In its current form, W11 presents two clear limitations – the fact that the model only includes neurons with positive encoding and the challenge posed by the baseline activity of *offer value* cells. Negative encoding is an intriguing phenomenon, partly because it is not observed in sensory systems (to our knowledge). More detailed neurophysiology work is necessary to establish the excitatory/inhibitory nature of neurons with negative encoding, whether these neurons are preferentially located in specific cortical layers, and whether they present specific patterns of connectivity. From a modeling perspective, the presence of cells with negative encoding can be viewed as a degree of freedom present in the empirical data but not used by the model. In principle, this degree of freedom could help resolve the challenge posed by the baseline activity in *offer value* cells, because two inputs encoding the same variable with opposite sign (excitatory and inhibitory) could be combined in a way that reduces or even eliminates any baseline. In the light of these considerations, we view W11 as a benchmark and a starting point for biophysically realistic models of economic decisions. The next steps are to examine empirically the excitatory/inhibitory nature of different groups of cells in OFC, to assess possible differences between cortical layers, and to establish the actual connectivity between the different groups of neurons (a set of challenging tasks!). The results of these enquiries should then inform new and more accurate neuro-computational models.
To conclude, Wang's model provides a biologically credible account for the neuronal mechanisms of economic decisions. In the model, decisions are generated by three groups of cells whose activity closely resembles the activity of the three groups of neurons previously found in the OFC. This close resemblance does not demonstrate that economic decisions take place in the OFC, or that the decision mechanisms are based on recurrent excitation and pooled inhibition. However, the model does provide an important proof of concepts that economic decisions could emerge from the activity of neurons identified in this area. In this framework, W11 suggests that chosen value cells are a key component of the decision process. The model makes several non-trivial predictions that require further empirical testing. Last but not least, Wang's model represents an important bridge between neuroscience and economics, providing a platform to investigate the implications of neuronal data for economic theory. We will examine some of these implications in future work.
**Table 1. Parameters used in simulations.**

<table>
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<th>Network parameters (same as in W11)</th>
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<tr>
<td>$N_E$</td>
<td>1,600</td>
</tr>
<tr>
<td>$N_I$</td>
<td>400</td>
</tr>
<tr>
<td>$C_{ext}$</td>
<td>800</td>
</tr>
<tr>
<td>$F$</td>
<td>0.15</td>
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<tr>
<td>$r_{ext}$</td>
<td>3.0 sp/s</td>
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<th>Time constants, synaptic efficacies and noise (same as in W11)</th>
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<tr>
<td>$\tau_{AMPA}$</td>
<td>2 ms</td>
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<td>$\tau_{NMDA}$</td>
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</tr>
<tr>
<td>$\tau_{GABA}$</td>
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<td>$J_{AMPA, \text{ext, pyr}}$</td>
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<td>$J_{GABA, \text{rec, pyr}}$</td>
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<td>$J_{AMPA, \text{ext, in}}$</td>
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<td>$J_{AMPA, \text{rec, in}}$</td>
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<td>$J_{GABA, \text{rec, in}}$</td>
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<td>$\gamma$</td>
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<td>$\sigma$</td>
<td>0.020 A</td>
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<th>Parameters of input-output function for integrate-and-fire neurons (same as in W11)</th>
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<td>$I_E$</td>
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<td>$g_E$</td>
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<td>$c_E$</td>
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<tr>
<td>$I_I$</td>
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<tr>
<td>$g_I$</td>
<td>0.087</td>
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<tr>
<td>$c_I$</td>
<td>615</td>
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<table>
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<th>Parameters used to model OV cells</th>
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<td>$r_0$</td>
<td>0 or 6 sp/s</td>
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<tr>
<td>$\Delta r$</td>
<td>8 sp/s</td>
</tr>
<tr>
<td>$a$</td>
<td>$t_{offer} + 175$ ms</td>
</tr>
<tr>
<td>$b$</td>
<td>30 ms</td>
</tr>
<tr>
<td>$c$</td>
<td>$t_{offer} + 400$ ms</td>
</tr>
<tr>
<td>$d$</td>
<td>100 ms</td>
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<tr>
<td>$J_{AMPA, \text{input}}$</td>
<td>$30 \times J_{AMPA, \text{ext, pyr}}$</td>
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Figure legends

Figure 1. Summary of experimental results. a. Neuronal response encoding the offer value B. In the left panel, the x-axis represents different offer types. Black dots represent the percent of trials in which the animal chose juice B and red symbols indicate the neuronal firing rate. Each symbol represents one trial type. Diamonds and circle indicate trials in which the animal chose juice A and juice B, respectively. In the right panel, the same neuronal response (y-axis) is plotted against the encoded variable (x-axis). It can be noted that the encoding is close to linear. b. Activity profile for offer value cells. The plot illustrates the population activity. For each neuron, trials were divided in three groups (tertiles) based on the value of the encoded juice (low, medium, high). A trace was computed for each group of by averaging the activity across trials, and the resulting traces were averaged across the population. Only cells with positive encoding are shown here. c. Neuronal response encoding the chosen juice. Same conventions as in panel a. In this case, the activity is roughly binary depending on the juice chosen by the animal. d. Activity profile for chosen juice cells. In this case, trials were divided depending on whether the animal chose the juice encoded by the cell (E chosen) or the other juice (O chosen). Here the "encoded" juice was defined as that which elicited higher activity. Traces are population averages. e. Neuronal response encoding the chosen value. Same conventions as in panel a. f. Activity profile for chosen value cells. In this case, trials were divided in three groups (tertiles) depending on the chosen value (low, medium, high). Traces are population averages. The figure is adapted from (Padoa-Schioppa 2013).

Figure 2. Schematics of W11. Excitatory and inhibitory connections are indicated in blue and red, respectively. Different arrow heads indicate synaptic efficacies between different groups of cells (see figure legend and corresponding numerical values in Table 1).

Figure 3. Illustration of inputs for random dot task (left) and economic choice task (right).

Figure 4. Initial test of Wang's model. a. OVB cells, activity profile. Trials were divided in three groups (tertiles) depending on the offer value B (same convention as in Fig.1b). b. OVB cells, tuning curve. The firing rate (z-axis) is plotted as a function of the two offers (x- and y-axes). Each data point represents one trial type. Red diamonds and blue circles represent trial types in
which the network chose juice A and juice B, respectively. c. OVB cells, reduced tuning curve. Down-sampled version of the data shown in b, displayed in the format of Fig.1a. d. OVB cells, activity versus encoded variable. Same firing rates shown in panel b plotted as a function of the encoded variable (offer value B). efgh. Activity of CJB cells. In panel e, trials were divided in two groups depending on the chosen juice (same convention as in Fig.1d). ijkl. Activity of CV cells. In panel i, trials were divided in three groups (tertiles) depending on the chosen value (same conventions as in Fig.1f). In all panels, red diamonds (blue circles) represent choices of juice A (juice B).

Figure 5. Choice patterns obtained for different balances of synaptic efficacies. a. Choice pattern for the simulation illustrated in Fig.4. Each data point (gray dots) represents one offer type. The percent of B choices (z-axis) is plotted as a function of the two offers (x- and y-axes). The surface illustrates the result of a logistic fit including all second order terms (Eq.1). b. Same surface as in panel a viewed from the z-axis. The indifference function, roughly corresponding to the green pixels (see color legend) is a straight line through the origin. In this simulation, $\delta_{\text{stim}} = (2, 1)$ and all other synaptic weights were balanced. c. Choice pattern obtained with an imbalanced NMDA-mediated reverberation. d. Choice pattern obtained with an imbalanced inhibition. In both panels c and d, parameters were chosen such that the model was roughly indifferent between 10A and 20B. In both cases, the indifference function did not cross the origin.

Figure 6. Robustness with respect to the strength of recurring synapses. abcd. Summary of results obtained with $w_+ = 1.55$. Panels a and b summarize the results obtained for CJB cells (same format as in Fig.4ef). Panels c and d summarize the results obtained for CV cells (same format as in Fig.4il). efgh. Summary of results obtained with $w_+ = 1.70$. ijkl. Summary of results obtained with $w_+ = 1.85$.

Figure 7. Predictive activity and choice hysteresis. a. Choice hysteresis (experimental data). In this representative session, trials were divided in two groups depending on whether the outcome of the previous trial was juice A (A• trials) or juice B (B• trials). It can be observed that the relative value of juice A was higher in A• trials (dark gray) compared to B• trials (light gray). In other words, all other things equal, the animal had a tendency to choose on any given trial the same juice as in the previous trial. b. Predictive activity of chosen juice cells (experimental data).
c. Predictive activity of CJ cells (model). Same simulation as in Fig.4. The activity immediately before and after the offer is enlarged in the insert. It can be noted that the predictive activity is significant but relatively modest compared to that measured experimentally (see main text).

d. Lack of choice hysteresis in initial simulations. Same simulation as in panel c and Fig.4. In this simulation, the initial conditions were reset at the beginning of each trial. In the analysis, trials were divided in A• trials and B• trials depending on the previous outcome, and the two data sets were separately fitted with a logistic function. The two fits are completely overlapping, indicating the lack of choice hysteresis. In other words, the predictive activity in panel c corresponds to the "residual" predictive activity measured in OFC (see main text).

ef. Predictive activity in the presence of choice hysteresis. In this simulation, we set the initial conditions in each trial equal to the final conditions in the previous trial (tail activity) and $w_+ = 1.82$. To ensure that each juice was chosen in ~50% of trials, we set in this simulation $\Delta A = [0, 10]$ and $\Delta B = [0, 20]$. Under these conditions, the predictive activity is much larger (compare panels e and c) and there is a robust choice hysteresis (panel f). In panel f, the two logistic fit refer to A• trials and B• trials, respectively. Panels a and b are reproduced from (Padoa-Schioppa 2013).

Figure 8. Activity overshooting. a. Overshooting of chosen value cells (experimental data). Only trials in which the animal chose 1A are included in the figure. See main text for details. b. Overshooting of CV cells (model). Same simulation as in Fig.4. For each quantity of A that induced some split decisions (i.e., offer value $A = 1...10$), trials in which the network chose juice A were divided in easy and split (see main text). The activity overshooting was observed for each value of A. The main plot here illustrates the effect averaged across values of A. The insert illustrates the effect for offer value $A = 8$. Panel a is reproduced from (Padoa-Schioppa 2013).

Figure 9. Stable points of W11. The leftmost column (panels a1-a3) represents the space of possible offers (as in Fig.3, right panel). The other three columns represent the results of three simulations, with $w_+ = 1.75$ and $\Delta J = 30$ (normal parameters, panels b1-b3), $w_+ = 1.85$ and $\Delta J = 30$ (panels c1-c3) and $w_+ = 1.75$ and $\Delta J = 15$ (panels d1-d3). In each panel, the two axes represent the "final" activity of CJA cells (y-axis) and CJB cells (x-axis) (time window 400-600 ms after the offer). Each data point represents one trial and data points are color-coded according to the legend in the corresponding leftmost column (see main text).
Figure 10. Effects of non-zero baseline in OV cells. a-f. Symmetric network. Panel a shows the simulated activity of OVB cells, panels be show the activity of CJB cells, panels de show the activity of CV cells and panel f shows the choice pattern. The presence of the baseline activity in OV cells changes dramatically the nature of the input. However, with small adjustments, the symmetric network is robust to this change. In this simulation, we set $w_+ = 1.65$, $J_{AMPA, \text{rec, in}} = 0.9$ $J_{AMPA, \text{rec, in}}$, $J_{NMDA, \text{rec, in}} = 0.9$, $J_{NMDA, \text{rec, in}}$, $J_{GABA, \text{rec, in}} = 0.8$, $J_{GABA, \text{rec, in}}$. The activity profiles of CJ and CV cells remain comparable to the empirical observations, and CV cells encode the chosen value. g-l. Asymmetric network. In this simulation, we set $\delta J_{\text{stim}} = (1.2, 1)$ all other parameters as in panels a-f. Although the activity profiles of CJ cells (panels hi) and CV cells (panels j) are realistic, the indifference function is non-homogenous. Tests in which we tried to recover this departure from homogeneity introducing imbalances at other stages of the network did not provide successful results.


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Figure 1

a) Offer value cells

b) Average firing rate (sp/s)

0% 100%
offer value B

0B:1A 1B:3A 1B:2A 1B:1A 2B:1A 3B:1A 4B:1A 6B:1A

R² = 0.92

1A = 3.1B

1B = 1.9B

0A 1B 2B 3B 4B 5B 6B 7B 8B 9B

Firing rate (sp/s)

offer

1A = 3.1B

R² = 0.92

1B = 1.9B

0A 1B 2B 3B 4B 5B 6B 7B 8B 9B

Firing rate (sp/s)

offer

1B = 1.9B

1B = 1.9B

0A 1B 2B 3B 4B 5B 6B 7B 8B 9B

Firing rate (sp/s)

offer

1B = 1.9B

R² = 0.95

1B = 1.9B

0A 1B 2B 3B 4B 5B 6B 7B 8B 9B

Firing rate (sp/s)

offer

1B = 1.9B

R² = 0.95

0B:1A 1B:3A 1B:2A 1B:1A 2B:1A 3B:1A 4B:1A 6B:1A

10B:1A 6B:1A 2B:0A

1A = 3.2B

R² = 0.90

1A = 3.2B

1A = 3.2B

0B:1A 1B:3A 1B:2A 1B:1A 2B:1A 3B:1A 4B:1A 6B:1A

10B:1A 6B:1A 2B:0A

chosen juice

offer

chosen value

offer

chosen value

offer

500 ms
Figure 2
Figure 3

random dot task

![Random Dot Task Graph](image)

economic choice task

![Economic Choice Task Graph](image)
Figure 4

OVB cells

a

Firing rate (sp/s)

b

A chosen

B chosen

0

5

10

15

offer A

offer B

OVB cells

CJB cells

e

Firing rate (sp/s)

f

A chosen

B chosen

0

5

10

15

offer A

offer B

CJB cells

CV cells

i

Firing rate (sp/s)

j

A chosen

B chosen

0

5

10

15

offer A

offer B

CV cells
Figure 6

CJB cells

A chosen
B chosen

CV cells

high
medium
low

A chosen
B chosen

w+ = 1.55

offer A
offer B

Firing rate (sp/s)

offer

w+ = 1.70

offer A
offer B

Firing rate (sp/s)

offer

w+ = 1.85

offer A
offer B

Firing rate (sp/s)

offer

CV cells

A chosen
B chosen

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer

CV cells

offer A
offer B

Firing rate (sp/s)

offer
Figure 8

(a) chosen value cells

(b) CV cells

Trials: A chosen
- easy
- split

Firing rate (sp/s) vs. offer
Figure 9

w+ = 1.75, ∆J = 30

w+ = 1.85, ∆J = 30

w+ = 1.75, ∆J = 15

offer A

offer B

w+ = 1.75,

w+ = 1.85,