Attractor Neural States: A Brain-Inspired Complementary Approach to Reinforcement Learning

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Abstract: It is widely accepted that reinforcement learning (RL) mechanisms are optimal only if there is a predefined set of distinct states that are predictive of reward. This poses a cognitive challenge as to which events or combinations of events could potentially predict reward in a non-stationary environment. In addition, the computational discrepancy between two families of RL algorithms, model-free and model-based RL, creates a stability-plasticity dilemma, which in the case of interactive and competitive multiple brain systems poses a question of how to guide optimal decision-making control when there is competition between two systems implementing different types of RL methods. We argue that both computational and cognitive challenges can be met by infusing the RL framework as an algorithmic theory of human behavior with the strengths of the attractor framework at the level of neural implementation. Our position is supported by the hypothesis that ‘attractor states’ which are stable patterns of self-sustained and reverberating brain activity, are a manifestation of the collective dynamics of neuronal populations in the brain. Hence, when neuronal activity is described at an appropriate level of abstraction, simulations of spiking neuronal populations capture the collective dynamics of the network in response to recurrent interactions between these populations.

1 INTRODUCTION

In machine learning and other artificial intelligence (AI) related disciplines, the theory of reinforcement learning (RL) provides an algorithmic account for gaining optimal action control in sequential decision-making processes when only limited feedback is available (Sutton and Barto, 1998; Daw et al., 2005; Lewis and Vrabie, 2009; van Otterlo and Wiering, 2012; Krigolson et al., 2014; Marsland, 2015). In the corresponding fields of cognitive science and psychology, RL describes the practice by which animals and humans probe reward contingencies while acting in a novel environment (Schultz et al., 1997; Doya, 2007; Niv and Montague, 2008; Shteingart et al., 2013).

Despite indisputable advances in RL research over the past two decades, two challenges still remain: a computational and a cognitive one (Gershman and Daw, 2017). One accumulating evidence from cognitive science and brain research suggests two quite different conceptual frameworks for thinking about learning (Gallistel and King, 2009). Though methods of both frameworks involve experience, they differ in the way and extent to which they impose computational load so as to achieve a high level performance (Dayan and Berridge, 2014). In the first framework, termed as model-free RL, learning is the ability of a plastic brain to modify itself (by experience) in order to operate more efficiently in a novel environment. Such a modification affects both the structure and the function of the brain (Davidson and Begley, 2012; Phelps et al., 2014). Synonym terms for this type of RL approach are habitual learning, retrospective revaluation, and reflexive decision-making (Dolan and Dayan, 2013). The second framework, called model-based RL, conceptualizes learning as the process of deducing (also from experience) structural characteristics of the operating environment in order to shape subsequent behavior (Hamid, 2015). Informing behavior takes place in that the derived information is carried forward in memory (Gallistel and King, 2009). Synonym terms for this type of RL approach are goal-directed behavior, prospective planning, and reflective decision-making (Friedel et al., 2015). In the human brain, mechanisms of model-
free RL were linked to corticostriatal circuits, involving ventral striatum and regions of the amygdala (Balleine et al., 2007), whereas the prefrontal cortex was found to be the substrate for model-based RL mechanisms (Mante et al., 2013). Though each of the two frameworks was successfully tested in its own domain, current research points out some of the challenging issues in rounding out the theory of RL as an integrative account of human behavior.

Computational considerations, too, are the focus of extensive comments on RL methods. Specifically, comparing the apparently simplistic laboratory conditions with the relatively more complex real-world decision-making scenarios, as encountered by biological agents in their natural environments, a number of technical factors turn out to be exceptionally decisive when considering the computational load that should be imposed by any of the RL learning paradigms on one side and the biological world on the other side (Dayan and Berridge, 2014). While laboratory experiments assume mainly low-dimensional, discrete, and almost fully treatable state spaces, real-world situations have high-dimensional, continuous and partially observable state spaces. This implies that the learning algorithm has to struggle with minimal sets of data and low frequency of probed situations, as it is almost impossible for a real-world situation to be encountered twice with all its circumstantial characteristics (Gershman and Daw, 2017). Moreover, the discrepancy between the two frameworks of RL in the way they perform computations creates a dilemma, the stability-plasticity dilemma, when state-reward contingencies are reversed (Mermillod et al., 2013). It turns out, although context-specificity is beneficial for learning in novel stationary environments, it is detrimental for flexibility in non-stationary environments. These inspections point to a fundamental limitation of the RL framework.

In the following, we shall be arguing that the above limitations can, nevertheless, be resolved by integrating the 'attractor' framework as a complementary approach at the level of neural implementation to the behavioral success of RL theory at the algorithmic level. Our position is supported by the hypothesis that 'attractor states', which are stable patterns of self-sustained and reverberating brain activity, are a manifestation of the collective dynamics of neuronal populations in the brain (Amit et al., 1994; Hopfield, 1982). Hence, when neuronal activity is described at an appropriate level of abstraction, simulations of spiking neuronal populations capture the collective dynamics of the network in response to recurrent interactions between these populations (Braun and Mattia, 2010). As an illuminating example for potency, we shall consider the field of associative learning.

The remainder of the paper is organized as follows. In Section 2 we introduce the theory of reinforcement learning including a formal description of the underlying Markov decision processes and the two major classes of RL algorithms. Section 3 points out to current challenges in linking human behavior to existing RL algorithms. Section 4 discusses how models in the attractor framework could complement the RL framework, accounting for the current cognitive challenges. We finally conclude and portray our plans for future work in Section 5.

2 RL: THEORY & FORMALISM

Historically, RL has its origin in mathematical psychology and operations research (Dayan and Niv, 2008). Inspired by the psychological literature on Pavlovian (classical) and instrumental conditioning, Richard Sutton developed, together with Andrew Barto, algorithms for agent-based learning that later became the core ideas for the theory of RL (Sutton and Barto, 1998). Parallel to their research, yet in a separate line, Dimitri Bertsekas and John Tsitsiklis, two electrical engineers working in the field of operations research, developed stochastic approximations to dynamic programming that allow a system to learn about its behavior through simulation (experience) and improve its performance through iterative reinforcement (Bertsekas and Tsitsiklis, 1996). These lines of research marked the emergence of RL as an algorithmic theory for optimal decision making on the basis of behavior and subsequent effects (Niv and Montague, 2008).

2.1 RL and Markov Decision Processes

In a typical RL setting, a goal-directed agent (this could be a natural or an artificial system) interacts with an environment via embedded sets of sensors and actuators. The sensors provide the agent with information about the state of the environment, whereas the actuators enable the agent to act upon the environment, causing its current state to change. As a consequence, the agent receives reinforcement in terms of a numerical signal that describes how close (or far) it moved to (or away from) its predefined goal (Sutton and Barto, 1998; van der Ree and Wiering, 2013; Castro-González et al., 2014). As the agent moves along the several states of the environment, the corresponding sequence of action-reward combinations moves on, too.
Decisions in RL can be modeled as a Markov decision process (MDP). When the state of the operating environment, however, is subject to inherent uncertainty, modeling takes the form of a partially observable Markov decision process (POMDP) (Kaelbling et al., 1996; Sutton and Barto, 1998). Formally, an MDP process consists of two functions, $R$ and $T$, defined over two sets, $S$ and $A$. The functions represent the resulting rewards and state transitions, whereas the two sets describe the available states and effective actions, respectively.

In an MDP process, the environment evolves stochastically under simple discrete temporal dynamics. At time $t$, the environment is in state $s_t = s$. The agent chooses some action $a_t \in A$, expecting to reap a certain reward $r$ (expected outcome). Nevertheless, it experiences the actual consequence of its choice (either immediately or later on) in terms of a numerical reinforcement $r \in \mathbb{R}$ (actual outcome). Subsequently, the state of the environment changes its instance into $s_{t+1} = s'$ at the next time step $t + 1$. Moreover, the agent updates its knowledge about reward contingencies within the operating environment as a reflection on its very experience. To denote the probability of moving from state $s$ into $s'$ when taking action $a$, we write $T(s,a,s')$. Analogously, the notion $R(s,a,r)$ refers to the probability of receiving a reward at state $s$, when taking action $a$. Note that formalizing the reward and transition functions in terms of the current state rather than the entire history of the environment, a characteristic referred to as Markov property, provides a computational advantage, for it requires the learning algorithm to remember and work with the parameters that are related only to the current state. This is definitely easier than dealing with all previous states of the environment (Maia, 2009; Hamid, 2014).

### 2.2 Model-Free and Model-Based RL

Based on the way they optimize their learning and decision-making processes, RL methods can be sorted into two main classes: model-free and model-based methods (Dayan and Berridge, 2014).

Though both use experience, model-free RL algorithms assume no a priori knowledge of the MDP but learn a state-action value function, known as the ‘value function’ (Dayan, 2008). One of the successful implementations of model-free RL methods is the well-known temporal difference (TD) learning algorithm. It utilizes a reward-prediction error, which is the discrepancy between the actual and expected rewards, to ‘cache’ actually observed information about the long-term rewarding potencies of the probed actions. This approach represents a computationally simple way to exploit experience, for the model needs only to learn one or two simple quantities (state/action values). However, it is statistically less efficient, because the cached information is stored as a scalar quantity without connecting outcomes to their direct causes in a distinguishable manner (Dayan and Niv, 2008). Consequently, the model’s performance is most likely to suffer from two shortcomings. First, the model cannot (later on) extricate insights about rewards or transitions from the cached value. Second, the cached information intermixes previous estimates or beliefs about state values regardless their sometimes erroneous valence. As a result, model-free RL methods lack an appropriately quick adaptation to sudden changes in reward contingencies (Hamid, 2015). Because of this characteristic, model-free RL was proposed as the underlying model for habitual controllers, in which actions are presumably based on habits (Daw et al., 2005). This key characteristic links model-free RL to corticostriatal circuits involving, in particular, the ventral striatum and regions of the amygdala in the human’s brain (Packard and Knowlton, 2002; Balleine et al., 2007; Dayan and Balleine, 2002).

Model-based RL is a family of algorithms that generate goal-directed choices by utilizing an explicit model of the underlying MDP process. This sums up representations of the environment, expectations, and prospective calculations to make cognitive predictions of future values (Daw et al., 2005; Dayan and Berridge, 2014). Specifically, model-based RL algorithms capture the dynamics of the MDP in terms of state transition probabilities. Such probabilities can be presented as a tree connecting short-term predictions about immediate outcomes of each action in an arbitrary sequence of actions. Deciding which action is more beneficial can then be done by exploring branching sets of possible future situations. There are several ‘tree search’ techniques that can do this (Daw et al., 2005). It turns out that exploiting experience in the case of model-based RL is more efficient than in model-free RL for two reasons. First, it provides more statistical reliability, especially when storing the sometimes unrelated morsels of information. Second, and importantly, it ensures more flexibility in terms of adaptive planning, which becomes necessary when changes occur in the learning environment. Hence, model-based RL accounts best for goal-directed behavior that contains more cognitive planning. This key characteristic links model-based RL to the prefrontal cortex in the primate’s brain (Owen, 1997).


3 COGNITIVE CHALLENGES

Besides the above mentioned computational considerations, RL faces cognitive challenges as well. On one hand, the theory could successfully explain several characteristics of human and animal learning, e.g., blocking (Kamin, 1969), overshadowing (Reynolds, 1961), and inhibitory conditioning (Rescorla and Lollordo, 1968). It also proved able to predict new phenomena such as over-expectation (Kremer, 1978) and managed to account for the relatively touchy ‘secondary conditioning’: a phenomenon in which a predictor of a predictor servers as a predictor (Dayan and Abbott, 2005). However, it still suffers from difficulties in accounting for some human instrumental learning behaviors. For example, analysis of the stock market suggests that a positively surprising obtained pay-off and a negatively surprising forgone payoffs trigger a reversal of choice behavior rather than repeating that behavior according to the recency assumption (Nevo and Erev, 2012).

Furthermore, consistent with the idea of interactive and competitive multiple memory systems in the brain (Poldrack and Packard, 2003), recent research addressed the question of how to guide optimal decision-making control in the face of a running competition between two systems, each of which implements a different type of RL methods (Dolan and Dayan, 2013).

Another challenge concerns the architecture of the theory itself. Specifically, it has been argued that RL mechanisms are optimal only if there is a predefined set of distinct states that are predictive of reward (Doya, 2007). This implies, for an agent behaving in an RL fashion to achieve successfully optimal action control, it is necessary to define the states such that they contain all reward-relevant information, even if task-irrelevant (Hamid, 2015). But the question becomes: how do such states emerge in the brain or more generally within the decision-making component of the agent in the first place (Rigotti et al., 2010)?

In a recent study of conditional associative learning, the authors accounted for the effect of temporal order on accelerating the learning of arbitrary associations by devising a model-free RL rule that sets a probabilistic response choice, reflecting reward expectations that have been accumulated in the form of ‘action values’ (Hamid et al., 2010). The reinforcement rule modifies these ‘action values’ in proportion to the reward-prediction error that corresponds to the chosen response. The key feature of the devised model is that ‘action values’ are expanded in time: some attach to the object of the current trial and others attach to objects of preceding trials. This provides them with a cumulative effect in the sense that the more ‘action values’ favor a particular response, the more likely this response is chosen. Accordingly, when successive objects appear in a consistent order, more than one ‘action value’ will favor the correct response, which will therefore be chosen more frequently.

Though the model could account qualitatively and quantitatively to the behavioral observations, it failed to account for the same associative task in a reversal learning paradigm (Hamid and Braun, 2010). The main goal of the reversal paradigm was to test the model’s key assumption, i.e., the reinforcement of pairings between past stimuli and present response. Specifically, let \( S_{t-2} \) and \( S_{t-1} \) be the visual stimuli presented at trials \( t-2 \) and \( t-1 \) in figure 1, respectively. \( S_t \) is the target stimulus at trial \( t \) with motor response \( R_t \). A: learned response \( R_t \) for the target stimulus is replaced by response \( R_{t'} \) in the second run of the object sequence (‘action reversal’). Before reversal, the model reinforces, in addition to the paring \( (S_t \rightarrow R_t) \), the pairings \( (S_{t-1} \rightarrow R_t) \) and \( (S_{t-2} \rightarrow R_t) \). After reversal, however, these pairings become invalid, as the model has to learn the new response \( R_{t'} \). Hence, the model’s performance is expected to fall to chance level. B: target stimulus \( S_t \) is replaced by stimulus \( S_{t'} \), which has the same response as that of \( S_t \). Before reversal, the model reinforces the pairings \( (S_t \rightarrow R_t), (S_{t-1} \rightarrow R_t), \) and \( (S_{t-2} \rightarrow R_t) \). These pairings remain valid after reversal. Hence, predicted performance remains above chance level.

The cognitive experiment was conducted using mixed sequences of visual objects as presented in figure 1 with type A, B, and C objects similar to experiment 2 in (Hamid et al., 2010). Specifically, Thirty two fractal objects were used to create sequences of 72 trials. Eight of these objects were recurring. Four of the recurring objects formed two consistent pairs \((5,6)\) and \((7,8)\), each of which appeared six times in the sequence. The ‘predecessor’ objects \((5,7)\) were termed type A and the ‘successor’ objects \((6,8)\) type B. Four additional recurring objects were used to form twelve random pairs \((1,2), (1,3), (1,4), \ldots, (4,1), (4,2), (4,3)\), each appearing once per sequence (type C). Random pairs and consistent pairs were alternated and separated by 24 one-time objects to form sequences of 72 trials. Human subjects learned by trial and error to associate each visual object with one of four possible motor responses: up, down, left, or right. Each of the 72 trials long temporal sequences was presented twice without interruption making up new sequences of 144 trials long. In the ‘action reversal’ condition, target objects were associated with a
new motor response in the second half of the temporal sequence. In the ‘object reversal’ condition, target objects were replaced by other objects in the second half of the temporal sequence. Figure 2 illustrates behavioral and modeling results of the cognitive experiment. We can briefly summarize the results as follows. First, contrary to the predictions of our reinforcement model (Hamid et al., 2010), any type of reversal reduced performance to chance level. Second, the rate of recovery seemed to differ between reversal types, appearing to be faster for an ‘object reversal’ than for an ‘action reversal’.

4 MODELS IN THE ATTRACTOR FRAMEWORK

It is widely accepted that reinforcement mechanisms are optimal only if there is a predefined set of distinct states that are predictive of reward (Sutton and Barto, 1998; Daw et al., 2005; Doya, 2007; Niv and Montague, 2008; Dayan and Niv, 2008). Thus, reinforcement models beg the question as to which events or combinations of events could potentially predict reward in a non-stationary environment. This brings us to the crucial question of how our brain selects and creates neural representations for potentially reward-predicting events. An interesting approach to this question is the attractor framework, which postulates that the formation of such representations is based on temporal statistics of the environment. The key idea is that mental representations are realized by stable patterns of reverberating activity, which are stable steady-states (‘attractors’) in the neural dynamics of the network (Hopfield, 1982; Amit et al., 1997; Fusi et al., 2005).

The central tenets of attractor theory are that (i) the network is plastic, that is, connection strengths develop in an activity-driven, Hebbian manner and that (ii) associations (e.g., stimulus-response pairings) are maintained as self-sustained, persistent patterns of activity that represent attractors of the neural dynamics. These tenets predict the formation of associative links whenever a set of events occurs repeatedly in a consistent temporal order (Griniasty et al., 1993; Amit, 1995; Brunel, 1996).

A recent study of behavioral flexibility in reversal situations exemplifies the attractor framework (Rigotti et al., 2010). The authors of this study postulate
two neural circuits, one for learning reward-relevant conditional associations (‘associative network’) and another for observing temporal contingencies (‘context network’). The interaction between these two networks leads to the formation of distinct neural representations for different contexts. More specifically, the associative network comprises two populations of excitatory neurons, which represent alternative stimulus-response associations. One population represents the stimulus-response associations appropriate for one context, whereas the other population codes the appropriate associations for another context. The two excitatory populations compete through a third, inhibitory population. As long as the reward predictions of one population are fulfilled, the currently dominant population will continue to suppress the other population, and new stimuli will be evaluated in the light of the experience encoded in the dominant population. However, when predicted rewards fail to materialize, the other population may gain ascendancy and behavior may now be governed by the experience accumulated in another, alternative context.

So how can a representation of context be formed, which can link all the stimulus-response associations that are rewarded in a particular context? The key idea is that different stimulus-response associations become linked on the basis of temporal statistics. Specifically, as long as one context holds for much longer than one trial, stimulus-response associations within this context follow each other more frequently than stimulus-response associations in diff-
fterent contexts. This correlational difference can be translated by Hebbian mechanisms into selective meta-associations among the stimulus-response associations of a given context. Mechanistically, the formation of these meta-associations relies on the temporal overlap between the representation of a current stimulus-response association and lingering representations of stimulus-response associations in the recent past. Further details can be found in (Rigotti et al., 2010).

5 CONCLUSIONS

Although most attempts to test the attractor framework experimentally have used single-unit recordings in behaving, non-human primates, we believe that this framework makes some predictions even at the behavioral level. For example, the neurophysiological findings of (Miyashita, 1988) and (Yakovlev et al., 1998) imply that reverberative delay activity exists only after an attractor representation has formed. In the context of (Hamid et al., 2010), this suggests that lingering representations of past events are available only after these past events have become familiar. On this basis, we would expect that the presence of consistent predecessor objects becomes influential only after these objects have become familiar and are recognized. Accordingly, it would be an interesting extension of the present study to examine whether the facilitative effect of temporal context is conditional on correct performance with regard to predecessor objects.

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REFERENCES


